Package 'jipApprox'

June 12, 2022

Title Approximate Inclusion Probabilities for Survey Sampling

Version 0.1.4

Date 2022-06-11

Description Approximate joint-inclusion probabilities in Unequal Probability Sampling, or compute Monte Carlo approximations of the first and second-order inclusion probabilities of a general sampling design as in Fattorini (2006) <doi:10.1093/biomet/93.2.269>.

Depends R (>= 4.0.0)

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Encoding UTF-8

BugReports https://github.com/rhobis/jipApprox/issues

RoxygenNote 7.1.1

Imports sampling

NeedsCompilation no

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Repository CRAN

Date/Publication 2022-06-12 07:00:02 UTC

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HTvar

Description

Compute or estimate the variance of the Horvitz-Thompson total estimator by the Horvitz-Thompson or Sen-Yates-Grundy variance estimators.

Usage

HTvar(y, pikl, sample = TRUE, method = "HT")

Arguments

У	numeric vector representing the variable of interest
pikl	matrix of second-order (joint) inclusion probabilities; the diagonal must contain the first-order inclusion probabilities.
sample	logical value indicating whether sample or population values are provided. If sample=TRUE, the function returns a sample estimate of the variance, while if sample=FALSE, the Variance is computed over all population units. Default is TRUE.
method	string, indicating if the Horvitz-Thompson ("HT") or the Sen-Yates-Grundy ("SYG") estimator should be computed.

Details

The Horvitz-Thompson variance is defined as

$$\sum_{i \in U} \sum_{j \in U} \frac{(\pi_{ij} - \pi_i \pi_j)}{\pi_i \pi_j} y_i y_j$$

which is estimated by

$$\sum_{i \in U} \sum_{j \in U} \frac{(\pi_{ij} - \pi_i \pi_j)}{\pi_i \pi_j \pi_{ij}} y_i y_j$$

The Sen-Yates-Grundy variance is obtained from the Horvitz-Thompson variance by conditioning on the sample size n, and is therefore only appliable to fixed size sampling designs:

$$\sum_{i \in U} \sum_{j > i} (\pi_i \pi_j - \pi_{ij}) \left(\frac{y_i}{\pi_i} - \frac{y_j}{\pi_j} \right)^2$$

Its estimator is

$$\sum_{i \in U} \sum_{j > i} \frac{(\pi_i \pi_j - \pi_{ij})}{\pi_{ij}} \left(\frac{y_i}{\pi_i} - \frac{y_j}{\pi_j} \right)^2$$

jipApprox

Examples

```
### Generate population data ---
N <- 500; n <- 50
set.seed(0)
x <- rgamma(500, scale=10, shape=5)
y <- abs( 2*x + 3.7*sqrt(x) * rnorm(N) )
pik <- n * x/sum(x)
pikl <- jip_approx(pik, method='Hajek')
### Dummy sample ---
s <- sample(N, n)
### Compute Variance ---
HTvar(y=y, pikl=pikl, sample=FALSE, method="HT")
HTvar(y=y, pikl=pikl, sample=FALSE, method="SYG")
### Estimate Variance ---
#' HTvar(y=y[s], pikl=pikl[s,s], sample=TRUE, method="HT")
#' HTvar(y=y[s], pikl=pikl[s,s], sample=TRUE, method="SYG")</pre>
```

jipApprox

jipApprox: Approximate inclusion probabilities for survey sampling

Description

Approximate joint-inclusion probabilities in Unequal Probability Sampling, or compute Monte Carlo approximations of the first and second-order inclusion probabilities of a general sampling design as in Fattorini (2006) <doi:10.1093/biomet/93.2.269>.

Approximation of Joint-inclusion probabilities

Function jip_approx provides a number of approximations of the second-order inclusion probabilities that require only the first-order inclusion probabilities. These approximations may be employed in unequal probability sampling design with high entropy. A more flexible approximation may be obtained by using function jip_MonteCarlo, which estimates inclusion probabilities through a Monte Carlo simulation.

The variance of the Horvitz-Thompson total estimator may be then estimated by plugging the approximated joint probabilities into the Horvitz-Thompson or Sen-Yates-Grundy variance estimator using function HTvar.

References

Matei, A.; Tillé, Y., 2005. Evaluation of variance approximations and estimators in maximum entropy sampling with unequal probability and fixed sample size. Journal of Official Statistics 21 (4), 543-570.

Haziza, D.; Mecatti, F.; Rao, J.N.K. 2008. Evaluation of some approximate variance estimators under the Rao-Sampford unequal probability sampling design. Metron LXVI (1), 91-108.

Fattorini, L. 2006. Applying the Horvitz-Thompson criterion in complex designs: A computerintensive perspective for estimating inclusion probabilities. Biometrika 93 (2), 269-278

jipDFtoM

Transform a Joint-Inclusion Probability data.frame to a matrix

Description

Transform a Joint-Inclusion Probability data.frame to a matrix

Usage

jipDFtoM(jip, symmetric = TRUE)

Arguments

jip	vector or data.frame containing the joint-inclusion probabilities
symmetric	boolean, if TRUE, returns a symmetric matrix, otherwise, an upper triangular
	matrix

Value

a symmetric matrix of joint-inclusion probabilities if TRUE, otherwise, an upper triangular matrix

jipMtoDF

Transform a matrix of Joint-Inclusion Probabilities to a data.frame

Description

Transform a matrix of Joint-Inclusion Probabilities to a data.frame

Usage

jipMtoDF(jip, id = NULL)

Arguments

jip	a square matrix of joint-inclusion probabilities, symmetric or upper-triangular
id	optional, vector of id labels, its length should be equal to ncol(jip) and nrow(jip)

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jip_approx

Description

Approximations of joint-inclusion probabilities by means of first-order inclusion probabilities.

Usage

```
jip_approx(pik, method)
```

Arguments

pik	numeric vector of first-order inclusion probabilities for all population units.
method	string representing one of the available approximation methods.

Details

Available methods are "Hajek", "HartleyRao", "Tille", "Brewer1", "Brewer2", "Brewer3", and "Brewer4". Note that these methods were derived for high-entropy sampling designs, therefore they could have low performance under different designs.

Hájek (1964) approximation [method="Hajek"] is derived under Maximum Entropy sampling design and is given by

$$\tilde{\pi}_{ij} = \pi_i \pi_j \frac{1 - (1 - \pi_i)(1 - \pi_j)}{d}$$

where $d = \sum_{i \in U} \pi_i (1 - \pi_i)$

Hartley and Rao (1962) proposed the following approximation under randomised systematic sampling [method="HartleyRao"]:

$$\begin{split} \tilde{\pi}_{ij} &= \frac{n-1}{n} \pi_i \pi_j + \frac{n-1}{n^2} (\pi_i^2 \pi_j + \pi_i \pi_j^2) - \frac{n-1}{n^3} \pi_i \pi_j \sum_{i \in U} \pi_j^2 \\ &+ \frac{2(n-1)}{n^3} (\pi_i^3 \pi_j + \pi_i \pi_j^3 + \pi_i^2 \pi_j^2) - \frac{3(n-1)}{n^4} (\pi_i^2 \pi_j + \pi_i \pi_j^2) \sum_{i \in U} \pi_i^2 \\ &+ \frac{3(n-1)}{n^5} \pi_i \pi_j \left(\sum_{i \in U} \pi_i^2 \right)^2 - \frac{2(n-1)}{n^4} \pi_i \pi_j \sum_{i \in U} \pi_j^3 \end{split}$$

Tillé (1996) proposed the approximation $\tilde{\pi}_{ij} = \beta_i \beta_j$, where the coefficients β_i are computed iteratively through the following procedure [method="Tille"]:

1.
$$\beta_i^{(0)} = \pi_i, \ \forall i \in U$$

jip_approx

2.
$$\beta_i^{(2k-1)} = \frac{(n-1)\pi_i}{\beta^{(2k-2)} - \beta_i^{(2k-2)}}$$

3. $\beta_i^{2k} = \beta_i^{(2k-1)} \left(\frac{n(n-1)}{(\beta^{(2k-1)})^2 - \sum_{i \in U} (\beta_k^{(2k-1)})^2} \right)^{(1/2)}$

with $\beta^{(k)} = \sum_{i \in U} \beta_i^i, \ k = 1, 2, 3, \dots$

Finally, Brewer (2002) and Brewer and Donadio (2003) proposed four approximations, which are defined by the general form

$$\tilde{\pi}_{ij} = \pi_i \pi_j (c_i + c_j)/2$$

where the c_i determine the approximation used:

• Equation (9) [method="Brewer1"]:

$$c_i = (n-1)/(n-\pi_i)$$

• Equation (10) [method="Brewer2"]:

$$c_i = (n-1)/\left(n - n^{-1} \sum_{i \in U} \pi_i^2\right)$$

• Equation (11) [method="Brewer3"]:

$$c_i = (n-1)/\left(n - 2\pi_i + n^{-1}\sum_{i \in U} \pi_i^2\right)$$

• Equation (18) [method="Brewer4"]:

$$c_i = (n-1) / \left(n - (2n-1)(n-1)^{-1} \pi_i + (n-1)^{-1} \sum_{i \in U} \pi_i^2 \right)$$

Value

A symmetric matrix of inclusion probabilities, which diagonal is the vector of first-order inclusion probabilities.

References

Hartley, H.O.; Rao, J.N.K., 1962. Sampling With Unequal Probability and Without Replacement. The Annals of Mathematical Statistics 33 (2), 350-374.

Hájek, J., 1964. Asymptotic Theory of Rejective Sampling with Varying Probabilities from a Finite Population. The Annals of Mathematical Statistics 35 (4), 1491-1523.

Tillé, Y., 1996. Some Remarks on Unequal Probability Sampling Designs Without Replacement. Annals of Economics and Statistics 44, 177-189.

Brewer, K.R.W.; Donadio, M.E., 2003. The High Entropy Variance of the Horvitz-Thompson Estimator. Survey Methodology 29 (2), 189-196.

jip_MonteCarlo

Examples

```
### Generate population data ---
N <- 20; n<-5
set.seed(0)
x <- rgamma(N, scale=10, shape=5)
y <- abs( 2*x + 3.7*sqrt(x) * rnorm(N) )
pik <- n * x/sum(x)
### Approximate joint-inclusion probabilities ---
pikl <- jip_approx(pik, method='Hajek')
pikl <- jip_approx(pik, method='HartleyRao')
pikl <- jip_approx(pik, method='HartleyRao')
pikl <- jip_approx(pik, method='Tille')
pikl <- jip_approx(pik, method='Brewer1')
pikl <- jip_approx(pik, method='Brewer2')
pikl <- jip_approx(pik, method='Brewer3')
pikl <- jip_approx(pik, method='Brewer4')</pre>
```

jip_MonteCarlo

```
Approximate inclusion probabilities by Monte Carlo simulation
```

Description

Approximate first and second-order inclusion probabilities by means of Monte Carlo simulation. Estimates are obtained as proportion of the number of occurrences of each unit or couple of units over the total number of replications. One unit is added to both numerator and denominator to assure strict positivity of estimates (Fattorini, 2006).

Usage

```
jip_MonteCarlo(
    x,
    n,
    replications = 1e+06,
    design,
    units,
    seed = NULL,
    as_data_frame = FALSE,
    design_pars,
    write_on_file = FALSE,
    filename,
    path,
    by = NULL,
```

```
progress_bar = TRUE
)
```

Arguments

x	size measure or first-order inclusion probabilities, a vector or single-column data.frame
n	sample size (for fixed-size designs), or expected sample size (for Poisson sampling)
replications	numeric value, number of independent Monte Carlo replications
design	sampling procedure to be used for sample selection. Either a string indicating the name of the sampling design or a function; see section "Details" for more information.
units	indices of units for which probabilities have to be estimated. Optional, if miss- ing, estimates are produced for the whole population
seed	a valid seed value for reproducibility
as_data_frame	logical, should output be in a data.frame form? if FALSE, a matrix is returned
design_pars	only used when a function is passed to argument design, named list of parame- ters to pass to the sampling design function.
write_on_file	logical, should output be written on a text file?
filename	string indicating the name of the file to create on disk, must include the .txt extension; only applies if write_on_file = TRUE.
path	string indicating the path to the directory where the output file should be created; only applies if write_on_file = TRUE.
by	optional; integer scalar indicating every how many replications a partial output should be saved
progress_bar	logical, indicating whether a progress bar is desired

Details

Argument design accepts either a string indicating the sampling design to use to draw samples or a function. Accepted designs are "brewer", "tille", "maxEntropy", "poisson", "sampford", "systematic", "randomSystematic". The user may also pass a function as argument; such function should take as input the parameters passed to argument design_pars and return either a logical vector or a vector of 0s and 1s, where TRUE or 1 indicate sampled units and FALSE or 0 indicate non-sample units. The length of such vector must be equal to the length of x if units is not specified, otherwise it must have the same length of units.

When write_on_file = TRUE, specifying a value for aurgument by will produce intermediate files with approximate inclusion probabilities every by number of replications. E.g., if replications=1e06 and by=5e05, two output files will be created: one with estimates at 5e05 and one at 1e06 replications. This option is particularly useful to assess convergence of the estimates.

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Value

A matrix of estimated inclusion probabilities if as_data_frame=FALSE, otherwise a data.frame with three columns: the first two indicate the ids of the the couple of units, while the third one contains the joint-inclusion probability values. Please, note that when as_data_frame=TRUE, first-order inclusion probabilities are not returned.

References

Fattorini, L. 2006. Applying the Horvitz-Thompson criterion in complex designs: A computerintensive perspective for estimating inclusion probabilities. Biometrika 93 (2), 269–278

Examples

```
### Generate population data ---
N <- 20; n<-5
set.seed(0)
x <- rgamma(N, scale=10, shape=5)</pre>
y <- abs( 2*x + 3.7*sqrt(x) * rnorm(N) )</pre>
pik <- n * x/sum(x)
### Approximate joint-inclusion probabilities
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "brewer")</pre>
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "tille")</pre>
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "maxEntropy")</pre>
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "randomSystematic")</pre>
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "systematic")</pre>
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "sampford")</pre>
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "poisson")</pre>
#Use an external function to draw samples
pikl <- jip_MonteCarlo(x=pik, n=n, replications=100,</pre>
                        design = sampling::UPmidzuno, design_pars = list(pik=pik))
#Write output on file after 50 and 100 replications
pikl <- jip_MonteCarlo(x=pik, n = n, replications = 100, design = "brewer",</pre>
                      write_on_file = TRUE, filename="test.txt", path=tempdir(), by = 50 )
```

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