

# Package ‘synthpop’

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**Type** Package

**Title** Generating Synthetic Versions of Sensitive Microdata for  
Statistical Disclosure Control

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**Date** 2018-06-27

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**Description** A tool for producing synthetic versions of microdata containing confidential information so that they are safe to be released to users for exploratory analysis. The key objective of generating synthetic data is to replace sensitive original values with synthetic ones causing minimal distortion of the statistical information contained in the data set. Variables, which can be categorical or continuous, are synthesised one-by-one using sequential modelling. Replacements are generated by drawing from conditional distributions fitted to the original data using parametric or classification and regression trees models. Data are synthesised via the function `syn()` which can be largely automated, if default settings are used, or with methods defined by the user. Optional parameters can be used to influence the disclosure risk and the analytical quality of the synthesised data. For a description of the implemented method see Nowok, Raab and Dibben (2016) <doi:10.18637/jss.v074.i11>.

**License** GPL-2 | GPL-3

**Depends** lattice, MASS, methods, nnet, ggplot2

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synthpop-package	<i>Generating synthetic versions of sensitive microdata for statistical disclosure control</i>
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## Description

Generate synthetic versions of a data set using parametric or CART methods.

## Details

Package: synthpop  
 Type: Package  
 Version: 1.4-4  
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 License: GPL-2 | GPL-3

Synthetic data are generated from the original (observed) data by the function `syn`. The package includes also tools to compare synthetic data with the observed data (`compare.synds`) and to fit (generalized) linear model to synthetic data (`lm.synds`, `glm.synds`) and compare the estimates with those for the observed data (`compare.fit.synds`). More extensive documentation with illustrative examples is provided in the package vignette.

### Author(s)

Beata Nowok, Gillian M Raab, Joshua Snoke and Chris Dibben based on package **mice** (2.18) by Stef van Buuren and Karin Groothuis-Oudshoorn

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### References

Nowok, B., Raab, G.M and Dibben, C. (2016). synthpop: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).

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compare

*Comparison of synthesised and observed data*

---

### Description

A generic function for comparison of synthesised and observed data. The function invokes particular methods which depend on the class of the first argument.

### Usage

```
compare(object, data, ...)
```

### Arguments

<code>object</code>	a synthetic data object of class <code>synds</code> or <code>fit.synds</code> .
<code>data</code>	an original observed data set.
<code>...</code>	additional arguments specific to a method.

### Details

Compare methods facilitate quality assessment of synthetic data by comparing them with the original observed data sets. The data themselves (for class `synds`) or models fitted to them (for class `fit.synds`) are compared.

### Value

The value returned by `compare` depends on the class of its argument. See the documentation of the particular methods for details.

### See Also

[compare.synds](#), [compare.fit.synds](#)

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 compare.fit.synds

*Compare model estimates based on synthesised and observed data*


---

## Description

The same model that was used for the synthesised data set is fitted to the observed data set. The coefficients with confidence intervals for the observed data is plotted together with their estimates from synthetic data. When more than one synthetic data set has been generated (`object$m>1`) combining rules are applied. Analysis-specific utility measures are used to evaluate differences between synthetic and observed data.

## Usage

```
## S3 method for class 'fit.synds'
compare(object, data, plot = "Z",
        print.coef = FALSE, return.plot = TRUE, plot.intercept = FALSE,
        lwd = 1, lty = 1, lcol = c("#1A3C5A", "#4187BF"),
        dodge.height = .5, point.size = 2.5, incomplete = FALSE,
        population.inference = FALSE, ci.level = 0.95, ...)

## S3 method for class 'compare.fit.synds'
print(x, print.coef = x$print.coef, ...)
```

## Arguments

<code>object</code>	an object of type <code>fit.synds</code> created by fitting a model to synthesised data set using function <code>glm.synds</code> or <code>lm.synds</code> .
<code>data</code>	an original observed data set.
<code>plot</code>	values to be plotted: "Z" (Z scores) or "coef" (coefficients).
<code>print.coef</code>	a logical value determining whether tables of estimates for the original and synthetic data should be printed.
<code>return.plot</code>	a logical value indicating whether a confidence interval plot should be returned.
<code>plot.intercept</code>	a logical value indicating whether estimates for intercept should be plotted.
<code>lwd</code>	the line type.
<code>lty</code>	the line width.
<code>lcol</code>	line colours.
<code>dodge.height</code>	size of vertical shifts for confidence intervals to prevent overlapping.
<code>point.size</code>	size of plotting symbols used to plot point estimates of coefficients.
<code>incomplete</code>	a logical value indicating whether the method of Reiter (2003) for what he terms partially synthetic data should be used for inference. It requires multiple syntheses with <code>m</code> greater than the number of coefficients, ideally at least 5 more.

population.inference	a logical value indicating whether intervals for inference to population quantities, as described by Karr et al. (2006), should be calculated and plotted. This option suppresses the lack-of-fit test and the standardised differences since these are based on differences standardised by the original interval widths.
ci.level	Confidence interval coverage as a proportion.
...	additional parameters passed to <code>ggplot</code> .
x	an object of class <code>compare.fit.synds</code> .

### Details

This function can be used to evaluate whether the method used for synthesis is appropriate for the fitted model. If this is the case the estimates from the synthetic data of what would be expected from the original data  $x_{pct}(\text{Beta})$   $x_{pct}(Z)$  should not differ from the estimates from the observed data (Beta and Z) by more than would be expected from the standard errors ( $se(\text{Beta})$  and  $se(Z)$ ). For more details see the vignette on inference.

### Value

An object of class `compare.fit.synds` which is a list with the following components:

call	the original call to fit the model to the synthesised data set.
coef.obs	a data frame including estimates based on the observed data: coefficients (Beta), their standard errors ( $se(\text{Beta})$ ) and Z scores (Z).
coef.syn	a data frame including (combined) estimates based on the synthesised data: point estimates of observed data coefficients (B.syn), standard errors of those estimates ( $se(\text{B.syn})$ ), estimates of the observed standard errors ( $se(\text{Beta})$ .syn), Z scores estimates (Z.syn) and their standard errors ( $se(\text{Z.syn})$ ). Note that $se(\text{B.syn})$ and $se(\text{Z.syn})$ give the standard errors of the mean of the $m$ syntheses and can be made very small by increasing $m$ (see the vignette on inference for more details).
coef.diff	a data frame containing standardized differences between the coefficients estimated from the original data and those calculated from the combined synthetic data. The difference is standardized by dividing by the estimated standard error of the fit from the original. The corresponding p-values are calculated from a standard Normal distribution and represent the probability of achieving differences as large as those found if the model use for synthesis is compatible with the model that generated the original data.
mean.abs.std.diff	Mean absolute standardized difference (over all coefficients).
ci.overlap	a data frame containing the percentage of overlap between the estimated synthetic confidence intervals and the original sample confidence intervals for each parameter. When <code>population.inference = TRUE</code> overlaps are calculated as suggested by Karr et al. (2006). Otherwise a simpler overlap measure with intervals of equal length is calculated.
mean.ci.overlap	Mean confidence interval overlap (over all coefficients).

lack.of.fit	lack-of-fit measure from all $m$ synthetic data sets combined, calculated as follows, when <code>incomplete.method = FALSE</code> . The vector of mean differences ( <code>diff</code> ) between the coefficients calculated from the synthetic and original data provides a standardised lack-of-fit = $t(\text{diff}) \%*\% V^{-1} t(\text{diff})$ , where $\%*\%$ represents the matrix product and $V^{-1}$ is the inverse of the variance-covariance matrix for the mean coefficients from the synthetic data. If the model used to synthesize the data is correct this quantity, which is a Mahalanobis distance measure, will follow a chi-squared distribution with degrees of freedom, and thus expectation, equal to the number of parameters ( $p$ ) in the fitted model. When <code>incomplete.method = TRUE</code> the function above follows a Hotelling's $T^2$ distribution and the lack-of-fit statistic is referred to an $F(p, m - p)$ .
lof.pvalue	p-value for the combined lack-of-fit test of the NULL hypothesis that the method used for synthesis retains all relationships between variables that influence the parameters of the fit.
ci.plot	ggplot of the the coefficients with confidence intervals for models based on observed and synthetic data. If <code>return.plot</code> was set to <code>FALSE</code> then <code>ci.plot</code> is NULL.
print.coef	a logical value determining whether tables of estimates for the original and synthetic data should be printed.
m	the number of synthetic versions of the original (observed) data.
ncoef	the number of coefficients in the fitted model (including an intercept).
incomplete	whether methods for incomplete synthesis due to Reiter (2003) have been used in calculations.
population.inference	whether intervals as described by Karr et al. (2016) have been calculated.

## References

- Karr, A., Kohnen, C.N., Oganian, A., Reiter, J.P. and Sanil, A.P. (2006). A framework for evaluating the utility of data altered to protect confidentiality. *The American Statistician*, **60**(3), 224-232.
- Nowok, B., Raab, G.M and Dibben, C. (2016). synthpop: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).
- Reiter, J.P. (2003) Inference for partially synthetic, public use microdata sets. *Survey Methodology*, **29**, 181-188.

## See Also

[summary.fit.synds](#)

## Examples

```
ods <- SD2011[,c("sex", "age", "edu", "smoke")]
s1 <- syn(ods, m = 3)
f1 <- glm.synds(smoke ~ sex + age + edu, data = s1, family = "binomial")
compare(f1, ods)
compare(f1, ods, print.coef = TRUE, plot = "coef")
```

---

compare.synds	<i>Compare univariate distributions of synthesised and observed data</i>
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---

## Description

Compare synthesised data set with the original (observed) data set using percent frequency tables and histograms. When more than one synthetic data set has been generated (`object$m>1`), by default pooled synthetic data are used for comparison.

## Usage

```
## S3 method for class 'synds'
compare(object, data, vars = NULL, msel = NULL,
        breaks = 20, nrow = 2, ncol = 2, rel.size.x = 1,
        cols = c("#1A3C5A", "#4187BF"), ...)

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

## Arguments

<code>object</code>	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn()</code> and it includes <code>object\$m</code> synthesised data set(s).
<code>data</code>	an original (observed) data set.
<code>vars</code>	variables to be compared. If <code>vars</code> is <code>NULL</code> (the default) all synthesised variables are compared.
<code>msel</code>	index or indices of synthetic data copies for which a comparison is to be made. If <code>NULL</code> pooled synthetic data copies are compared with the original data.
<code>breaks</code>	the number of cells for the histogram.
<code>nrow</code>	the number of rows for the plotting area.
<code>ncol</code>	the number of columns for the plotting area.
<code>rel.size.x</code>	a number representing the relative size of x-axis labels.
<code>cols</code>	bar colors.
<code>...</code>	additional parameters.
<code>x</code>	an object of class <code>compare.synds</code> .

## Details

Missing data categories for numeric variables are plotted on the same plot as non-missing values. They are indicated by `miss.` suffix.

## Value

An object of class `compare.synds` which is a list including a list of comparative percent frequency tables (`tables`) and a `ggplot` object (`plots`) with bar charts/histograms. If multiple plots are produced they and their corresponding frequency tables are stored as a list.

## References

Nowok, B., Raab, G.M and Dibben, C. (2016). synthpop: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).

## Examples

```
ods <- SD2011[ , c("sex", "age", "edu", "marital", "ls", "income")]
s1 <- syn(ods)
compare(s1, ods, vars = "ls")
compare(s1, ods, vars = "income")
```

---

glm.synds, lm.synds     *Fitting (generalized) linear models to synthetic data*

---

## Description

Fits generalized linear models or simple linear models to the synthesised data set(s) using [glm](#) and [lm](#) function respectively.

## Usage

```
glm.synds(formula, family = "binomial", data, ...)
lm.synds(formula, data, ...)

## S3 method for class 'fit.synds'
print(x, msel = NULL, ...)
```

## Arguments

formula	a symbolic description of the model to be estimated. A typical model has the form response $\sim$ predictors. See the documentation of <a href="#">glm</a> and <a href="#">formula</a> for details.
family	a description of the error distribution and link function to be used in the model. See the documentation of <a href="#">glm</a> and <a href="#">family</a> for details.
data	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <a href="#">syn</a> and it includes <code>data\$m</code> synthesised data set(s).
...	additional parameters passed to <a href="#">glm</a> or <a href="#">lm</a> .
x	an object of class <code>fit.synds</code> .
msel	index or indices of synthetic data copies for which coefficient estimates are to be displayed. If NULL (default) the combined (average) coefficient estimates are printed.



**Value**

An object of class `fit.synds`. It is a list with the following components:

<code>call</code>	the original call to <code>glm.synds</code> or <code>lm.synds</code> .
<code>mcoefavg</code>	combined (average) coefficient estimates.
<code>mvaravg</code>	combined (average) variance estimates of <code>mcoef</code> .
<code>analyses</code>	<code>summary.glm</code> or <code>summary.lm</code> object respectively or a list of <code>m</code> such objects.
<code>fitting.function</code>	function used to fit the model.
<code>n</code>	a number of cases in the original data.
<code>k</code>	a number of cases in the synthesised data.
<code>proper</code>	a logical value indicating whether synthetic data were generated using proper synthesis.
<code>m</code>	the number of synthetic versions of the observed data.
<code>method</code>	a vector of synthesising methods applied to each variable in the saved synthesised data.
<code>mcoef</code>	a matrix of coefficients estimates from all <code>m</code> syntheses.
<code>mvar</code>	a matrix of variance estimates from all <code>m</code> syntheses.

**See Also**

[glm,lm,multinom.synds,compare](#)

**Examples**

```
### Logit model
ods <- SD2011[1:1000, c("sex", "age", "edu", "marital", "ls", "smoke")]
s1 <- syn(ods, m = 3)
f1 <- glm.synds(smoke ~ sex + age + edu + marital + ls, data = s1, family = "binomial")
f1
print(f1, msel = 1:2)

### Linear model
ods <- SD2011[1:1000, c("sex", "age", "income", "marital", "depress")]
ods$income[ods$income == -8] <- NA
s2 <- syn(ods, m = 3)
f2 <- lm.synds(depress ~ sex + age + log(income) + marital, data = s2)
f2
print(f2, 1:3)
```

---

 multi.compare

*Multivariate comparison of synthesised and observed data*


---

### Description

Graphical comparisons of a variable (`var`) in the synthesised data set with the original (observed) data set within subgroups defined by the variables in a vector `by`. `var` can be a factor or a continuous variable and the plots produced will depend on the class of `var`. The variables in `by` will usually be factors or variables with only a few values.

### Usage

```
multi.compare(object, data, var = NULL, by = NULL, msel = NULL,
  barplot.position = "fill", cont.type = "hist", y.hist = "count",
  boxplot.point = TRUE, binwidth = NULL, ...)
```

### Arguments

<code>object</code>	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn()</code> and it includes <code>object\$m</code> synthesised data set(s).
<code>data</code>	an original (observed) data set.
<code>var</code>	variable to be compared between observed and synthetic data within subgroups.
<code>by</code>	variables to be tabulated or cross-tabulated to form groups.
<code>barplot.position</code>	type of barplot. The default "fill" gives a single bar with the proportions in each group while "dodge" gives side-by-side bars with the numbers in each category.
<code>cont.type</code>	default "hist" gives histograms and "boxplot" gives boxplots.
<code>y.hist</code>	defines y scale for histograms - "count" is default; "density" gives proportions.
<code>boxplot.point</code>	default (TRUE) adds individual points to boxplots.
<code>msel</code>	numbers of synthetic data sets to be used - must be numbers in the range <code>1 : object\$m</code> - defaults to <code>1 : object\$m</code>
<code>binwidth</code>	sets width of a bin for histograms.
<code>...</code>	additional parameters that can be supplied to <a href="#">ggplot</a> .

### Value

Plots as specified above. A table of the numbers in the subgroups is printed to the R console.

### See Also

[compare.synds](#), [compare.fit.synds](#)

**Examples**

```
### default synthesis of selected variables
vars <- c("sex", "age", "edu", "smoke")
ods <- na.omit(SD2011[1:1000, vars])
s1 <- syn(ods)

### categorical var
multi.compare(s1, ods, var = "smoke", by = c("sex","edu"))

### numeric var
multi.compare(s1, ods, var = "age", by = c("sex"), y.hist = "density", binwidth = 5)
multi.compare(s1, ods, var = "age", by = c("sex", "edu"), cont.type = "boxplot")
```

---

multinom.synds

*Fitting multinomial models to synthetic data*


---

**Description**

Fits multinomial models to the synthesised data set(s) using the `multinom` function.

**Usage**

```
multinom.synds(formula, data, ...)
```

**Arguments**

formula	a symbolic description of the model to be estimated. A typical model has the form <code>response ~ predictors</code> . See the documentation of <code>multinom</code> and <code>formula</code> for details.
data	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn</code> and it includes <code>data\$m</code> synthesised data set(s).
...	additional parameters passed to <code>multinom</code> .

**Value**

An object of class `fit.synds`. It is a list with the following components:

call	the original call to <code>multinom.synds</code> .
mcoefavg	combined (average) coefficient estimates.
mvaravg	combined (average) variance estimates of <code>mcoef</code> .
analyses	an object summarising the fit to each synthetic data set or a list of <code>m</code> such objects. Note that this is different from the object created by <code>summary.multinom</code> to make it compatible with other fitting methods. In particular the coefficients are vectors, not matrices.
fitting.function	function used to fit the model.

n	a number of cases in the original data.
k	a number of cases in the synthesised data.
proper	a logical value indicating whether synthetic data were generated using proper synthesis.
m	the number of synthetic versions of the observed data.
method	a vector of synthesising methods applied to each variable in the saved synthesised data.
mcoef	a matrix of coefficients estimates from all m syntheses.
mvar	a matrix of variance estimates from all m syntheses.

**See Also**

[multinom](#), [glm.synds](#), [compare](#)

**Examples**

```
ods <- SD2011[1:1000, c("sex", "age", "edu", "marital", "ls", "smoke")]
s1 <- syn(ods, m = 3)
f1 <- multinom.synds(edu ~ sex + age, data = s1)
summary(f1)
print(summary(f1, msel = 1:2))
compare(f1,ods)
```

---

read.obs

*Importing original data sets form external files*

---

**Description**

Imports data data sets form external files into a data frame. Currently supported files include: sav (SPSS), dta (Stata), xpt (SAS), csv (comma-separated file), tab (tab-delimited file) and txt (delimited text files). For SPSS, Stata and SAS it uses functions from the `foreign` package with some adjustments where necessary.

**Usage**

```
read.obs(file, convert.factors = TRUE, lab.factors = FALSE,
export.lab = FALSE, ...)
```

**Arguments**

file	the name of the file (including extension) which the data are to be read from.
convert.factors	a logical value indicating whether variables with value labels in Stata and SPSS should be converted into R factors with those levels.

lab.factors	a logical value indicating whether variables with complete value labels but imported using their numeric codes ( <code>convert.factors = FALSE</code> ) should be converted from numeric to factor variables.
export.lab	a logical variable indicating whether labels from SPSS or Stata should be exported to an external file.
...	additional parameters passed to read functions.

**Value**

A data frame with an imported data set. For SPSS, Stata and SAS it has attributes with labels.

**See Also**

[write.syn](#)

---

replicated.uniques	<i>Replications in synthetic data</i>
--------------------	---------------------------------------

---

**Description**

Determines which unique units in the synthesised data set(s) replicates unique units in the original observed data set.

**Usage**

```
replicated.uniques(object, data, exclude = NULL)
```

**Arguments**

object	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn()</code> and it includes <code>object\$m</code> synthesised data set(s).
data	the original observed data set.
exclude	a single string or a vector of strings with name(s) of variable(s) to be excluded from the identification of uniques.

**Value**

A list with the following components:

replications	a vector (for <code>object\$m = 1</code> ) or a data frame with <code>object\$m</code> columns (for <code>object\$m &gt; 1</code> ) with logical values indicateing duplicates in <code>m</code> th synthetic data set.
no.replications	a single number or a vector of <code>object\$m</code> integers indicating the number of duplicates in the synthetic data set(s).
no.uniques	a number of unique individuals in the original data set.
per.replications	a single number or a vector of <code>object\$m</code> numeric values indicating the percentage of duplicates in the synthetic data set(s).

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

```
ods <- SD2011[1:1000,c("sex","age","edu","marital","smoke")]
s1 <- syn(ods, m = 2)
replicated.uniques(s1,ods)
```

SD2011

*Social Diagnosis 2011 - Objective and Subjective Quality of Life in Poland*

**Description**

Sample of 5,000 individuals from the Social Diagnosis 2011 survey; selected variables only.

**Usage**

```
SD2011
```

**Format**

A data frame with 5,000 observations on the following 35 variables:

**sex** Sex

**age** Age of person, 2011

**agegr** Age group, 2011

**placesize** Category of the place of residence

**region** Region (voivodeship)

**edu** Highest educational qualification, 2011

**eduspec** Discipline of completed qualification

**socprof** Socio-economic status, 2011

**unempdur** Total duration of unemployment in the last 2 years (in months)

**income** Personal monthly net income

**marital** Marital status

**mmarr** Month of marriage

**ymarr** Year of marriage

**msepddiv** Month of separation/divorce

**ysepddiv** Year of separation/divorce

**ls** Perception of life as a whole

**depress** Depression symptoms indicator

**trust** View on interpersonal trust  
**trustfam** Trust in own family members  
**trustneigh** Trust in neighbours  
**sport** Active engagement in some form of sport or exercise  
**nofriend** Number of friends  
**smoke** Smoking cigarettes  
**nociga** Number of cigarettes smoked per day  
**alcabuse** Drinking too much alcohol  
**alcsol** Starting to use alcohol to cope with troubles  
**workab** Working abroad in 2007-2011  
**wkabdur** Total time spent on working abroad  
**wkabint** Plans to go abroad to work in the next two years  
**wkabintdur** Intended duration of working abroad  
**emcc** Intended destination country  
**englang** Knowledge of English language  
**height** Height of person  
**weight** Weight of person  
**bmi** Body mass index

### Note

Please note that the original variable names have been changed to make them more self-explanatory. Some variable labels have been adjusted as well.

### Source

Council for Social Monitoring. Social Diagnosis 2000-2011: integrated database. <http://www.diagnoza.com/index-en.html> [downloaded on 13/12/2013]

### References

Czapinski J. and Panek T. (Eds.) (2011). Social Diagnosis 2011. Objective and Subjective Quality of Life in Poland - full report. Contemporary Economics, Volume 5, Issue 3 (special issue) <http://ce.vizja.pl/en/issues/volume/5/issue/3#art254>

### Examples

```

spineplot(englang ~ agegr, data = SD2011, xlab = "Age group", ylab = "Knowledge of English")
boxplot(income ~ sex, data = SD2011[SD2011$income != -8,])

```

sdc

*Tools for statistical disclosure control (sdc)***Description**

Labeling and removing unique replicates of unique actual (observed) individuals.

**Usage**

```
sdc(object, data, label = NULL, rm.replicated.uniques = FALSE,
    uniques.exclude = NULL, recode.vars = NULL, bottom.top.coding = NULL,
    recode.exclude = NULL, smooth.vars = NULL)
```

**Arguments**

<code>object</code>	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn()</code> and it includes <code>object\$<i>m</i></code> synthesised data set(s).
<code>data</code>	the original (observed) data set.
<code>label</code>	a single string with a label to be added to the synthetic data sets as a new variable to make it clear that the data are synthetic/fake.
<code>rm.replicated.uniques</code>	a logical value indicating whether unique replicates of units that are unique also in the original data set should be removed.
<code>uniques.exclude</code>	a single string or a vector of strings with name(s) of variable(s) to be excluded from the identification of uniques.
<code>recode.vars</code>	a single string or a vector of strings with name(s) of variable(s) to be bottom-or/and top-coded.
<code>bottom.top.coding</code>	a list of two-element vectors specifying bottom and top codes for each variable in <code>recode.vars</code> . If there is no need for bottom or top coding NA should be used. If only one variable is to be recoded, codes can be given as a two-element vector.
<code>recode.exclude</code>	a list specifying for each variable in <code>recode.vars</code> values to be excluded from recoding, e.g. missing data codes. If all values should be considered for recoding NA should be used. If only one variable is to be recoded, code(s) can be given as a single number or a vector.
<code>smooth.vars</code>	a single string or a vector of strings with name(s) of numeric variable(s) to be smoothed ( <a href="#"><code>smooth.spline</code></a> function is used).

**Value**

An object provided as an argument adjusted in accordance with the other parameters' values.

**See Also**

[replicated.uniques](#)



**Examples**

```
ods <- SD2011[1:1000,c("sex","age","edu","marital","income")]
s1 <- syn(ods, m = 2)
s1.sdc <- sdc(s1, ods, label="false_data", rm.replicated.uniques = TRUE,
  recode.vars = c("age","income"),
  bottom.top.coding = list(c(20,80),c(NA,2000)),
  recode.exclude = list(NA,c(NA,-8)))
```

---

summary.fit.synds      *Inference from synthetic data*

---

**Description**

Combines the results of models fitted to each of the `m` synthetic data sets.

**Usage**

```
## S3 method for class 'fit.synds'
summary(object, population.inference = FALSE, msel = NULL,
  incomplete = FALSE, real.varcov = NULL, ...)

## S3 method for class 'summary.fit.synds'
print(x, ...)
```

**Arguments**

<code>object</code>	an object of class <code>fit.synds</code> created by fitting a model to synthesised data set using function <a href="#">glm.synds</a> or <a href="#">lm.synds</a> .
<code>population.inference</code>	a logical value indicating whether inference should be made to population quantities. If <code>FALSE</code> inference is made to the results that would be expected from an analysis of the original data. This option should be selected if the synthetic data are being used for exploratory analysis, but the final published results will be obtained by running code on the original confidential data. If <code>population.inference = TRUE</code> results would allow population inference to be made from the synthetic data. In both cases the inference will depend on the synthesising model being correct, but this can be checked by running the same analysis on the real data, see <a href="#">compare.fit.synds</a> .
<code>msel</code>	index or indices of the synthetic datasets (1, . . . , <code>m</code> ), for which summaries of fitted models are to be produced. If <code>NULL</code> (default) only the summary of combined estimates is produced.
<code>incomplete</code>	a logical value indicating whether inference is to use the method proposed by Reiter (2003) for what he terms partially synthetic data. This method is valid for any synthesis, but requires multiple syntheses. It is only necessary, when the dependent variable in a model is not completely synthesised and it only makes any difference to the results when <code>population.inference = TRUE</code> .

<code>real.varcov</code>	the estimated variance-covariance matrix of the fit of the model to the original data. This parameter is used in the function <code>compare.fit.synds</code> which has the original data as one of its parameters.
<code>...</code>	additional parameters.
<code>x</code>	an object of class <code>summary.fit.synds</code> .

## Details

The mean of the estimates from each of the  $m$  synthetic data sets yields asymptotically unbiased estimates of the coefficients if the observed data conform to the distribution used for synthesis. The standard errors are estimated differently depending whether inference is made for the results that we would expect to obtain from the observed data or for the parameters of the population that we assume the observed data are sampled from. The standard errors also differ according to whether synthetic data were produced using simple or proper synthesis (for details see Raab et al. (2017)).

## Value

An object of class `summary.fit.synds` which is a list with the following components:

<code>call</code>	the original call to <code>glm.synds</code> or <code>lm.synds</code> .
<code>proper</code>	a logical value indicating whether synthetic data were generated using proper synthesis.
<code>population.inference</code>	a logical value indicating whether inference is made to population coefficients or to the results that would be expected from an analysis of the original data (see above).
<code>incomplete</code>	a logical value indicating whether inference is to use the method proposed by Reiter (2003) for what he terms partially synthetic data.
<code>fitting.function</code>	function used to fit the model.
<code>m</code>	the number of synthetic versions of the original (observed) data.
<code>coefficients</code>	a matrix with combined estimates. If inference is required to the results that would be obtained from an analysis of the original data, ( <code>population.inference = FALSE</code> ) the coefficients are given by <code>xpct(Beta)</code> , the standard errors by <code>xpct(se.Beta)</code> and the corresponding Z-statistic by <code>xpct(Z)</code> . If the synthetic data are to be used to make inferences to population quantities ( <code>population.inference = TRUE</code> ), the coefficients are given by <code>Beta.syn</code> , their standard errors by <code>se.Beta.syn</code> and the Z-statistic by <code>Z.syn</code> (see vignette on inference for more details).
<code>n</code>	a number of cases in the original data.
<code>k</code>	the number of cases in the synthesised data. Note that if $k$ and $n$ are not equal and <code>population.inference = FALSE</code> (the default), then the standard errors produced will estimate what would be expected by an analysis of the original data set of size $n$ .
<code>analyses</code>	<code>summary.glm</code> or <code>summary.lm</code> object respectively or a list of $m$ such objects.
<code>mset</code>	index or indices of synthetic data copies for which summaries of fitted models are produced. If <code>NULL</code> only a summary of combined estimates is produced.

## References

- Nowok, B., Raab, G.M and Dibben, C. (2016). synthpop: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).
- Raab, G.M., Nowok, B. and Dibben, C. (2017). Practical data synthesis for large samples. *Journal of Privacy and Confidentiality*, **7**(3), 67-97. Available at: <http://repository.cmu.edu/jpc/vol7/iss3/4>
- Reiter, J.P. (2003) Inference for partially synthetic, public use microdata sets. *Survey Methodology*, **29**, 181-188.

## See Also

[compare.fit.synds](#), [summary](#), [print](#)

## Examples

```
ods <- SD2011[1:1000,c("sex","age","edu","ls","smoke")]

### simple synthesis
s1 <- syn(ods, m = 5)
f1 <- glm.synds(smoke ~ sex + age + edu + ls, data = s1, family = "binomial")
summary(f1)
summary(f1, population.inference = TRUE)

### proper synthesis
s2 <- syn(ods, m = 5, method = "parametric", proper = TRUE)
f2 <- glm.synds(smoke ~ sex + age + edu + ls, data = s2, family = "binomial")
summary(f2)
summary(f2, population.inference = TRUE)
```

---

summary.synds

*Synthetic data object summaries*

---

## Description

Produces summaries of the synthesised variables. When more than one synthetic data set has been generated (`object$m>1`), by default summaries are calculated by averaging summary values for all synthetic data copies (see `mse1` argument).

## Usage

```
## S3 method for class 'synds'
summary(object, mse1 = NULL, maxsum = 7,
digits = max(3, getOption("digits")-3), ...)

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

**Arguments**

object	an object of class <code>synds</code> ; a result of a call to <code>syn</code> .
mselect	index or indices of synthetic data copies for which a summary is desired. If NULL (default) summaries are calculated by averaging summary values for all synthetic data copies.
maxsum	integer, indicating how many levels should be shown for factors.
digits	integer, used for number formatting with <code>format</code> .
...	additional arguments passed to <code>summary</code> .
x	an object of class <code>summary.synds</code> .

**Details**

See `summary` for more details.

**Value**

An object of class `summary.synds`, which is a list with the following components:

m	the number of synthetic versions of the original (observed) data.
mselect	index or indices of synthetic data copies for which a summary is produced. If NULL summaries are calculated by averaging summary values for all synthetic data copies.
method	a vector of synthesising methods applied to each variable in the saved synthesised data.
result	a table or a list of tables (if more than one synthetic data set is selected) with summaries of synthesised variables.

**References**

Nowok, B., Raab, G.M and Dibben, C. (2016). `synthpop`: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).

**See Also**

[summary,print](#)

**Examples**

```
s1 <- syn(SD2011[,c("sex","age","edu","marital")], m = 3)
summary(s1)
summary(s1, mselect = c(1,3))
```

---

 syn *Generating synthetic data sets*


---

**Description**

Generates synthetic version(s) of a data set. `syn.strata` performs stratified synthesis.

**Usage**

```
syn(data, method = vector("character", length = ncol(data)),
     visit.sequence = (1:ncol(data)), predictor.matrix = NULL,
     m = 1, k = nrow(data), proper = FALSE, minnumlevels = -1,
     maxfaclevels = 60, rules = NULL, rvalues = NULL,
     cont.na = NULL, semicont = NULL, smoothing = NULL,
     event = NULL, denom = NULL, drop.not.used = FALSE, drop.pred.only = FALSE,
     default.method = c("normrank", "logreg", "polyreg", "polr"), models = FALSE,
     print.flag = TRUE, seed = "sample", ...)
```

```
syn.strata(data, strata = NULL,
            minstratumsize = 10 + 10 * length(visit.sequence),
            tab.strataobs = TRUE, tab.stratasyn = FALSE,
            method = vector("character", length = ncol(data)),
            visit.sequence = (1:ncol(data)), predictor.matrix = NULL,
            m = 1, k = nrow(data), proper = FALSE, minnumlevels = 5,
            maxfaclevels = 60, rules = NULL, rvalues = NULL,
            cont.na = NULL, semicont = NULL, smoothing = NULL,
            event = NULL, denom = NULL, drop.not.used = FALSE, drop.pred.only = FALSE,
            default.method = c("normrank", "logreg", "polyreg", "polr"), models = FALSE,
            print.flag = TRUE, seed = "sample", ...)
```

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

**Arguments**

data	a data frame or a matrix (n x p) containing the original data. Observations are in rows and variables are in columns.
method	a single string or a vector of strings of length <code>ncol(data)</code> specifying the synthesising method to be used for each variable in the data. Order of variables is exactly the same as in data. If specified as a single string, the same method is used for all variables in a visit sequence unless a data type or a position in a visit sequence requires a different method. If <code>method</code> is set to "parametric" the default synthesising method specified by the <code>default.method</code> argument are applied. Variables that are transformations of other variables can be synthesised using a passive method that is specified as a string starting with <code>~</code> (see <a href="#">syn.passive</a> ). Variables that need not to be synthesised have the empty method <code>""</code> . By default

all variables are synthesised using "cart" method, which is rpart implementation of a CART model (see [syn.cart](#)). See details for more information on method.

<code>visit.sequence</code>	a character vector of names of variables or an integer vector of their column indices specifying the order of synthesis. The default sequence <code>1:ncol(data)</code> implies that column variables are synthesised from left to right. See details for more information.
<code>predictor.matrix</code>	a square matrix of size <code>ncol(data)</code> specifying the set of column predictors to be used for each target variable in the row. Each entry has value 0 or 1. A value of 1 means that the column variable is used as a predictor for the row variable. Order of variables is exactly the same as in data. By default all variables that are earlier in the visit sequence are used as predictors. For the default visit sequence ( <code>1:ncol(data)</code> ) the default <code>predictor.matrix</code> will have values of 1 in the lower triangle. See details for more information.
<code>m</code>	number of synthetic copies of the original (observed) data to be generated. The default is <code>m = 1</code> .
<code>k</code>	a size of the synthetic data set ( $k \times p$ ), which can be smaller or greater than the size of the original data set ( $n \times p$ ). The default is <code>nrow(data)</code> which means that the number of individuals in the synthesised data is the same as in the original (observed) data ( $k = n$ ).
<code>proper</code>	a logical value with default set to FALSE. If TRUE proper synthesis is conducted.
<code>minnumlevels</code>	a minimum number of values a numeric variable should have to be treated as numeric. Numeric variables with fewer levels than <code>minnumlevels</code> are changed into factors. If set to -1 (default) numeric variables are left unchanged regardless of the number of values.
<code>maxfaclevels</code>	a maximum number of factor levels that can be handled. It can be increased but it may cause computational problems, especially for parametric methods.
<code>rules</code>	a named list of rules for restricted values. Restricted values are those that are determined explicitly by values of other variables. The names of the list elements must correspond to the variables names for which the rules need to be specified.
<code>rvalues</code>	a named list of the values corresponding to the rules specified by <code>rules</code> .
<code>cont.na</code>	a named list of codes for missing values for continuous variables if different from the R missing data code NA. The names of the list elements must correspond to the variables names for which the missing data codes need to be specified.
<code>semicont</code>	a named list of values at which semi-continuous variables have spikes. The names of the list elements must correspond to the names of the semi-continuous variables.
<code>smoothing</code>	a named list specifying smoothing method ("density" or "") to be used for selected variables. Smoothing can only be applied to continuous variables synthesised using <code>sample</code> , <code>ctree</code> , <code>cart</code> or <code>normrank</code> method. The names of the list elements must correspond to the names of the variables whose values are to be smoothed. Smoothing is applied to the synthesised values. For "density" smoothing a Gaussian kernel density estimator is applied with bandwidth selected using the Sheather-Jones 'solve-the-equation' method (see <a href="#">bw.SJ</a> ).

<code>event</code>	a named list specifying for survival data the names of corresponding event indicators. The names of the list elements must correspond to the names of the survival variables.
<code>denom</code>	a named list specifying for variables to be modelled using binomial regression the names of corresponding denominator variables. The names of the list elements must correspond to the names of the variables to be modelled using binomial regression.
<code>drop.not.used</code>	a logical value. If TRUE (default) variables not used in synthesis are not saved in the synthesised data and are not included in the corresponding synthesis parameters.
<code>drop.pred.only</code>	a logical value. If TRUE (default) variables not synthesised and used as predictors only are not saved in the synthesised data.
<code>default.method</code>	a vector of four strings containing the default parametric synthesising methods for numerical variables, factors with two levels, unordered factors with more than two levels and ordered factors with more than two levels respectively. They are used when <code>method</code> is set to "parametric" or when there is an inconsistency between variable type and provided method.
<code>models</code>	if TRUE parameters of models fitted to the original data and used to generate the synthetic values are stored.
<code>print.flag</code>	if TRUE (default) synthesising history and information messages will be printed at the console. For silent computation use <code>print.flag = FALSE</code> .
<code>seed</code>	an integer to be used as an argument for the <code>set.seed()</code> . If no integer is provided, the default "sample" will generate one and it will be stored. To prevent generating an integer set seed to NA.
<code>...</code>	additional arguments to be passed to synthesising functions. See section 'Details' below for more information.
<code>strata</code>	a numeric vector with strata identifiers or a string vector with names of stratifying variable(s).
<code>minstratumsize</code>	minimum size of each stratum.
<code>tab.strataobs</code>	a logical value indicating whether a frequency table of the number of observations in strata in the original data set should be printed.
<code>tab.stratasyn</code>	a logical value indicating whether a frequency table of the number of observations in strata in the synthetic data set(s) should be printed.
<code>x</code>	an object of class <code>synds</code> ; a result of a call to <code>syn</code> .

## Details

Only variables that are in `visit.sequence` with corresponding non-empty `method` are synthesised. The only exceptions are event indicators. They are synthesised along with the corresponding time to event variables and should not be included in `visit.sequence`. All other variables (not in `visit.sequence` or in `visit.sequence` with a corresponding blank `method`) can be used as predictors. Including them in `visit.sequence` generates a default `predictor.matrix` reflecting the order of variables in the `visit.sequence` otherwise `predictor.matrix` has to be adjusted accordingly. All predictors of the variables that are not in `visit.sequence` or are in `visit.sequence` but with a blank `method` are removed from `predictor.matrix`.

Variables to be synthesised that are not synthesised yet cannot be used as predictors. Also all variables used in passive synthesis or in restricted values rules (rules) have to be synthesised before the variables they apply to.

Mismatch between data type and synthesising method stops execution and print an error message but numeric variables with number of levels less than `minnumlevels` are changed into factors and methods are changed automatically, if necessary, to methods for categorical variables. Methods for variables not in a visit sequence will be changed into blank.

The built-in elementary synthesising methods include:

**ctree, cart** classification and regression trees (CART), see [syn.cart](#)

**bagging, random forests** methods using ensembles of CART trees, see [syn.bag](#) and [syn.rf](#)

**survctree** classification and regression trees (CART) for duration time data (parametric methods for survival data are not implemented yet), see [syn.survctree](#)

**norm** normal linear regression, see [syn.norm](#)

**normrank** normal linear regression preserving the marginal distribution, see [syn.normrank](#)

**lognorm, sqrtnorm, cubertnorm** normal linear regression after natural logarithmic, square root and cube root transformation of a dependent variable respectively, see [syn.lognorm](#)

**logreg** logistic regression, see [syn.logreg](#)

**polyreg** unordered polytomous regression, see [syn.polyreg](#)

**polr** ordered polytomous regression, see [syn.polr](#)

**pmm** predictive mean matching, see [syn.pmm](#)

**sample** random sample from the observed data, see [syn.sample](#)

**passive** function of other synthesised data, see [syn.passive](#)

**nested** bootstrap sample within each category of the original grouping variable, see [syn.nested](#)

**satcat** bootstrap sample within each category of the crosstabulation of all the predictor variables, see [syn.satcat](#)

The functions corresponding to these methods are called `syn.method`, where `method` is a string with the name of a synthesising method. For instance a function corresponding to `ctree` function is called `syn.ctree`. A new synthesising method can be introduced by writing a function named `syn.newmethod` and then specifying `method` parameter of `syn` function as `"newmethod"`.

In order to use "nested" sampling, `method` parameter of `syn` function has to be specified as `"nested.varname"`, where `"varname"` is the name of the grouped (less detailed) variable, the only one used in nested synthesis. A variable synthesised using "nested" method is excluded from synthesising other variables except when used for "nested" method.

Additional parameters can be passed to synthesising methods as part of the `dots` argument. They have to be named using period-separated method and parameter name (`method.parameter`). For instance, in order to set a `minbucket` (minimum number of observations in any terminal node of a CART model) for a `ctree` synthesising method, `ctree.minbucket` has to be specified. The parameters are method-specific and will be used for all variables to be synthesised using that method. See help for `syn.method` for further details about the allowed parameters for a specific method.



**Value**

An object of class `synds`, which stands for 'synthesised data set'. It is a list with the following components:

<code>call</code>	an original call to <code>syn</code> .
<code>m</code>	number of synthetic versions of the original (observed) data.
<code>syn</code>	a data frame (for $m = 1$ ) or a list of $m$ data frames (for $m > 1$ ) with synthetic data set(s).
<code>method</code>	a vector of synthesising methods applied to each variable in the saved synthesised data.
<code>visit.sequence</code>	a vector of column indices of the visiting sequence. The indices refer to the columns in the saved synthesised data.
<code>predictor.matrix</code>	a matrix specifying the set of predictors used for each variable in the saved synthesised data.
<code>smoothing</code>	a vector specifying smoothing methods applied to each variable in the saved synthesised data.
<code>event</code>	a vector of integers specifying for survival data the column indices for corresponding event indicators. The indices refer to the columns in the saved synthesised data.
<code>denom</code>	a vector of integers specifying for variables modelled using binomial regression the column indices for corresponding denominator variables. The indices refer to the columns in the saved synthesised data.
<code>proper</code>	a logical value indicating whether proper synthesis was conducted.
<code>n</code>	a number of cases in the original data.
<code>k</code>	a number of cases in the synthesised data.
<code>rules</code>	a list of rules for restricted values applied to the synthetic data.
<code>rvalues</code>	a list of the values corresponding to the rules specified by <code>rules</code> .
<code>cont.na</code>	a list of codes for missing values for continuous variables.
<code>semicont</code>	a list of values for semi-continuous variables at which they have spikes.
<code>drop.not.used</code>	a logical value indicating whether variables not used in synthesis are saved in the synthesised data and corresponding synthesis parameters.
<code>drop.pred.only</code>	a logical value indicating whether variables not synthesised and used as predictors only are saved in the synthesised data.
<code>seed</code>	an integer used as a <code>set.seed()</code> argument.
<code>var.lab</code>	a vector of variable labels for data imported from SPSS using <code>read.obs()</code> .
<code>val.lab</code>	a list value labels for factors for data imported from SPSS using <code>read.obs()</code> .
<code>obs.vars</code>	a vector of all variable names in the observed data set.
<code>models</code>	estimates of models fitted to the original data and used to generate the synthetic values.

**Note**

See package vignette for additional information.

**References**

Nowok, B., Raab, G.M and Dibben, C. (2016). synthpop: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).

**See Also**

[compare.synnds](#), [summary.synnds](#)

**Examples**

```
### selection of variables
vars <- c("sex", "age", "marital", "income", "ls", "smoke")
ods <- SD2011[1:1000, vars]

### default synthesis
s1 <- syn(ods)
s1

### synthesis with default parametric methods
s2 <- syn(ods, method = "parametric", seed = 1)
s2$method

### multiple synthesis of selected variables with customised methods
s3 <- syn(ods, visit.sequence = c(2, 1, 4, 5), m = 2,
         method = c("logreg", "sample", "", "normrank", "ctree", ""),
         ctree.minbucket = 10)
summary(s3)
summary(s3, msel = 1:2)

### adjustment to the default predictor matrix
s4.ini <- syn(data = ods, visit.sequence = c(1, 2, 5, 3),
            m = 0, drop.not.used = FALSE)
pM.cor <- s4.ini$predictor.matrix
pM.cor["marital", "ls"] <- 0
s4 <- syn(data = ods, visit.sequence = c(1, 2, 5, 3),
         predictor.matrix = pM.cor)

### handling missing values in continuous variables
s5 <- syn(ods, cont.na = list(income = c(NA, -8)))

### rules for restricted values - marital status of males under 18 should be 'single'
s6 <- syn(ods, rules = list(marital = "age < 18 & sex == 'MALE'"),
         rvalues = list(marital = 'SINGLE'), method = "parametric", seed = 1)
with(s6$syn, table(marital[age < 18 & sex == 'MALE']))
### results for default parametric synthesis without the rule
with(s2$syn, table(marital[age < 18 & sex == 'MALE']))

### stratified synthesis
```

```
s7 <- syn.strata(ods, strata = "sex")
```

---

syn.bag

*Synthesis with bagging*

---

## Description

Generates univariate synthetic data using bagging. It uses [randomForest](#) function from the **randomForest** package with number of sampled predictors equal to number of all predictors.

## Usage

```
syn.bag(y, x, xp, smoothing, proper = FALSE, ntree = 10, ...)
```

## Arguments

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
smoothing	smoothing method for continuous variables.
proper	...
ntree	number of trees to grow.
...	additional parameters passed to <a href="#">randomForest</a> .

## Details

...

## Value

A vector of length k with synthetic values of y.

## References

...

## See Also

[syn](#), [syn.rf](#), [syn.cart](#), [randomForest](#)

---

syn.ctree, syn.cart     *Synthesis with classification and regression trees (CART)*

---

### Description

Generates univariate synthetic data using classification and regression trees (without or with bootstrap).

### Usage

```
syn.ctree(y, x, xp, smoothing, proper = FALSE, minbucket = 5, ...)
syn.cart(y, x, xp, smoothing, proper = FALSE, minbucket = 5, cp = 1e-08, ...)
```

### Arguments

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
smoothing	smoothing method for continuous variables.
proper	for proper synthesis (proper = TRUE) a CART model is fitted to a bootstrapped sample of the original data.
minbucket	the minimum number of observations in any terminal node. See <a href="#">rpart.control</a> and <a href="#">ctree.control</a> for details.
cp	complexity parameter. Any split that does not decrease the overall lack of fit by a factor of cp is not attempted. See <a href="#">rpart.control</a> for details.
...	additional parameters passed to <a href="#">ctree.control</a> for syn.ctree and <a href="#">rpart.control</a> for syn.cart.

### Details

The procedure for synthesis by a CART model is as follows:

1. Fit a classification or regression tree by binary recursive partitioning.
2. For each xp find the terminal node.
3. Randomly draw a donor from the members of the node and take the observed value of y from that draw as the synthetic value.

syn.ctree uses [ctree](#) function from the **party** package and syn.cart uses [rpart](#) function from the **rpart** package. They differ, among others, in a selection of a splitting variable and a stopping rule for the splitting process.

A Gaussian kernel smoothing can be applied to continuous variables by setting smoothing parameter to "density". It is recommended as a tool to decrease the disclosure risk. Increasing minbucket is another means of data protection.

CART models were suggested for generation of synthetic data by Reiter (2005) and then evaluated by Drechsler and Reiter (2011).

**Value**

A vector of length  $k$  with synthetic values of  $y$ .

**References**

Reiter, J.P. (2005). Using CART to generate partially synthetic, public use microdata. *Journal of Official Statistics*, **21**(3), 441–462.

Drechsler, J. and Reiter, J.P. (2011). An empirical evaluation of easily implemented, nonparametric methods for generating synthetic datasets. *Computational Statistics and Data Analysis*, **55**(12), 3232–3243.

**See Also**

[syn](#), [syn.survctree](#), [rpart](#), [ctree](#)

---

syn.lognorm, syn.sqrtnorm, syn.cubertnorm

*Synthesis by linear regression after transformation of a dependent variable*

---

**Description**

Generates univariate synthetic data using linear regression of an outcome variable transformed by natural logarithm (lognorm), square root (sqrtnorm) or cube root (cubertnorm).

**Usage**

```
syn.lognorm(y, x, xp, proper = FALSE, ...)
syn.sqrtnorm(y, x, xp, proper = FALSE, ...)
syn.cubertnorm(y, x, xp, proper = FALSE, ...)
```

**Arguments**

<code>y</code>	an original data vector of length $n$ .
<code>x</code>	a matrix ( $n \times p$ ) of original covariates.
<code>xp</code>	a matrix ( $k \times p$ ) of synthesised covariates.
<code>proper</code>	a logical value specifying whether proper synthesis should be conducted. See details.
<code>...</code>	additional parameters.

**Details**

Generates synthetic values using the spread around the fitted linear regression line of transformed  $y$  given  $x$ . For proper synthesis first the regression coefficients are drawn from normal distribution with mean and variance from the fitted model. The synthetic values are transformed back to the original scale.

**Value**

A vector of length k with synthetic values of y.

**See Also**

[syn](#), [syn.norm](#), [syn.normrank](#)

---

syn.logreg

*Synthesis by logistic regression*

---

**Description**

Generates univariate synthetic data for binary or binomial response variable using logistic regression model.

**Usage**

```
syn.logreg(y, x, xp, denom = NULL, denomp = NULL, proper = FALSE, ...)
```

**Arguments**

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
denom	an original denominator vector of length n for a binomial regression model.
denomp	a synthesised denominator vector of length k for a binomial regression model.
proper	a logical value specifying whether proper synthesis should be conducted. See details.
...	additional parameters.

**Details**

Synthesis for binary response variables by the non-Bayesian or approximate Bayesian logistic regression model. The non-Bayesian method consists of the following steps:

1. Fit a logistic regression to the original data.
2. Calculate predicted inverse logits for synthesised covariates.
3. Compare the inverse logits to a random (0,1) deviate and get synthetic values.

The Bayesian version (for proper synthesis) includes additional step before computing inverse logits:

- Draw coefficients from normal distribution with mean and variance estimated in step 1.

The method relies on the standard `glm.fit` function. Warnings from `glm.fit` are suppressed. Perfect prediction is handled by the data augmentation method.

**Value**

A vector of length  $k$  with synthetic values (0 or 1) of  $y$ .

**See Also**

[syn](#), [glm](#), [glm.fit](#)

---

syn.nested

*Synthesis for a variable nested within another variable.*

---

**Description**

Synthesizes one variable ( $y$ ) from another one ( $x$ ) when  $y$  is nested in the categories of  $x$ . A bootstrap sample is created from the original values of  $y$  within each category of  $x$  (the synthesised values of the grouping variable).

**Usage**

```
syn.nested(y, x, xp, ...)
```

**Arguments**

$y$	an original data vector of length $n$ for the nested variable.
$x$	an original data vector of length $n$ for the variable within which $y$ is nested.
$xp$	a vector of length $k$ with synthetic values of $x$ .
$\dots$	additional parameters.

**Details**

An example would be when  $x$  is a detailed classification of occupations and  $y$  is more detailed sub-classification. It is intended that both  $x$  and  $y$  are categorical (factor) variables.

A warning will be issued if the original  $y$  is not nested within  $x$ .

**Value**

A vector of length  $k$  with synthetic values of  $y$ .

---

syn.norm	<i>Synthesis by linear regression</i>
----------	---------------------------------------

---

**Description**

Generates univariate synthetic data using linear regression analysis.

**Usage**

```
syn.norm(y, x, xp, proper = FALSE, ...)
```

**Arguments**

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
proper	a logical value specifying whether proper synthesis should be conducted. See details.
...	additional parameters.

**Details**

Generates synthetic values using the spread around the fitted linear regression line of y given x. For proper synthesis first the regression coefficients are drawn from normal distribution with mean and variance from the fitted model.

**Value**

A vector of length k with synthetic values of y.

**See Also**

[syn](#), [syn.normrank](#), [syn.lognorm](#)

---

syn.normrank	<i>Synthesis by normal linear regression preserving the marginal distribution</i>
--------------	-----------------------------------------------------------------------------------

---

**Description**

Generates univariate synthetic data using linear regression analysis and preserves the marginal distribution. Regression is carried out on Normal deviates of ranks in the original variable. Synthetic values are assigned from the original values based on the synthesised ranks that are transformed from their synthesised Normal deviates.



**Usage**

```
syn.normrank(y, x, xp, smoothing, proper = FALSE, ...)
```

**Arguments**

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
smoothing	smoothing method. See details.
proper	a logical value specifying whether proper synthesis should be conducted. See details.
...	additional parameters.

**Details**

First generates synthetic values of Normal deviates of ranks of the values in y using the spread around the fitted linear regression line of Normal deviates of ranks given x. Then synthetic Normal deviates of ranks are transformed back to get synthetic ranks which are used to assign values from y. For proper synthesis first the regression coefficients are drawn from normal distribution with mean and variance from the fitted model. A Gaussian kernel smoothing can be applied by setting smoothing parameter to "density". It is recommended as a tool to decrease the disclosure risk.

**Value**

A vector of length k with synthetic values of y.

**See Also**

[syn](#), [syn.norm](#), [syn.lognorm](#)

---

syn.passive

*Passive synthesis*

---

**Description**

Derives a new variable according to a specified function of synthesised data.

**Usage**

```
syn.passive(data, func)
```

**Arguments**

data	a data frame with synthesised data.
func	a formula specifying transformations on data. It is specified as a string starting with ~.

**Details**

Any function of the synthesised data can be specified. Note that several operators such as +, -, \* and ^ have different meanings in formula syntax. Use the identity function I() if they should be interpreted as arithmetic operators, e.g. "~I(age^2)".

**Value**

A vector including the result of applying the formula.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Van Buuren, S. and Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[syn](#)

---

syn.pmm

*Synthesis by predictive mean matching*

---

**Description**

Generates univariate synthetic data using predictive mean matching.

**Usage**

```
syn.pmm(y, x, xp, proper = FALSE, ...)
```

**Arguments**

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
proper	a logical value specifying whether proper synthesis should be conducted. See details.
...	additional parameters.

**Details**

Synthesis of  $y$  by predictive mean matching. The procedure is as follows:

1. Fit a linear regression to the original data.
2. Compute predicted values  $y.\hat{}$  and  $y_{\text{syn}}.\hat{}$  for the original  $x$  and synthesised  $x_p$  covariates respectively.
3. For each predicted value  $y_{\text{syn}}.\hat{}$  find donor observations with the closest predicted values  $y.\hat{}$  (ties are broken by random selection), randomly sample one of them and take its observed value  $y$  as the synthetic value.

The Bayesian version (for proper synthesis) includes additional step before computing predicted values:

- Draw coefficients from normal distribution with mean and variance estimated in step 1 and use them to calculate predicted values for the synthesised covariates.

**Value**

A numeric vector of length  $k$  with synthetic values of  $y$ .

**See Also**

[syn](#)

---

syn.polr

*Synthesis by ordered polytomous regression*

---

**Description**

Generates a synthetic categorical variable using ordered polytomous regression (without or with bootstrap).

**Usage**

```
syn.polr(y, x, xp, proper = FALSE, maxit = 1000, trace = FALSE,
        MaxNWts = 10000, ...)
```

**Arguments**

<code>y</code>	an original data vector of length $n$ .
<code>x</code>	a matrix ( $n \times p$ ) of original covariates.
<code>xp</code>	a matrix ( $k \times p$ ) of synthesised covariates.
<code>proper</code>	for proper synthesis ( <code>proper = TRUE</code> ) a model is fitted to a bootstrapped sample of the original data.
<code>maxit</code>	the maximum number of iterations for <a href="#">nnet</a> .
<code>trace</code>	switch for tracing optimization for <a href="#">nnet</a> .
<code>MaxNWts</code>	the maximum allowable number of weights for <a href="#">nnet</a> .
<code>...</code>	additional parameters passed to <a href="#">optim</a> or <a href="#">nnet</a> .

**Details**

Generates synthetic ordered categorical variables by the proportional odds logistic regression (polr) model. The function repeatedly applies logistic regression on the successive splits. The model is also known as the cumulative link model.

The algorithm of `syn.polr` uses the function `polr` from the **MASS** package.

In order to avoid bias due to perfect prediction, the data are augmented by the method of White, Daniel and Royston (2010).

In case the call to `polr` fails, usually because the data are very sparse, `multinom` function is used instead.

**Value**

A vector of length `k` with synthetic values of `y`.

**References**

White, I.R., Daniel, R. and Royston, P. (2010). Avoiding bias due to perfect prediction in multiple imputation of incomplete categorical variables. *Computational Statistics and Data Analysis*, **54**, 2267–2275.

**See Also**

[syn](#), [syn.polyreg](#) [multinom](#), [polr](#)

---

syn.polyreg

*Synthesis by unordered polytomous regression*

---

**Description**

Generates a synthetic categorical variable using unordered polytomous regression (without or with bootstrap).

**Usage**

```
syn.polyreg(y, x, xp, proper = FALSE, maxit = 1000, trace = FALSE,
            MaxNWts = 10000, ...)
```

**Arguments**

<code>y</code>	an original data vector of length <code>n</code> .
<code>x</code>	a matrix ( <code>n x p</code> ) of original covariates.
<code>xp</code>	a matrix ( <code>k x p</code> ) of synthesised covariates.
<code>proper</code>	for proper synthesis ( <code>proper = TRUE</code> ) a multinomial model is fitted to a bootstrapped sample of the original data.
<code>maxit</code>	the maximum number of iterations for <a href="#">nnet</a> .

trace	switch for tracing optimization for <a href="#">nnet</a> .
MaxNWts	the maximum allowable number of weights for <a href="#">nnet</a> .
...	additional parameters passed to <a href="#">nnet</a> .

### Details

Generates synthetic categorical variables by the polytomous regression model. The method consists of the following steps:

1. Fit categorical response as a multinomial model.
2. Compute predicted categories.
3. Add appropriate noise to predictions.

The algorithm of `syn.polyreg` uses the function `multinom` from the `nnet` package. Any numerical variables are scaled to cover the range (0,1) before fitting. Warnings are printed if the algorithm fails to converge in `maxit` iterations and also if the synthesised data has only one category. The latter may occur if the variable being synthesised is sparse so that the algorithm fails to iterate.

In order to avoid bias due to perfect prediction, the data are augmented by the method of White, Daniel and Royston (2010).

### Value

A vector of length `k` with synthetic values of `y`.

### References

White, I.R., Daniel, R. and Royston, P. (2010). Avoiding bias due to perfect prediction in multiple imputation of incomplete categorical variables. *Computational Statistics and Data Analysis*, **54**, 2267–2275.

### See Also

[syn](#), [syn.polr](#), [multinom](#), [polr](#)

---

syn.rf

*Synthesis with random forest*

---

### Description

Generates univariate synthetic data using Breiman's random forest algorithm classification and regression. It uses `randomForest` function from the `randomForest` package.

### Usage

```
syn.rf(y, x, xp, smoothing, proper = FALSE, ntree = 10, ...)
```

**Arguments**

y	an original data vector of length n.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
smoothing	smoothing method for continuous variables.
proper	...
ntree	number of trees to grow.
...	additional parameters passed to <a href="#">randomForest</a> .

**Details**

...

**Value**

A vector of length k with synthetic values of y.

**References**

...

**See Also**

[syn](#), [syn.bag](#), [syn.cart](#), [randomForest](#)

---

 syn.sample

*Synthesis by simple random sampling*


---

**Description**

Generates a random sample from the observed data.

**Usage**

```
syn.sample(y, xp, smoothing, cont.na, proper = FALSE, ...)
```

**Arguments**

y	an original data vector of length n.
xp	a target length k of a synthetic data vector.
smoothing	smoothing method for a continuous variable.
cont.na	a vector of codes for missing values for continuous variables that should be excluded from smoothing.
proper	if proper = TRUE values are sampled from a bootstrapped sample of the original data.
...	additional parameters passed to <code>sample</code> .

**Details**

A simple random sample with replacement is taken from the observed values in  $y$  and used as synthetic values. A Gaussian kernel smoothing can be applied to continuous variables by setting smoothing parameter to "density". It is recommended as a tool to decrease the disclosure risk.

**Value**

A vector of length  $k$  with synthetic values.

**See Also**

[syn](#)

---

syn.satcat	<i>Synthesis from a saturated model based on all combinations of the predictor variables.</i>
------------	-----------------------------------------------------------------------------------------------

---

**Description**

Synthesises one variable ( $y$ ) from all possible combinations of its precitors ( $x$ ). A bootstrap sample is created from the original values of  $y$  within each unique combinations of of  $x_p$  (the synthesised values of the grouping variable).

**Usage**

```
syn.satcat(y, x, xp, proper = FALSE, ...)
```

**Arguments**

$y$	an original data vector of length $n$ for the satcat variable.
$x$	a matrix ( $n \times p$ ) with the original predictor variables for $y$ .
$x_p$	a matrix ( $k \times p$ ) with synthetic values of $x$ .
proper	if proper = TRUE values are a bootstrapped sample from the synthesised data.
...	additional parameters.

**Details**

It is intended that the variables in  $x$  are categorical (factor) variables. If  $y$  is also a categorical variable `syn.satcat` will give the same results as fitting a saturated polychotomous regression model but will usually be much faster. `syn.satcat` will fail with an error message if previous syntheses have generated a combination of variables in  $x_p$  that was not present in  $x$ .

**Value**

A vector of length  $k$  with synthetic values of  $y$ .

**Examples**

```
ods <- SD2011[, c("region", "sex", "agegr", "placesize")]

s1 <- syn(ods, method = c("sample", "cart", "satcat", "cart"))

### mostly fails because too many small categories
s2 <- syn(ods, method = c("sample", "cart", "cart", "satcat"))
```

---

syn.survctree	<i>Synthesis of survival time by classification and regression trees (CART)</i>
---------------	---------------------------------------------------------------------------------

---

**Description**

Generates synthetic event indicator and time to event data using classification and regression trees (without or with bootstrap).

**Usage**

```
syn.survctree(y, yevent, x, xp, proper = FALSE, minbucket = 5, ...)
```

**Arguments**

y	a vector of length n with original time data.
yevent	a vector of length n with original event indicator data.
x	a matrix (n x p) of original covariates.
xp	a matrix (k x p) of synthesised covariates.
proper	for proper synthesis (proper = TRUE) a CART model is fitted to a bootstrapped sample of the original data.
minbucket	the minimum number of observations in any terminal node. See <a href="#">ctree_control</a> for details.
...	additional parameters passed to <a href="#">ctree</a> .

**Details**

The procedure for synthesis by a CART model is as follows:

1. Fit a tree-structured survival model by binary recursive partitioning (the terminal nodes include Kaplan-Meier estimates of the survival time).
2. For each xp find the terminal node.
3. Randomly draw a donor from the members of the node and take the observed value of yevent and y from that draw as the synthetic values.



**Value**

A list with the following components:

syn.time            a vector of length k with synthetic time values.  
 syn.event           a vector of length k with synthetic event indicator values.

**See Also**

[syn](#), [syn.ctree](#)

---

 utility.gen

*Distributional comparison of synthesised and observed data*


---

**Description**

Distributional comparison of synthesised data set with the original (observed) data set using propensity scores.

**Usage**

```
utility.gen(object, data, method = "logit", maxorder = 1,
            tree.method = "rpart", resamp.method = NULL,
            nperms = 50, cp = 1e-3, minbucket = 5, mincriterion = 0,
            vars = NULL, aggregate = FALSE, maxit = 200, ngroups = NULL,
            print.every = 10, digits = 2, print.zscores = FALSE, zthresh = 1.6,
            print.ind.results = TRUE, print.variable.importance = FALSE, ...)
```

```
## S3 method for class 'utility.gen'
print(x, digits = x$digits,
      print.zscores = x$print.zscores, zthresh = x$zthresh,
      print.ind.results = x$print.ind.results,
      print.variable.importance = x$print.variable.importance, ...)
```

**Arguments**

object	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn()</code> and it includes <code>object\$m</code> synthesised data set(s) as <code>object\$syn</code> . This a single data set when <code>object\$m = 1</code> or a list of length <code>object\$m</code> when <code>object\$m &gt; 1</code> .
data	the original (observed) data set.
method	a single string specifying the method for modeling the propensity scores. Method can be selected from "logit" and "cart".
maxorder	maximum order of interactions to be considered in "logit" method. For model without interactions 0 should be provided.
tree.method	implementation of "cart" method, can be "rpart" or "ctree".

resamp.method	method used for resampling estimate of the null pMSE, can be "perm" or "pairs". For pMSEs calculated with method "cart" it defaults to "perm" if all the variables from "object" in "vars" have been synthesised or to "pairs" if some have not (i.e. have method = "").
nperms	number of permutations for the permutation test to obtain the null distribution of the utility measure when resamp.method = "perm".
cp	complexity parameter for classification with tree.method "rpart". Small values grow bigger trees.
minbucket	minimum number of observations allowed in a leaf for classification when method = "cart".
mincriterion	criterion between 0 and 1 to use to control tree.method = "ctree" when the tree will not be allowed to split further. A value of 0.95 would be equivalent to a 5% significance test. Here we set it to 0 to effectively disable this test and grow large trees.
vars	variables to be included in the utility comparison. If none are specified all the variables in the synthesised data will be included.
aggregate	logical flag as to whether the data should be aggregated by collapsing identical rows before computation. This can lead to much faster computation when all the variables are categorical. Only works for method = "logit".
maxit	maximum iterations to use when method = "logit". If the model does not converge in this number a warning will suggest increasing it.
ngroups	target number of groups for categorisation of each numeric variable: final number may differ if there are many repeated values. If NULL (default) variables are not categorised into groups.
print.every	controls the printing of progress of resampling when resamp.method is not NULL. When print.every = 0 no progress is reported, otherwise the resample number is printed every print.every.
...	additional parameters passed to <a href="#">glm</a> , <a href="#">rpart</a> , or <a href="#">ctree</a> .
x	an object of class utility.gen.
digits	number of digits to print in the default output, excluding pMSE values.
print.zscores	logical value as to whether z-scores for coefficients of the logit model should be printed.
zthresh	threshold value to use to suppress the printing of z-scores under +/- this value for method = "logit". If set to NA all z-scores are printed.
print.ind.results	logical value as to whether utility score results from individual syntheses should be printed.
print.variable.importance	logical value as to whether the variable importance measure should be printed when tree.method = "rpart".

## Details

This function follows the method for evaluating the utility of masked data as given in Snoke et al. (forthcoming) and originally proposed by Woo et al. (2009). The original and synthetic data are

combined into one dataset and propensity scores, as detailed in Rosenbaum and Rubin (1983), are calculated to estimate the probability of membership in the synthetic data set. The utility measure is based on the mean squared difference between these probabilities and the probability expected if the data did not distinguish the synthetic data from the original. The expected probability is just the proportion of synthetic data in the combined data set, 0.5 when the original and synthetic data have the same number of records.

Propensity scores can be modeled by logistic regression method = "logit" or by two different implementations of classification and regression trees as method "cart". For logistic regression the predictors are all variables in the data and their interactions up to order maxorder. The default of 1 gives all main effects and first order interactions. For logistic regression the null distribution of the propensity score is derived and is used to calculate ratios and standardised values.

For method = "cart" the expectation and variance of the null distribution is calculated from a permutation test.

If missing values exist, indicator variables are added and included in the model as recommended by Rosenbaum and Rubin (1984). For categorical variables, NA is treated as a new category.

## Value

An object of class utility.gen which is a list including the utility measures their expected null values for each synthetic set with the following components:

call	the call that produced the result.
m	number of synthetic data sets in object.
method	method used to fit propensity score.
tree.method	cart function used to fit propensity score when method = "cart".
pMSE	Propensity score mean square error from the utility model or a vector of these values if object\$m > 1.
utilVal	utility value(s). Calculated from the pMSE as $pMSE * (n1+n2)^3 / n1^2 / n2$ or a vector of these values if object\$m > 1. For method = "logit" the null distribution of this quantity will be chi-squared with degrees of freedom equal to the number of parameters involving synthesised variables in the propensity score minus 1. For method = "cart" its distribution will be obtained by resampling.
utilExp	expected value(s) of the utility score if the synthesis method is correct.
utilR	ratio(s) of utilVal(s) to utilExp.
utilStd	utility value standardised by expressing it as z-scores, difference(s) from the expected value divided by the expected standard deviation.
fit	the fitted model for the propensity score or a list of fitted models of length m if m > 0.

## References

- Woo, M-J., Reiter, J.P., Oganian, A. and Karr, A.F. (2009). Global measures of data utility for microdata masked for disclosure limitation. *Journal of Privacy and Confidentiality*, 1(1), 111-124.
- Rosenbaum, P.R. and Rubin, D.B. (1984). Reducing bias in observational studies using subclassification on the propensity score. *Journal of the American Statistical Association*, 79(387), 516-524.
- Snoke, J., Raab, G.M., Nowok, B., Dibben, C. and Slavkovic, A. (forthcoming). General and specific utility measures for synthetic data.

**See Also**[utility.tab](#)**Examples**

```
## Not run:
ods <- SD2011[1:1000, c("age", "bmi", "depress", "alcabuse", "englang")]
s1 <- syn(ods, m = 5)
utility.gen(s1, ods)
u1 <- utility.gen(s1, ods)
print(u1, print.zscores = TRUE, usethresh = TRUE)
u2 <- utility.gen(s1, ods, groups = TRUE)
print(u2, print.zscores = TRUE)
u3 <- utility.gen(s1, ods, method = "cart", nperms = 20)
print(u3, print.variable.importance = TRUE)
## End(Not run)
```

utility.tab

*Tabular utility***Description**

Produce tables from observed and synthesized data and calculates utility measures to compare them with their expectation if the synthesising model is correct.

**Usage**

```
utility.tab(object, data, vars = NULL, ngroups = 5,
            print.tables = TRUE, print.zdiff = FALSE,
            digits = 2, ...)

## S3 method for class 'utility.tab'
print(x, print.tables = x$print.tables,
      print.zdiff = x$print.zdiff, digits = x$digits, ...)
```

**Arguments**

object	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <code>syn()</code> or <code>syn.strata()</code> and it includes <code>object\$m</code> number of synthesised data set(s), as well as <code>object\$syn</code> the synthesised data set, if <code>m = 1</code> , or a list of <code>m</code> such data sets.
data	the original (observed) data set.
vars	a single string or a vector of strings with the names of variables to be used to form the table.

ngroups	if numerical (non-factor) variables are included they will be classified into this number of groups to form tables. Classification is performed using <code>classIntervals()</code> function for $n = \text{ngroups}$ . By default, to avoid problems for variables with a small number of unique values, <code>style = "fisher"</code> . Arguments of <code>classIntervals()</code> may be, however, specified in the call to <code>utility.tab()</code> .
digits	an integer indicating the number of decimal places for printing statistics, <code>tab.zdiff</code> and mean results for $m > 1$ .
print.tables	a logical value that determines if tables of observed and synthesised are to be printed.
print.zdiff	a logical value that determines if tables of Z scores for differences between observed and expected are to be printed.
...	additional parameters; can be passed to <code>classIntervals()</code> function.
x	an object of class <code>utility.tab</code> .

### Details

Forms tables of observed and synthesised values for the variables specified in `vars`. Two utility measures are calculated from the cells of the tables, a measure of fit proposed by Voas and Williams  $\text{sum}((\text{observed} - \text{synthesised})^2 / [(\text{observed} + \text{synthesised})/2])$  and one proposed by Freeman and Tukey  $4 * \text{sum}((\text{observed}^{(0.5)} - \text{synthesised}^{(0.5)})^2)$ . In both cases those cells where observed and synthesised are both zero do not contribute to the sum. If the synthesising model is correct both of these measures should have chi-square distributions for large samples.

### Value

An object of class `utility.tab` which is a list with the following components:

<code>m</code>	number of synthetic data sets in object, i.e. <code>object\$m</code> .
<code>tab.obs</code>	a table from the observed data.
<code>UtabFT</code>	a vector with <code>object\$m</code> values for the Freeman Tukey utility measure.
<code>UtabVW</code>	a vector with <code>object\$m</code> values for the Voas Williamson utility measure.
<code>df</code>	a vector of degrees of freedom for the chi-square tests which equal to one minus the number of cells in the table with any observed or synthesised counts.
<code>ratioFT</code>	a vector with ratios of <code>UtabFT</code> to <code>df</code> .
<code>ratioVW</code>	a vector with ratios of <code>UtabVW</code> to <code>df</code> .
<code>pvalFT</code>	a vector with <code>object\$m</code> p-values for the chi-square tests for the Freeman Tukey utility measure.
<code>pvalVW</code>	a vector with <code>object\$m</code> p-values for the chi-square tests for the Voas Williamson utility measure.
<code>nempty</code>	a vector of length <code>object\$m</code> with number of cells not contributing to the statistics.
<code>tab.obs</code>	a table from the observed data.
<code>tab.syn</code>	a table or a list of <code>m</code> tables from the synthetic data.

tab.zdiff      a table or a list of  $m$  tables of  $Z$  statistics for differences between observed and synthesised cells of the tables. Large absolute values indicate a large contribution to lack-of-fit.

n                number of observation in the original dataset.

## References

Nowok, B., Raab, G.M and Dibben, C. (2016). synthpop: Bespoke creation of synthetic data in R. *Journal of Statistical Software*, **74**(11), 1-26. doi: [10.18637/jss.v074.i11](https://doi.org/10.18637/jss.v074.i11).

Read, T.R.C. and Cressie, N.A.C. (1988) *Goodness-of-Fit Statistics for Discrete Multivariate Data*, Springer-Verlag, New York.

Voas, D. and Williamson, P. (2001) Evaluating goodness-of-fit measures for synthetic microdata. *Geographical and Environmental Modelling*, **5**(2), 177-200.

## See Also

[utility.gen](#)

## Examples

```
ods <- SD2011[1:1000, c("sex", "age", "edu", "marital")]

s1 <- syn(ods, m = 10)
utility.tab(s1, ods, vars = c("marital", "sex"))

s2 <- syn(ods, m = 1)
utility.tab(s2, ods, vars = c("marital", "age"), ngroups = 3, print.tables = TRUE)
u2 <- utility.tab(s2, ods, vars = c("marital", "age"), style = "pretty")
print(u2, print.tables = TRUE, print.zdiff = TRUE)
```

---

write.syn

*Exporting synthetic data sets to external files*

---

## Description

Exports synthetic data set(s) from synthesised data set (synds) object to external files of selected format. Currently supported file formats include: SPSS, Stata, SAS, csv, tab, rda, RData and txt. For SPSS, Stata and SAS it uses functions from the foreign package with some adjustments where necessary. Information about the synthesis is written into a separate text file.

NOTE: Currently numeric codes and labels can be preserved correctly only for SPSS files imported into R using [read.obs](#) function.

## Usage

```
write.syn(object, filename,
filetype = c("SPSS", "Stata", "SAS", "csv", "tab", "rda", "RData", "txt"),
convert.factors = "numeric", data.labels = NULL, save.complete = TRUE,
extended.info = TRUE, ...)
```

**Arguments**

<code>object</code>	an object of class <code>synds</code> , which stands for 'synthesised data set'. It is typically created by function <a href="#">syn</a> and it includes <code>object\$m</code> synthesised data set(s).
<code>filename</code>	the name of the file (excluding extension) which the synthetic data are to be written into. For multiple synthetic data sets it will be used as a prefix followed respectively by <code>_1</code> , <code>_...</code> , <code>_m</code> .
<code>filetype</code>	a desired format of the output files.
<code>convert.factors</code>	a single string indicating how to handle factors in Stata output files. The default value is set to "numeric" in order to preserve the numeric codes from the original data. See <a href="#">write.dta</a> for other possible values.
<code>data.labels</code>	a list with variable labels and value labels.
<code>save.complete</code>	a logical value indicating whether a complete 'synthesised data set' ( <code>synds</code> ) object should be saved into a file ( <code>synobject_filename.RData</code> ).
<code>extended.info</code>	a logical value indicating whether extended information should be saved into an information file.
<code>...</code>	additional parameters passed to write functions.

**Value**

File(s) with synthesised data set(s) and a text file with information about synthesis are produced. Optionally a complete synthesised data set object is saved into `synobject_filename.RData` file.

**See Also**

[read.obs](#)

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