Package 'surveysd'

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Type Package

Title Survey Standard Error Estimation for Cumulated Estimates and their Differences in Complex Panel Designs

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Description Calculate point estimates and their standard errors in complex household surveys using bootstrap replicates. Bootstrapping considers survey design with a rotating panel. A comprehensive description of the methodology can be found under <https://statistikat.github.io/surveysd/articles/methodology.html>.

Encoding UTF-8

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Imports Rcpp (>= 0.12.12),data.table,ggplot2,laeken,methods

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BugReports https://github.com/statistikat/surveysd/issues

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```
calc.stError
```

Calcualte point estimates and their standard errors using bootstrap weights.

Description

Calculate point estimates as well as standard errors of variables in surveys. Standard errors are estimated using bootstrap weights (see draw.bootstrap and recalib). In addition the standard error of an estimate can be calculated using the survey data for 3 or more consecutive periods, which results in a reduction of the standard error.

Usage

```
calc.stError(
  dat,
 weights = attr(dat, "weights"),
 b.weights = attr(dat, "b.rep"),
  period = attr(dat, "period"),
  var,
  fun = weightedRatio,
  national = FALSE,
  group = NULL,
  fun.adjust.var = NULL,
  adjust.var = NULL,
  period.diff = NULL,
  period.mean = NULL,
  bias = FALSE,
  size.limit = 20,
  cv.limit = 10,
```

```
p = NULL,
add.arg = NULL
)
```

dat	either data.frame or data.table containing the survey data. Surveys can be a panel survey or rotating panel survey, but does not need to be. For rotating panel survey bootstrap weights can be created using draw.bootstrap and recalib.	
weights	character specifying the name of the column in dat containing the original sam- ple weights. Used to calculate point estimates.	
b.weights	character vector specifying the names of the columns in dat containing boot- strap weights. Used to calculate standard errors.	
period	character specifying the name of the column in dat containing the sample peri- ods.	
var	character vector containing variable names in dat on which fun shall be applied for each sample period.	
fun	function which will be applied on var for each sample period. Predefined func- tions are weightedRatio, weightedSum, but can also take any other function which returns a double or integer and uses weights as its second argument.	
national	boolean, if TRUE point estimates resulting from fun will be divided by the point estimate at the national level.	
group	character vectors or list of character vectors containing variables in dat. For each list entry dat will be split in subgroups according to the containing variables as well as period. The pointestimates are then estimated for each subgroup seperately. If group=NULL the data will split into sample periods by default.	
fun.adjust.var	can be either NULL or a function. This argument can be used to apply a function for each period and bootstrap weight to the data. The resulting estimates will be passed down to fun. See details for more explanations.	
adjust.var	can be either NULL or a character specifying the first argument in fun.adjust.var.	
period.diff	character vectors, defining periods for which the differences in the point esti- mate as well it's standard error is calculated. Each entry must have the form of "period1 - period2". Can be NULL	
period.mean	odd integer, defining the range of periods over which the sample mean of point estimates is additionally calcualted.	
bias	boolean, if TRUE the sample mean over the point estimates of the bootstrap weights is returned.	
size.limit	integer defining a lower bound on the number of observations on dat in each group defined by period and the entries in group. Warnings are returned if the number of observations in a subgroup falls below size.limit. In addition the concerned groups are available in the function output.	
cv.limit	non-negativ value defining a upper bound for the standard error in relation to the point estimate. If this relation exceed cv.limit, for a point estimate, they are flagged and available in the function output.	

р	numeric vector containing values between 0 and 1. Defines which quantiles for the distribution of var are additionally estimated.
add.arg	additional arguments which will be passed to fun. Can be either a named list or vector. The names of the object correspond to the function arguments and the values to column names in dat, see also examples.

calc.stError takes survey data (dat) and returns point estimates as well as their standard Errors defined by fun and var for each sample period in dat. dat must be household data where household members correspond to multiple rows with the same household identifier. The data should at least contain the following columns:

- Column indicating the sample period;
- Column indicating the household ID;
- Column containing the household sample weights;
- Columns which contain the bootstrap weights (see output of recalib);
- Columns listed in var as well as in group

For each variable in var as well as sample period the function fun is applied using the original as well as the bootstrap sample weights.

The point estimate is then selected as the result of fun when using the original sample weights and it's standard error is estimated with the result of fun using the bootstrap sample weights.

fun can be any function which returns a double or integer and uses sample weights as it's second argument. The predifined options are weightedRatio and weightedSum.

For the option weightedRatio a weighted ratio (in \ calculated for var equal to 1, e.g sum(weight[var==1])/sum(weight[! Additionally using the option national=TRUE the weighted ratio (in \ divided by the weighted ratio at the national level for each period.

If group is not NULL but a vector of variables from dat then fun is applied on each subset of dat defined by all combinations of values in group.

For instance if group = "sex" with "sex" having the values "Male" and "Female" in dat the point estimate and standard error is calculated on the subsets of dat with only "Male" or "Female" value for "sex". This is done for each value of period. For variables in group which have NAs in dat the rows containing the missings will be discarded.

When group is a list of character vectors, subsets of dat and the following estimation of the point estimate, including the estimate for the standard error, are calculated for each list entry.

The optional parameters fun.adjust.var and adjust.var can be used if the values in var are dependent on the weights. As is for instance the case for the poverty thershhold calculated from EU-SILC. In such a case an additional function can be supplied using fun.adjust.var as well as its first argument adjust.var, which needs to be part of the data set dat. Then, before applying fun on variable var for all period and groups, the function fun.adjust.var is applied to adjust.var using each of the bootstrap weights seperately (NOTE: weight is used as the second argument of fun.adjust.var). Thus creating i=1,...,length(b.weights) additional variables. For applying fun on var the estimates for the bootstrap replicate will now use each of the corresponding new

calc.stError

additional variables. So instead of

 $fun(var, weights, \ldots), fun(var, b.weights[1], \ldots), fun(var, b.weights[2], \ldots), \ldots$

the function fun will be applied in the way

fun(var, weights, ...), fun(var.1, b.weights[1], ...), fun(var.2, b.weights[2], ...), ...

where var.1, var.2, ... correspond to the estimates resulting from fun.adjust.var and adjust.var. NOTE: This procedure is especially usefull if the var is dependent on weights and fun is applied on subgroups of the data set. Then it is not possible to capture this procedure with fun and var, see examples for a more hands on explanation.

When defining period.diff the difference of point estimates between periods as well their standard errors are calculated.

The entries in period.diff must have the form of "period1 - period2" which means that the results of the point estimates for period2 will be substracted from the results of the point estimates for period1.

Specifying period.mean leads to an improvement in standard error by averaging the results for the point estimates, using the bootstrap weights, over period.mean periods. Setting, for instance, period.mean = 3 the results in averaging these results over each consecutive set of 3 periods.

Estimating the standard error over these averages gives an improved estimate of the standard error for the central period, which was used for averaging.

The averaging of the results is also applied in differences of point estimates. For instance defining period.diff = "2015-2009" and period.mean = 3 the differences in point estimates of 2015 and 2009, 2016 and 2010 as well as 2014 and 2008 are calcualated and finally the average over these 3 differences is calculated. The periods set in period.diff are always used as the middle periods around which the mean over period.mean years is build.

Setting bias to TRUE returns the calculation of a mean over the results from the bootstrap replicates. In the output the corresponding columns is labeled *_mean* at the end.

If fun needs more arguments they can be supplied in add.arg. This can either be a named list or vector.

The parameter size.limit indicates a lower bound of the sample size for subsets in dat created by group. If the sample size of a subset falls below size.limit a warning will be displayed.

In addition all subsets for which this is the case can be selected from the output of calc.stError with \$smallGroups.

With the parameter cv.limit one can set an upper bound on the coefficient of variantion. Estimates which exceed this bound are flagged with TRUE and are available in the function output with cv.limit nust be a positive integer and is treated internally as \ for cv.limit=1 the estimate will be flagged if the coefficient of variantion exceeds 1\

When specifying period.mean, the decrease in standard error for choosing this method is internally calcualted and a rough estimate for an implied increase in sample size is available in the output with stEDecrease. The rough estimate for the increase in sample size uses the fact that for a sample of size n the sample estimate for the standard error of most point estimates converges with a factor $1/\sqrt{n}$ against the true standard error σ .

Value

Returns a list containing:

- Estimates: data.table containing period differences and/or k period averages for estimates of fun applied to var as well as the corresponding standard errors, which are calculated using the bootstrap weights. In addition the sample size, n, and population size for each group is added to the output.
- smallGroups: data.table containing groups for which the number of observation falls below size.limit.
- cvHigh: data.table containing a boolean variable which indicates for each estimate if the estimated standard error exceeds cv.limit.
- stEDecrease: data.table indicating for each estimate the theoretical increase in sample size which is gained when averaging over k periods. Only returned if period.mean is not NULL.

Author(s)

Johannes Gussenbauer, Alexander Kowarik, Statistics Austria

See Also

draw.bootstrap recalib

Examples

Import data and calibrate

```
set.seed(1234)
eusilc <- demo.eusilc(n = 4,prettyNames = TRUE)</pre>
dat_boot <- draw.bootstrap(eusilc, REP = 3, hid = "hid", weights = "pWeight",</pre>
                            strata = "region", period = "year")
dat_boot_calib <- recalib(dat_boot, conP.var = "gender", conH.var = "region")</pre>
# estimate weightedRatio for povertyRisk per period
err.est <- calc.stError(dat_boot_calib, var = "povertyRisk",</pre>
                         fun = weightedRatio)
err.est$Estimates
# calculate weightedRatio for povertyRisk and fraction of one-person
# households per period
dat_boot_calib[, onePerson := .N == 1, by = .(year, hid)]
err.est <- calc.stError(dat_boot_calib, var = c("povertyRisk", "onePerson"),</pre>
                         fun = weightedRatio)
err.est$Estimates
## Not run:
# estimate weightedRatio for povertyRisk per period and gender and
# period x region x gender
group <- list("gender", c("gender", "region"))</pre>
err.est <- calc.stError(dat_boot_calib, var = "povertyRisk",</pre>
                         fun = weightedRatio, group = group)
```

computeLinear

```
err.est$Estimates
# use average over 3 periods for standard error estimation
# and calculate estimate for difference of
# period 2011 and 2012 inclulding standard errors
period.diff <- c("2012-2011")
err.est <- calc.stError(
    dat_boot_calib, var = "povertyRisk", fun = weightedRatio,
    period.diff = period.diff, # <- take difference of periods 2012 and 2011
    period.mean = 3) # <- average over 3 periods
err.est$Estimates
## End(Not run)</pre>
```

for more examples see https://statistikat.github.io/surveysd/articles/error_estimation.html

computeLinear Numerical weighting functions

Description

Customize weight-updating within factor levels in case of numerical calibration. The functions described here serve as inputs for ipf.

Usage

```
computeLinear(curValue, target, x, w, boundLinear = 10)
computeLinearG1(curValue, target, x, w, boundLinear = 10)
computeFrac(curValue, target, x, w)
```

Arguments

curValue	Current summed up value. Same as sum(x*w)
target	Target value. An element of conP in ipf
х	Vector of numeric values to be calibrated against
W	Vector of weights
boundLinear	The output f will satisfy 1/boundLinear <= f <= boundLinear. See bound in ipf

Details

computeFrac provides the "standard" IPU updating scheme given as

f = target/curValue

which means that each weight inside the level will be multiplied by the same factor when doing the actual update step (w := f*w). computeLinear on the other hand calculates f as

$$f_i = ax_i + b$$

where a and b are chosen, so f satisfies the following two equations.

$$\sum f_i * w_i * x_i = target$$
$$\sum f_i * w_i = \sum w_i$$

computeLinearG1 calculates f in the same way as computeLinear, but if $f_i*w_i<1$ f_i will be set to $1/w_i$.

Value

A weight multiplier f

```
cpp_mean
```

Calculate mean by factors

Description

These functions calculate the arithmetic and geometric mean of the weight for each class. geometric_mean and arithmetic_mean return a numeric vector of the same length as w which stores the averaged weight for each observation. geometric_mean_reference returns the same value by reference, i.e. the input value w gets overwritten by the updated weights. See examples.

Usage

geometric_mean_reference(w, classes)

Arguments

W	An numeric vector. All entries should be positive.
classes	A factor variable. Must have the same length as w.

Examples

Not run:

```
## create random data
nobs <- 10
classLabels <- letters[1:3]
dat = data.frame(
   weight = exp(rnorm(nobs)),
   household = factor(sample(classLabels, nobs, replace = TRUE))</pre>
```

demo.eusilc

```
) dat
## calculate weights with geometric_mean
geom_weight <- geometric_mean(dat$weight, dat$household)
cbind(dat, geom_weight)
## calculate weights with arithmetic_mean
arith_weight <- arithmetic_mean(dat$weight, dat$household)
cbind(dat, arith_weight)
## calculate weights "by reference"
geometric_mean_reference(dat$weight, dat$household)
dat
## End(Not run)</pre>
```

demo.eusilc

Generate multiple years of EU-SILC data

Description

Create a dummy dataset to be used for demonstrating the functionalities of the surveysd package based on laeken::eusilc. Please refer to the documentation page of the original data for details about the variables.

Usage

demo.eusilc(n = 8, prettyNames = FALSE)

Arguments

n	Number of years to generate. Should be at least 1
prettyNames	Create easy-to-read names for certain variables. Recommended for demonstra-
	tion purposes. Otherwise, use the original codes documented in laeken::eusilc.

Details

If prettyNames is TRUE, the following variables will be available in an easy-to-read manner.

- hid Household id. Consistent with respect to the reference period (year)
- hsize Size of the household. derived from hid and period
- · region Federal state of austria where the household is located
- pid Personal id. Consistent with respect to the reference period (year)
- age Age-class of the respondent
- gender A persons gender ("male", "Female")
- ecoStat Ecnomic status ("part time", "full time", "unemployed", ...)

- citizenship Citizenship ("AT", "EU", "other")
- pWeight Personal sample weight inside the reference period
- year. Simulated reference period
- povertyRisk. Logical variable determining whether a respondent is at risk of poverty

Examples

```
demo.eusilc(n = 1, prettyNames = TRUE)[, c(1:8, 26, 28:30)]
```

draw.bootstrap Draw bootstrap replicates

Description

Draw bootstrap replicates from survey data with rotating panel design. Survey information, like ID, sample weights, strata and population totals per strata, should be specified to ensure meaningfull survey bootstraping.

Usage

```
draw.bootstrap(
    dat,
    REP = 1000,
    hid = NULL,
    weights,
    period = NULL,
    strata = NULL,
    cluster = NULL,
    totals = NULL,
    single.PSU = c("merge", "mean"),
    boot.names = NULL,
    split = FALSE,
    pid = NULL,
    new.method = FALSE
)
```

Arguments

dat	either data.frame or data.table containing the survey data with rotating panel design.
REP	integer indicating the number of bootstrap replicates.
hid	character specifying the name of the column in dat containing the household id. If NULL (the default), the household structure is not regarded.
weights	character specifying the name of the column in dat containing the sample weights

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- strata character vector specifying the name(s) of the column in dat by which the population was stratified. If strata is a vector stratification will be assumed as the combination of column names contained in strata. Setting in addition cluster not NULL stratification will be assumed on multiple stages, where each additional entry in strata specifies the stratification variable for the next lower stage. see Details for more information.
- cluster character vector specifying cluster in the data. If not already specified in cluster household ID is taken es the lowest level cluster.
- totals character specifying the name of the column in dat containing the the totals per strata and/or cluster. Is ONLY optional if cluster is NULL or equal hid and strata contains one columnname! Then the households per strata will be calcualted using the weights argument. If clusters and strata for multiple stages are specified totals needs to be a vector of length(strata) specifying the column on dat that contain the total number of PSUs at each stage. totals is interpreted from left the right, meaning that the first argument corresponds to the number of PSUs at the first and the last argument to the number of PSUs at the last stage.
- single.PSU either "merge" or "mean" defining how single PSUs need to be dealt with. For single.PSU="merge" single PSUs at each stage are merged with the strata or cluster with the next least number of PSUs. If multiple of those exist one will be select via random draw. For single.PSU="mean" single PSUs will get the mean over all bootstrap replicates at the stage which did not contain single PSUs.
- boot.names character indicating the leading string of the column names for each bootstrap replica. If NULL defaults to "w".
- split logical, if TRUE split households are considered using pid, for more information see Details.
- pid column in dat specifying the personal identifier. This identifier needs to be unique for each person throught the whole data set.
- new.method logical, if TRUE bootstrap replicates will never be negative even if in some strata the whole population is in the sample. WARNING: This is still experimental and resulting standard errors might be underestimated! Use this if for some strata the whole population is in the sample!

draw.bootstrap takes dat and draws REP bootstrap replicates from it. dat must be household data where household members correspond to multiple rows with the same household identifier. For most practical applications, the following columns should be available in the dataset and passed via the corresponding parameters:

- Column indicating the sample period (parameter period).
- Column indicating the household ID (parameter hid)
- Column containing the household sample weights (parameter weights);

Columns by which population was stratified during the sampling process (parameter: strata).

For single stage sampling design a column the argument totals is optional, meaning that a column of the number of PSUs at the first stage does not need to be supplied. For this case the number of PSUs is calculated and added to dat using strata and weights. By setting cluster to NULL single stage sampling design is always assumed and if strata contains of multiple column names the combination of all those column names will be used for stratification.

In the case of multi stage sampling design the argument totals needs to be specified and needs to have the same number of arguments as strata.

If cluster is NULL or does not contain hid at the last stage, hid will automatically be used as the final cluster. If, besides hid, clustering in additional stages is specified the number of column names in strata and cluster (including hid) must be the same. If for any stage there was no clustering or stratification one can set "1" or "I" for this stage.

For example strata=c("REGION", "I"), cluster=c("MUNICIPALITY", "HID") would speficy a 2 stage sampling design where at the first stage the municipalities where drawn stratified by regions and at the 2nd stage housholds are drawn in each municipality without stratification.

Bootstrap replicates are drawn for each survey period (period) using the function rescaled.bootstrap. Afterwards the bootstrap replicates for each household are carried forward from the first period the household enters the survey to all the censecutive periods it stays in the survey.

This ensures that the bootstrap replicates follow the same logic as the sampled households, making the bootstrap replicates more comparable to the actual sample units.

If split ist set to TRUE and pid is specified, the bootstrap replicates are carried forward using the personal identifiers instead of the houshold identifier. This takes into account the issue of a houshold splitting up. Any person in this new split household will get the same bootstrap replicate as the person that has come from an other household in the survey. People who enter already existing households will also get the same bootstrap replicate as the other households members had in the previous periods.

Value

the survey data with the number of REP bootstrap replicates added as columns.

Returns a data.table containing the original data as well as the number of REP columns containing the bootstrap replicates for each repetition.

The columns of the bootstrap replicates are by default labeled "w*Number*" where *Number* goes from 1 to REP. If the column names of the bootstrap replicates should start with a different character or string the parameter boot.names can be used.

Author(s)

Johannes Gussenbauer, Alexander Kowarik, Statistics Austria

See Also

data.table for more information on data.table objects.

draw.bootstrap

Examples

```
## Not run:
eusilc <- demo.eusilc(prettyNames = TRUE)</pre>
## draw sample without stratification or clustering
dat_boot <- draw.bootstrap(eusilc, REP = 10, weights = "pWeight",</pre>
                            period = "year")
## use stratification w.r.t. region and clustering w.r.t. households
dat_boot <- draw.bootstrap(</pre>
 eusilc, REP = 10, hid = "hid", weights = "pWeight",
 strata = "region", period = "year")
## use multi-level clustering
dat_boot <- draw.bootstrap(</pre>
 eusilc, REP = 10, hid = "hid", weights = "pWeight",
 strata = c("region", "age"), period = "year")
# create spit households
eusilc[, pidsplit := pid]
year <- eusilc[, unique(year)]</pre>
year <- year[-1]
leaf_out <- c()</pre>
for(y in year) {
 split.person <- eusilc[</pre>
   year == (y-1) & !duplicated(hid) & !(hid %in% leaf_out),
   sample(pid, 20)
 ٦
 overwrite.person <- eusilc[</pre>
    (year == (y)) & !duplicated(hid) & !(hid %in% leaf_out),
    .(pid = sample(pid, 20))
 ٦
 overwrite.person[, c("pidsplit", "year_curr") := .(split.person, y)]
 eusilc[overwrite.person, pidsplit := i.pidsplit,
         on = .(pid, year >= year_curr)]
 leaf_out <- c(leaf_out,</pre>
                eusilc[pid %in% c(overwrite.person$pid,
                                   overwrite.person$pidsplit),
                unique(hid)])
}
dat_boot <- draw.bootstrap(</pre>
 eusilc, REP = 10, hid = "hid", weights = "pWeight",
 strata = c("region", "age"), period = "year", split = TRUE,
 pid = "pidsplit")
# split households were considered e.g. household and
# split household were both selected or not selected
dat_boot[, data.table::uniqueN(w1), by = pidsplit][V1 > 1]
## End(Not run)
```

generate.HHID

Generate new houshold ID for survey data with rotating panel design taking into account split households

Description

Generating a new houshold ID for survey data using a houshold ID and a personal ID. For surveys with rotating panel design containing housholds, houshold members can move from an existing household to a new one, that was not originally in the sample. This leads to the creation of so called split households. Using a peronal ID (that stays fixed over the whole survey), an indicator for different time steps and a houshold ID, a new houshold ID is assigned to the original and the split household.

Usage

```
generate.HHID(dat, period = "RB010", pid = "RB030", hid = "DB030")
```

Arguments

dat	data table of data frame containing the survey data
period	column name of dat containing an indicator for the rotations, e.g years, quarters, months, ect
pid	column name of dat containing the personal identifier. This needs to be fixed for an indiviual throught the whole survey
hid	column name of dat containing the household id. This needs to for a household throught the whole survey

Value

the survey data dat as data.table object containing a new and an old household ID. The new household ID which considers the split households is now named hid and the original household ID has a trailing "_orig".

Examples

```
## Not run:
library(surveysd)
library(laeken)
library(data.table)
eusilc <- surveysd:::demo.eusilc(n=4)
# create spit households
eusilc[,rb030split:=rb030]
year <- eusilc[,unique(year)]
year <- year[-1]</pre>
```

```
leaf_out <- c()</pre>
for(y in year) {
 split.person <- eusilc[year==(y-1)&!duplicated(db030)&!db030%in%leaf_out,</pre>
                          sample(rb030,20)]
 overwrite.person <- eusilc[year==(y)&!duplicated(db030)&!db030%in%leaf_out,</pre>
                              .(rb030=sample(rb030,20))]
 overwrite.person[,c("rb030split","year_curr"):=.(split.person,y)]
 eusilc[overwrite.person,
         rb030split:=i.rb030split,on=.(rb030,year>=year_curr)]
 leaf_out <- c(</pre>
    leaf_out,
    eusilc[rb030%in%c(overwrite.person$rb030,overwrite.person$rb030split),
    unique(db030)])
}
# pid which are in split households
eusilc[,.(uniqueN(db030)),by=list(rb030split)][V1>1]
eusilc.new <- generate.HHID(eusilc, period = "year", pid = "rb030split",</pre>
                             hid = "db030")
# no longer any split households in the data
eusilc.new[,.(uniqueN(db030)),by=list(rb030split)][V1>1]
## End(Not run)
```

ipf

Iterative Proportional Fitting

Description

Adjust sampling weights to given totals based on household-level and/or individual level constraints.

Usage

```
ipf(
  dat,
  hid = NULL,
  conP = NULL,
  conH = NULL,
  epsP = 1e-06,
  epsH = 0.01,
  verbose = FALSE,
  w = NULL,
  bound = 4,
  maxIter = 200,
```

```
meanHH = TRUE,
allPthenH = TRUE,
returnNA = TRUE,
looseH = FALSE,
numericalWeighting = computeLinear,
check_hh_vars = TRUE,
conversion_messages = FALSE,
nameCalibWeight = "calibWeight",
minMaxTrim = NULL
)
```

Arguments

dat	a data.table containing household ids (optionally), base weights (optionally), household and/or personal level variables (numerical or categorical) that should be fitted.
hid	name of the column containing the household-ids within dat or NULL if such a variable does not exist.
conP	list or (partly) named list defining the constraints on person level. The list ele- ments are contingency tables in array representation with dimnames correspond- ing to the names of the relevant calibration variables in dat. If a numerical vari- able is to be calibrated, the respective list element has to be named with the name of that numerical variable. Otherwise the list element shoud NOT be named.
conH	list or (partly) named list defining the constraints on household level. The list elements are contingency tables in array representation with dimnames corre- sponding to the names of the relevant calibration variables in dat. If a numeri- cal variable is to be calibrated, the respective list element has to be named with the name of that numerical variable. Otherwise the list element shoud NOT be named.
epsP	numeric value or list (of numeric values and/or arrays) specifying the conver- gence limit(s) for conP. The list can contain numeric values and/or arrays which must appear in the same order as the corresponding constraints in conP. Also, an array must have the same dimensions and dimnames as the corresponding constraint in conP.
epsH	numeric value or list (of numeric values and/or arrays) specifying the conver- gence limit(s) for conH. The list can contain numeric values and/or arrays which must appear in the same order as the corresponding constraints in conH. Also, an array must have the same dimensions and dimnames as the corresponding constraint in conH.
verbose	if TRUE, some progress information will be printed.
W	name if the column containing the base weights within dat or NULL if such a variable does not exist. In the latter case, every observation in dat is assigned a starting weight of 1.
bound	numeric value specifying the multiplier for determining the weight trimming boundary if the change of the base weights should be restricted, i.e. if the weights should stay between 1/bound*w and bound*w.

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maxIter	numeric value specifying the maximum number of iterations that should be per- formed.	
meanHH	if TRUE, every person in a household is assigned the mean of the person weights corresponding to the household. If "geometric", the geometric mean is used rather than the arithmetic mean.	
allPthenH	if TRUE, all the person level calibration steps are performed before the houshold level calibration steps (and meanHH, if specified). If FALSE, the houshold level calibration steps (and meanHH, if specified) are performed after everey person level calibration step. This can lead to better convergence properties in certain cases but also means that the total number of calibration steps is increased.	
returnNA	if TRUE, the calibrated weight will be set to NA in case of no convergence.	
looseH	if FALSE, the actual constraints conH are used for calibrating all the hh weights. If TRUE, only the weights for which the lower and upper thresholds defined by conH and epsH are exceeded are calibrated. They are however not calibrated against the actual constraints conH but against these lower and upper thresholds, i.e. conH-conH*epsH and conH+conH*epsH.	
numericalWeigh	-	
	See numericalWeighting	
check_hh_vars	If TRUE check for non-unique values inside of a household for variables in household constraints	
conversion_mes	-	
	show a message, if inputs need to be reformatted. This can be useful for speed optimizations if ipf is called several times with similar inputs (for example boot- strapping)	
nameCalibWeight		
	character defining the name of the variable for the newly generated calibrated weight.	
minMaxTrim	numeric vector of length2, first element a minimum value for weights to be trimmed to, second element a maximum value for weights to be trimmed to.	

This function implements the weighting procedure described here: doi:10.17713/ajs.v45i3.120. Usage examples can be found in the corresponding vignette (vignette("ipf")).

conP and conH are contingency tables, which can be created with xtabs. The dimnames of those tables should match the names and levels of the corresponding columns in dat.

maxIter, epsP and epsH are the stopping criteria. epsP and epsH describe relative tolerances in the sense that w_{i+1}

$$1 - epsP < \frac{w_{i+1}}{w_i} < 1 + epsP$$

will be used as convergence criterium. Here i is the iteration step and wi is the weight of a specific person at step i.

The algorithm performs best if all varables occuring in the constraints (conP and conH) as well as the household variable are coded as factor-columns in dat. Otherwise, conversions will be necessary which can be monitored with the conversion_messages argument. Setting check_hh_vars to FALSE can also incease the performance of the scheme.

The function will return the input data dat with the calibrated weights calibWeight as an additional column as well as attributes. If no convergence has been reached in maxIter steps, and returnNA is TRUE (the default), the column calibWeights will only consist of NAs. The attributes of the table are attributes derived from the data.table class as well as the following.

converged	Did the algorithm converge in maxIter steps?
iterations	The number of iterations performed.
conP, conH, epsP, epsH	See Arguments.
conP_adj, conH_adj	Adjusted versions of conP and conH
formP,formH	Formulas that were used to calculate conP_adj and conH_adj based on the output table.

Author(s)

Alexander Kowarik, Gregor de Cillia

Examples

```
## Not run:
# load data
eusilc <- demo.eusilc(n = 1, prettyNames = TRUE)</pre>
# personal constraints
conP1 <- xtabs(pWeight ~ age, data = eusilc)</pre>
conP2 <- xtabs(pWeight ~ gender + region, data = eusilc)</pre>
conP3 <- xtabs(pWeight*eqIncome ~ gender, data = eusilc)</pre>
# household constraints
conH1 <- xtabs(pWeight ~ hsize + region, data = eusilc)</pre>
# simple usage ------
calibweights1 <- ipf(</pre>
 eusilc,
 conP = list(conP1, conP2, eqIncome = conP3),
 bound = NULL,
 verbose = TRUE
)
# compare personal weight with the calibweigth
calibweights1[, .(hid, pWeight, calibWeight)]
# advanced usage ------
# use an array of tolerances
epsH1 <- conH1
epsH1[1:4, ] <- 0.005
epsH1[5, ] <- 0.2
```

create an initial weight for the calibration

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ipf_step

```
eusilc[, regSamp := .N, by = region]
eusilc[, regPop := sum(pWeight), by = region]
eusilc[, baseWeight := regPop/regSamp]
calibweights2 <- ipf(</pre>
  eusilc,
  conP = list(conP1, conP2),
  conH = list(conH1),
  epsP = 1e-6,
  epsH = list(epsH1),
  bound = 4,
  w = "baseWeight",
  verbose = TRUE
)
# show an adjusted version of conP and the original
attr(calibweights2, "conP_adj")
attr(calibweights2, "conP")
## End(Not run)
```

ipf_step

Perform one step of iterative proportional updating

Description

C++ routines to invoke a single iteration of the Iterative proportional updating (IPU) scheme. Targets and classes are assumed to be one dimensional in the ipf_step functions. combine_factors aggregates several vectors of type factor into a single one to allow multidimensional ipu-steps. See examples.

Usage

```
ipf_step_ref(w, classes, targets)
ipf_step(w, classes, targets)
ipf_step_f(w, classes, targets)
combine_factors(dat, targets)
```

W	a numeric vector of weights. All entries should be positive.
classes	a factor variable. Must have the same length as w.
targets	key figure to target with the ipu scheme. A numeric verctor of the same length as levels(classes). This can also be a table produced by xtabs. See examples.
dat	a data.frame containing the factor variables to be combined.

ipf_step returns the adjusted weights. ipf_step_ref does the same, but updates w by reference rather than returning. ipf_step_f returns a multiplicator: adjusted weights divided by unadjusted weights. combine_factors is designed to make ipf_step work with contingency tables produced by xtabs.

Examples

```
## create random data
nobs <- 10
classLabels <- letters[1:3]</pre>
dat = data.frame(
  weight = exp(rnorm(nobs)),
  household = factor(sample(classLabels, nobs, replace = TRUE))
)
dat
## create targets (same lenght as classLabels!)
targets <- 3:5
## calculate weights
new_weight <- ipf_step(dat$weight, dat$household, targets)</pre>
cbind(dat, new_weight)
## check solution
xtabs(new_weight ~ dat$household)
## calculate weights "by reference"
ipf_step_ref(dat$weight, dat$household, targets)
dat
## load data
factors <- c("time", "sex", "smoker", "day")</pre>
tips <- data.frame(sex=c("Female","Male","Male"), day=c("Sun","Mon","Tue"),</pre>
time=c("Dinner","Lunch","Lunch"), smoker=c("No","Yes","No"))
tips <- tips[factors]</pre>
## combine factors
con <- xtabs(~., tips)</pre>
cf <- combine_factors(tips, con)</pre>
cbind(tips, cf)[sample(nrow(tips), 10, replace = TRUE),]
## adjust weights
weight <- rnorm(nrow(tips)) + 5</pre>
adjusted_weight <- ipf_step(weight, cf, con)</pre>
## check outputs
```

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kishFactor

```
con2 <- xtabs(adjusted_weight ~ ., data = tips)
sum((con - con2)^2)</pre>
```

kishFactor

Description

Compute the design effect due to unequal weighting.

Kish Factor

Usage

kishFactor(w, na.rm = FALSE)

Arguments

W	a numeric vector with weights
na.rm	a logical value indicating whether NA values should be stripped before the com-
	putation proceeds.

Details

The factor is computed acording to 'Weighting for Unequal P_i', Leslie Kish, Journal of Official Statistics, Vol. 8. No. 2, 1992

$$deff = \sqrt{n} \sum_{j} w_j^2 / (\sum_{j} w_j)^2$$

Value

The function will return the the kish factor

Author(s)

Alexander Kowarik

Examples

kishFactor(rep(1,10))
kishFactor(rlnorm(10))

plot.surveysd

Description

Plot results of calc.stError()

Usage

```
## S3 method for class 'surveysd'
plot(
    x,
    variable = x$param$var[1],
    type = c("summary", "grouping"),
    groups = NULL,
    sd.type = c("dot", "ribbon"),
    ...
)
```

х	object of class 'surveysd' output of function calc.stError
variable	Name of the variable for which standard errors have been calcualated in dat
type	can bei either "summary" or "grouping", default value is "summary". For "summary" a barplot is created giving an overview of the number of estimates having the flag smallGroup, cvHigh, both or none of them. For 'grouping' results for point estimate and standard error are plotted for pre defined groups.
groups	If type='grouping' variables must be defined by which the data is grouped. Only 2 levels are supported as of right now. If only one group is defined the higher group will be the estimate over the whole period. Results are plotted for the first argument in groups as well as for the combination of groups[1] and groups[2].
sd.type	can bei either 'ribbon' or 'dot' and is only used if type='grouping'. Default is "dot" For sd.type='dot' point estimates are plotted and flagged if the corre- sponding standard error and/or the standard error using the mean over k-periods exceeded the value cv.limit (see calc.stError). For sd.type='ribbon' the point estimates including ribbons, defined by point estimate +- estimated stan- dard error are plotted. The calculated standard errors using the mean over k peri- ods are plotted using less transparency. Results for the higher level (~groups[1]) are coloured grey.
	additional arguments supplied to plot.

PointEstimates

Examples

```
library(surveysd)
set.seed(1234)
eusilc <- demo.eusilc(n = 3, prettyNames = TRUE)</pre>
dat_boot <- draw.bootstrap(eusilc, REP = 3, hid = "hid", weights = "pWeight",</pre>
                            strata = "region", period = "year")
# calibrate weight for bootstrap replicates
dat_boot_calib <- recalib(dat_boot, conP.var = "gender", conH.var = "region")</pre>
# estimate weightedRatio for povmd60 per period
group <- list("gender", "region", c("gender", "region"))</pre>
err.est <- calc.stError(dat_boot_calib, var = "povertyRisk",</pre>
                         fun = weightedRatio,
                         group = group , period.mean = NULL)
plot(err.est)
# plot results for gender
# dotted line is the result on the national level
plot(err.est, type = "grouping", groups = "gender")
# plot results for rb090 in each db040
# with standard errors as ribbons
plot(err.est, type = "grouping", groups = c("gender", "region"), sd.type = "ribbon")
```

PointEstimates Weighted Point Estimates

Description

Predefined functions for weighted point estimates in package surveysd.

Usage

weightedRatio(x, w)

weightedSum(x, w)

Х	numeric vector
W	weight vector

recalib

Details

Predefined functions are weighted ratio and weighted sum.

Value

Each of the functions return a single numeric value

Examples

```
x <- 1:10
w <- 10:1
weightedRatio(x,w)
x <- 1:10
w <- 10:1
weightedSum(x,w)
```

print.surveysd Print function for surveysd objects

Description

Prints the results of a call to calc.stError. Shows used variables and function, number of point estiamtes as well as properties of the results.

Usage

S3 method for class 'surveysd'
print(x, ...)

Arguments

х	an object of class 'surveysd'
	additonal parameters

|--|

Description

Calibrate weights for bootstrap replicates by using iterative proportional updating to match population totals on various household and personal levels.

recalib

Usage

```
recalib(
  dat,
  hid = attr(dat, "hid"),
  weights = attr(dat, "weights"),
  b.rep = attr(dat, "b.rep"),
  period = attr(dat, "period"),
  conP.var = NULL,
  conH.var = NULL,
  conH = NULL,
  conH = NULL,
  epsP = 0.01,
  epsH = 0.02,
  ...
)
```

recalib takes survey data (dat) containing the bootstrap replicates generated by draw.bootstrap and calibrates weights for each bootstrap replication according to population totals for person- or household-specific variables.

dat must be household data where household members correspond to multiple rows with the same household identifier. The data should at least containt the following columns:

- Column indicating the sample period;
- Column indicating the household ID;
- Column containing the household sample weights;
- Columns which contain the bootstrap replicates (see output of draw.bootstrap);
- Columns indicating person- or household-specific variables for which sample weight should be adjusted.

For each period and each variable in conP.var and/or conH.var contingency tables are estimated to get margin totals on personal- and/or household-specific variables in the population. Afterwards the bootstrap replicates are multiplied with the original sample weight and the resulting product ist then adjusted using ipf() to match the previously calcualted contingency tables. In this process the columns of the bootstrap replicates are overwritten by the calibrated weights.

Value

Returns a data.table containing the survey data as well as the calibrated weights for the bootstrap replicates. The original bootstrap replicates are overwritten by the calibrated weights. If calibration of a bootstrap replicate does not converge the bootsrap weight is not returned and numeration of the returned bootstrap weights is reduced by one.

Author(s)

Johannes Gussenbauer, Alexander Kowarik, Statistics Austria

See Also

ipf() for more information on iterative proportional fitting.

Examples

```
## Not run:
```

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

rescaled.bootstrap Draw bootstrap replicates

Description

Draw bootstrap replicates from survey data using the rescaled bootstrap for stratified multistage sampling, presented by Preston, J. (2009).

Usage

```
rescaled.bootstrap(
    dat,
    REP = 1000,
    strata = "DB050>1",
    cluster = "DB060>DB030",
    fpc = "N.cluster>N.households",
    single.PSU = c("merge", "mean"),
    return.value = c("data", "replicates"),
    check.input = TRUE,
    new.method = FALSE
)
```

dat	either data frame or data table containing the survey sample
REP	integer indicating the number of bootstraps to be drawn
strata	string specifying the column name in dat that is used for stratification. For mul- tistage sampling multiple column names can be specified by strata=c("strata1>strata2>strata3"). See Details for more information.

cluster	string specifying the column name in dat that is used for clustering. For in- stance given a household sample the column containing the household ID should be supplied. For multistage sampling multiple column names can be specified by cluster=c("cluster1>cluster2>cluster3"). See Details for more in- formation.
fpc	string specifying the column name in dat that contains the number of PSUs at the first stage. For multistage sampling the number of PSUs at each stage must be specified by strata=c("fpc1>fpc2>fpc3").
single.PSU	either "merge" or "mean" defining how single PSUs need to be dealt with. For single.PSU="merge" single PSUs at each stage are merged with the strata or cluster with the next least number of PSUs. If multiple of those exist one will be select via random draw. For single.PSU="mean" single PSUs will get the mean over all bootstrap replicates at the stage which did not contain single PSUs.
return.value	either "data" or "replicates" specifying the return value of the function. For "data" the survey data is returned as class data.table, for "replicates" only the bootstrap replicates are returned as data.table.
check.input	logical, if TRUE the input will be checked before applying the bootstrap procedure
new.method	logical, if TRUE bootstrap replicates will never be negative even if in some strata the whole population is in the sample. WARNING: This is still experimental and resulting standard errors might be underestimated! Use this if for some strata the whole population is in the sample!

For specifying multistage sampling designs the column names in strata, cluster and fpc need to separated by ">".

For multistage sampling the strings are read from left to right meaning that the column name before the first ">" is taken as the column for stratification/clustering/number of PSUs at the first and the column after the last ">" is taken as the column for stratification/clustering/number of PSUs at the last stage. If for some stages the sample was not stratified or clustered one must specify this by "1" or "I", e.g. strata=c("strata1>I>strata3") if there was no stratification at the second stage or cluster=c("cluster1>cluster2>I") if there were no clusters at the last stage.

The number of PSUs at each stage is not calculated internally and must be specified for any sampling design. For single stage sampling using stratification this can usually be done by adding over all sample weights of each PSU by each strata-code.

Spaces in each of the strings will be removed, so if column names contain spaces they should be renamed before calling this procedure!

Value

returns the complete data set including the bootstrap replicates or just the bootstrap replicates, depending on return.value="data" or return.value="replicates" respectively.

Author(s)

Johannes Gussenbauer, Statistics Austria

rescaled.bootstrap

References

Preston, J. (2009). Rescaled bootstrap for stratified multistage sampling. Survey Methodology. 35. 227-234.

Examples

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