

Package ‘NestedCategBayesImpute’

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Type Package

Title Modeling and Generating Synthetic Versions of Nested Categorical Data in the Presence of Impossible Combinations

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Description This tool set provides a set of functions to fit the nested Dirichlet process mixture of products of multinomial distributions (NDPMPM) model for nested categorical household data in the presence of impossible combinations. It has direct applications in generating synthetic nested household data.

License GPL(>=3)

LazyData TRUE

Imports stats, coda, dplyr, Rcpp (>= 0.12.0)

LinkingTo Rcpp

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R topics documented:

checkconstraints	2
checkconstraints_HHhead_at_group_level	3
GetImpossibleHouseholds	4
groupcount	5
groupcount1D	6
households2individuals	7
initData	7
initOutput	8
initParameters	9
RunModel	10
sampleG	13
samplehouseholds	14
samplehouseholds_HHhead_at_group_level	15
sampleM	16
UpdateAlpha	17
UpdateBeta	17
UpdateLambda	18

UpdateLambdaWeighted	19
UpdateOmega	19
UpdateOmegaWeighted	20
UpdatePhi	21
UpdatePhiWeighted	22
UpdatePi	23
UpdatePiWeighted	23

Index	25
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checkconstraints	<i>Checking a data matrix of households for the possible/impossible status under a predefined set of structural zeros.</i>
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Description

Checking a data matrix of households for the possible/impossible status under a predefined set of structural zeros.

Usage

```
checkconstraints(data, neededpossiblehh, hh_size)
```

Arguments

data	A household data matrix generated by calling <code>samplinghouseholds</code> .
neededpossiblehh	The number of possible households needed before checking is stopped.
hh_size	The household size for the households in data.

Details

Given an input household data matrix, these functions will check the possible/impossible status of each household and also output the desired number of possible and impossible households separately. `checkconstraints` checks constraints when the household head is included as an individual within the household.

The predefined list of structural zeros currently included is completely data driven. It does not fully represent the true list of impossible combinations in the true population from the American Community Survey (ACS) but rather represents a primary list of combinations not satisfied by our samples of sizes 10000 from the 2012 ACS. In fact, some of the structural zeros included here do not generalize to more recent ACS data sets. For example, we expect the true population to contain households with same sex couple but do not observe any in our samples and so we leave this in to be able to generate synthetic data with the properties of our sample data sets.

The structural zeros included are:

- Valid Head: Each household must contain exactly one head and he/she must be at least 16 years old.
- Valid Spouse: Each household cannot contain more than one spouse and he/she must be at least 16 years old.
- Valid Couple: No same sex couple and age difference between couples cannot exceed 49.
- Valid Children: The household head must be older than the oldest child by at least 7.

- Valid Parents: The youngest parent must be older than the household head by at least 10.
- Valid Parents-in-Law: The youngest parent-in-law must be older than the household head by at least 4.
- Valid Siblings: The age difference between the household head and siblings cannot exceed 37.
- Valid Grandchildren: The household head must be at least 34 years old. Also, the household head must be older than the oldest grandchild by at least 26.

Users can modify this list by downloading the package source, making changes only to the checkconstraints_imp.cpp file and re-building the package.

Value

A list containing information on checking result.

outcome	An indicator vector for the possible/impossible household status under constraints.
Households	A data matrix for impossible households.
Index	A vector for the original indexes of households when possible households are found. Generally not to be used.
synHouseholds	A data matrix for possible households.
possible	The actual number of possible households returned.

Author(s)

Quanli Wang, Olanrewaju Akande

checkconstraints_HHhead_at_group_level

Checking a data matrix of households for the possible/impossible status under a predefined set of structural zeros.

Description

Checking a data matrix of households for the possible/impossible status under a predefined set of structural zeros.

Usage

```
checkconstraints_HHhead_at_group_level(data, neededpossiblehh, hh_size)
```

Arguments

data	A household data matrix generated by calling samplinghouseholds.
neededpossiblehh	The number of possible households needed before checking is stopped.
hh_size	The household size for the households in data.

Details

Given an input household data matrix, these functions will check the possible/impossible status of each household and also output the desired number of possible and impossible households separately. `checkconstraints_HHhead_at_group_level` checks constraints when the household head is moved to the household level. For the list of structural zeros currently included, see the documentation for `checkconstraints`.

Value

A list containing information on checking result.

<code>outcome</code>	An indicator vector for the possible/impossible household status under constraints.
<code>Households</code>	A data matrix for impossible households.
<code>Index</code>	A vector for the original indexes of households when possible households are found. Generally not to be used.
<code>synHouseholds</code>	A data matrix for possible households.
<code>possible</code>	The actual number of possible households returned.

Author(s)

Quanli Wang, Olanrewaju Akande

`GetImpossibleHouseholds`

Generate the desired number of impossible households required to observe a given number of possible households.

Description

Given model parameters, generate the desired number of impossible households required to observe a given number of possible households. Also generate synthetic (and valid) data of the same size as the observed data when required.

Usage

```
GetImpossibleHouseholds(d, n_star_h, lambda, omega, phi, pi, howmany, n, synindex,
                        HHhead_at_group_level)
```

Arguments

<code>d</code>	Vector containing the number of levels for each individual-level variable.
<code>n_star_h</code>	Vector containing the number of observed households for the different household sizes in the original data.
<code>lambda</code>	Multinomial probabilities for each group-level variable.
<code>omega</code>	Latent class probabilities for the group-level and individual-level latent class pairs.
<code>phi</code>	Multinomial probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.

pi	Latent class probabilities for the group-level latent classes.
howmany	Number of households to be generated at a time; batch sampling is used to improve computing speed.
n	Number of households in the original input data and the sum of n_star_h.
synindex	Logical indicator for sampling synthetic data. Set to TRUE when synthetic data is needed.
HHhead_at_group_level	Logical indicator for data structure with respect to the household head. Set to TRUE if the household head has been moved to the household level and FALSE otherwise.

Value

G_Individuals_and_M_extra	A data matrix containing both the group-level (in long format) and individual-level latent classes for the impossible households.
G_extra	A vector containing the group-level latent classes for the impossible households.
IndividualData_extra	A data matrix containing the individual-level data for the impossible households.
HHdata_extra	A data matrix containing the group-level data for the impossible households.
hh_size_new	A vector for the number of impossible households for the different household sizes.
synIndividuals_all	Synthetic data when synindex is TRUE. NULL otherwise.

Author(s)

Quanli Wang

groupcount	<i>Generate 2D count table for two integer-valued vectors.</i>
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Description

Similar to 'table' function, this function builds a contingency table of the counts at each combination of all possible values from two integer-valued input vectors.

Usage

```
groupcount(g1, g2, n1, n2)
```

Arguments

g1	The first integer-valued input vector. The max value in g1 is n1.
g2	The second integer-valued input vector. The max value in g1 is n2.
n1	The maximum value in g1.
n2	The maximum value in g2.

Details

This is implemented as an utility function to build a 2D histogram count table. For efficiency, it does not check if the maximum values in input vectors exceed the maximum values specified.

Value

The count table.

Author(s)

Quanli Wang

Examples

```
n1 <- 20
n2 <- 10
g1 <- sample.int(n1,1000, replace = TRUE)
g2 <- sample.int(n2,1000, replace = TRUE)
counts <- groupcount(g1,g2,n1,n2)
```

groupcount1D

Generate histogram count for an integer-valued vector.

Description

Generate histogram count for an integer-valued vector.

Usage

```
groupcount1D(g, n)
```

Arguments

g An integer-valued input vector. The max value in g is n.
n The max value in g.

Details

This is implemented as an utility function for 1D histogram count. For efficiency, it does not check if the maximum value in the input vector exceeds the maximum value specified.

Value

The count values.

Author(s)

Quanli Wang

Examples

```
n <- 20
g <- sample.int(n,1000, replace = TRUE)
counts <- groupcount1D(g,n)
```

`households2individuals`

Convert a household data matrix to the corresponding individual member data matrix.

Description

Convert a household data matrix to the corresponding individual member data matrix.

Usage

```
households2individuals(data, hh_size)
```

Arguments

<code>data</code>	Household data matrix.
<code>hh_size</code>	The household size for the households in data.

Value

Individual member data matrix.

Author(s)

Quanli Wang

`initData`

Initialize the input data structure.

Description

Initialize the input data structure.

Usage

```
initData(household, individual_variable_index, household_variable_index)
```

Arguments

<code>household</code>	A data matrix for the input household data.
<code>individual_variable_index</code>	Vector of column indexes for the individual-level variables.
<code>household_variable_index</code>	Vector of column indexes for the group-level variables.

Value

A list object including all the necessary data variables needed by the sampler.

origdata	Original data.
n_i	Vector containing the number of individuals in each household in the data.
n	Number of households in the data
HHdataorigT	The transposed household level data – each column now represents each household.
HHserial	Vector containing the household index for each individual in the data.
n_individuals	The total number of individuals N across all n households in the input data.
n_individuals_real	The real total number of individuals N across all n households. The is the same as n_individuals if the household head hasn't been moved to the household level and different otherwise.
p	Number of individual-level variables.
d	Vector containing the number of levels for each of the p variables.
dataT	The transposed individual level data – each column now represents each individual.
maxd	The max value in d
n_star_h	Vector containing the number of observed households for the different household sizes in the original data.

Author(s)

Quanli Wang

initOutput	<i>Set the output structure for saving posterior samples of parameters.</i>
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Description

Set the output structure for saving posterior samples of parameters.

Usage

```
initOutput(data, hyper, mc)
```

Arguments

data	A list object including all the necessary data variables needed by the sampler.; output of the initData function.
hyper	Hyper parameters for priors.
mc	MCMC parameters.

Value

A list of output parameters to be saved.

alphaout	Vector of posterior samples for the concentration parameter in the Dirichlet process for the group-level latent classes.
betaout	Vector of posterior samples for the concentration parameter in the Dirichlet process for the individual-level latent classes. Currently, this is assumed to be the same within all group-level classes.
piout	Matrix of posterior samples for the vector of probabilities for the group-level latent classes.
omegaout	3D array of posterior samples for the matrix of probabilities for the group-level and individual-level latent class pairs.
nout	Vector of posterior samples for the total number of impossible households sampled.
extrasize	Matrix of posterior samples for the number of impossible households sampled, split by household size.
F_occupied	Vector of posterior samples for the number of occupied household-level latent classes.
S_occupied_max	Vector of posterior samples for the max number of occupied individual-level latent classes.
elapsed_time	Vector of time taken to run each iteration.
newphiout	3D array of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.
lambdaout	A list of an array of posterior samples for the group-level probabilities for each group-level variable. Each array in the list is for each group-level variable.

Author(s)

Quanli Wang, Olanrewaju Akande

initParameters	<i>Initialize the model parameters for the MCMC.</i>
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Description

Initialize the model parameters for the MCMC.

Usage

```
initParameters(data, hyper, HHhead_at_group_level)
```

Arguments

data	A list object including all the necessary data variables needed by the sampler; output of the initData function.
hyper	Hyper parameters for the prior distributions.
HHhead_at_group_level	Logical indicator for data structure with respect to the household head. Set to TRUE if the household head has been moved to the household level and FALSE otherwise.

Value

A list of the initial values of the parameters.

alpha	Concentration parameter in the Dirichlet process for the group-level latent classes.
beta	Concentration parameter in the Dirichlet process for the individual-level latent classes. Currently, this is assumed to be the same within all group-level classes.
phi	Matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.
HHdata_all	The transposed household level data – each column represents each household.
lambda	A list of matrices of the group-level probabilities for each group-level variable by the group-level latent classes. Each matrix in the list is for each group-level variable.
u	Vector of the beta-distributed variables in the stick breaking representation of the group-level latent classes.
pi	Vector of the probabilities for the group-level latent classes.
v	Matrix of the beta-distributed variables in the stick breaking representation of the individual-level latent classes by the group-level latent classes.
omega	Matrix of the probabilities for the individual-level latent classes by the group-level latent classes.

Author(s)

Quanli Wang

RunModel

Run the mcmc sampler for the model.

Description

Run the mcmc sampler for the model.

Usage

```
RunModel(orig,mc,hyper,para,output,synindex,individual_variable_index,
household_variable_index,HHhead_at_group_level,weight_option,struc_weight)
```

Arguments

orig	A list object including all the necessary data variables needed by the sampler; output of the initData function.
mc	A list specifying the number of mcmc iterations, burn-in, thinning and the effective sample size.
hyper	Hyper parameters for the prior distributions.
para	A list of the initial values of the parameters; output of the initParameters function.
output	A list of output parameters to be saved; output of the initOutput function.

synindex	A vector of iteration indexes for sampling synthetic data. <code>length(synindex)</code> is the number of synthetic data needed.
individual_variable_index	Vector of column indexes for the individual-level variables.
household_variable_index	Vector of column indexes for the group-level variables.
HHhead_at_group_level	Logical indicator for whether or not to move the household head to the household level. Set to <code>TRUE</code> to move the household head and <code>FALSE</code> otherwise.
weight_option	Logical indicator for whether or not to cap the number of impossible households to sample and re-weight the multinomial counts within each latent class back to the expected truth. Set to <code>TRUE</code> to use the weighting option and <code>FALSE</code> otherwise.
struc_weight	Vector specifying the weights to be used for each household size. The weights must be ordered by household sizes and no household must be excluded.

Details

This function runs the mcmc sampler for the NDPMPM model and generates posterior samples of parameters. It also generates synthetic data when needed.

Please note that:

- The minimum household size for this mcmc sampler is 2 because households of size 1 do not violate the structural zeros specified in this package. Also, moving the household head to the household level is not possible for households of size 1.
- Each variable included must be recoded to start from 1.
- Moving the household head to the household level and setting the `HHhead_at_group_level` option to `TRUE` speeds up the sampler significantly.
- Setting the `weight_option` to `TRUE` and specifying weights also speeds up the sampler but the exact rate of speedup depends on the specific weights.

Value

synData	The list of synthetic data when the <code>length(synindex) > 0</code> .
output	The list of posterior samples for the parameters included in output.

Author(s)

Quanli Wang, Olanrewaju Akande

Examples

```
### We now show how the sampler works using a sample from the 2012 ACS.
### The sample consists of 2000 households of sizes 2 to 6.
### Empty environment and load required libraries
rm(list = ls())
library(NestedCategBayesImpute)
library(dplyr)

### Set indicator for whether of not to move the household head
### Also set indicator for the weighting/capping option
```

```

HHhead_at_group_level <- TRUE #move head to the group level
weight_option <- TRUE #use weighting/capping option

### Use data included in package; head has been moved to the group level
### prepare data and specify variable indexes
orig.file <- system.file("extdata","origdata_newFormat.txt",
                        package="NestedCategBayesImpute")
orig.data <- read.table(orig.file,header = TRUE, sep = " ")
orig.data$relate <- orig.data$relate - 1L #recode relate to 11 levels
household.size <- as.data.frame(table(orig.data$Hhindex))
household.size[,1] <- as.numeric(household.size[,1])
names(household.size) <- c("Hhindex", 'householdsize')
household <- orig.data %>% inner_join(household.size)
individual_variable_index = c(3:7)
household_variable_index = c(8:13) #last column must be household size

### Initialize the input data structure
orig <- initData(household,individual_variable_index,household_variable_index)

### Check first few lines of data; data contains households of sizes 2 to 6
head(household)

### Supply weights; one for each household size
### Also,must be ordered & no household size must be excluded
if(weight_option){
  struc_weight <- c(1/2,1/2,1/3,1/3,1/3)
} else {
  struc_weight <-rep(1,length(orig$n_star_h)) #just a dummy vector of ones
}

### Set mcmc parameters
mc <- list(nrun = 10, burn = 5, thin = 1)
mc$eff.sam <- (mc$nrun-mc$burn)/mc$thin

### Set number of categories for each household level variable
dHH <- rep(0,length(household_variable_index))
for (i in 1:length(dHH)) {
  dHH[i] <- max(household[,household_variable_index[i]])
  if (i == length(dHH) & !HHhead_at_group_level) {
    #When leaving head within household, model assumes that the HH size starts from 2
    dHH[length(dHH)] <- dHH[length(dHH)] - 1
  }
}

### Set hyper parameters
#aa, ab, ba & bb are gamma hyperparameters for alpha and beta
#blocksize is the number of households to sample at once (batch sampling)
#FF is the max number of group-level latent classes
#SS is the max number of individual-level classes
hyper <- list(FF=20 , SS=15, aa=0.25, ab=0.25,

```

```

ba=0.25,bb=0.25,dHH = dHH, blocksize = 5000)

### Initialize parameters and output
para <- initParameters(orig,hyper,HHhead_at_group_level)
output <- initOutput(orig,hyper,mc)

### Set number of synthetic data and the mcmc indexes for them
mm <- 5
synindex <- sort(sample(seq((mc$burn +1),mc$nrun,by=mc$thin),mm,replace=FALSE))

### Run model
ModelResults <- RunModel(orig,mc,hyper,para,output,synindex,
                          individual_variable_index,household_variable_index,
                          HHhead_at_group_level,weight_option,struc_weight)

### View first few lines of the first synthetic data.
head((ModelResults$synData)[[1]])

### Some posterior summaries and plots
library(coda)
names(ModelResults$output)
dim(ModelResults$output$alphaout)
alpha_output <- mcmc(ModelResults$output$alphaout)
plot(alpha_output)
summary(alpha_output)

dim(ModelResults$output$betaout)
beta_output <- mcmc(ModelResults$output$betaout)
plot(beta_output)
summary(beta_output)

```

sampleG

*Update household (group) level latent class indexes.***Description**

Update household (group) level latent class indexes.

Usage

```
sampleG(phi, data, omega, pi, ni, HHdata, lambda)
```

Arguments

phi	Matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.
data	Individual level data.

omega	Matrix of the probabilities for the individual-level latent classes by the group-level latent classes.
pi	Vector of the probabilities for the group-level latent classes.
ni	Vector containing the number of individuals in each household in the data..
HHdata	Household level data.
lambda	A list of matrices of the group-level probabilities for each group-level variable by the group-level latent classes. Each matrix in the list is for each group-level variable.

Details

Function for obtaining a posterior sample of the household-level latent class indexes for all households in the input data based on the corresponding full conditional distribution.

Value

A list with two variables.

G	A vector for the updated values of the household-level latent class indexes for all households in the input data.
G_Individuals	The vector G expanded to a long format to match the number of individuals in data.

Author(s)

Quanli Wang

samplehouseholds	<i>Rcpp implementation for sampling household data without constraints.</i>
------------------	---

Description

Rcpp implementation for sampling household data without constraints.

Usage

```
samplehouseholds(phi, omega, pi, d, lambda, currrentbatch, nHouseholds, householdsize)
```

Arguments

phi	Matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.
omega	Matrix of the probabilities for the individual-level latent classes by the group-level latent classes.
pi	Vector of the probabilities for the group-level latent classes.
d	Vector containing the number of levels for each of the individual-level variables.
lambda	A list of matrices of the group-level probabilities for each group-level variable by the group-level latent classes. Each matrix in the list is for each group-level variable.

currentbatch	The current batch number for the household data to be generated. The household ID will be generated based on this batch number.
nHouseholds	The number of households to be generated by one call to this function.
householdsize	The size of the households to be generated.

Details

This function allows the model to generate a batch of nHouseholds with each household of size householdsize. The generated household data will include both possible and impossible households. Use samplehouseholds when the household head is included as an individual within the household.

Value

A data matrix with each row for one household.

Author(s)

Quanli Wang

samplehouseholds_HHhead_at_group_level

Rcpp implementation for sampling household data without constraints.

Description

Rcpp implementation for sampling household data without constraints.

Usage

```
samplehouseholds_HHhead_at_group_level(phi, omega, pi, d, lambda, currentbatch,
                                         nHouseholds, householdsize)
```

Arguments

phi	Matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.
omega	Matrix of the probabilities for the individual-level latent classes by the group-level latent classes.
pi	Vector of the probabilities for the group-level latent classes.
d	Vector containing the number of levels for each of the individual-level variables.
lambda	A list of matrices of the group-level probabilities for each group-level variable by the group-level latent classes. Each matrix in the list is for each group-level variable.
currentbatch	The current batch number for the household data to be generated. The household ID will be generated based on this batch number.
nHouseholds	The number of households to be generated by one call to this function.
householdsize	The size of the households to be generated.

Details

This function allows the model to generate a batch of nHouseholds with each household of size householdsize. The generated household data will include both possible and impossible households. Use samplehouseholds_HHhead_at_group_level when the household head is moved to the household level.

Value

A data matrix with each row for one household.

Author(s)

Quanli Wang

sampleM	<i>Update individual level latent class indexes.</i>
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Description

Update individual level latent class indexes.

Usage

```
sampleM(phi, data, omega, G, serial)
```

Arguments

phi	Matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes.
data	Input individual-level data.
omega	Matrix of the probabilities for the individual-level latent classes by the group-level latent classes.
G	Household-level latent class indexes.
serial	Vector containing the household index for each individual in the data.

Details

Function for obtaining a posterior sample of the individual-level latent class indexes for all individuals in the input data based on the corresponding full conditional distribution.

Value

A vector for the updated values of the individual-level latent class indexes for all individuals in the input data.

Author(s)

Quanli Wang

UpdateAlpha	<i>Update alpha.</i>
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Description

Update alpha – the concentration parameter in the Dirichlet process for the group-level latent classes.

Usage

```
UpdateAlpha(aa, ab, u)
```

Arguments

aa	Hyper-parameter a for alpha.
ab	Hyper-parameter b for alpha.
u	Vector of the beta-distributed variables in the stick breaking representation of the group-level latent classes.

Value

Updated (posterior) value for alpha based on the corresponding full conditional distribution.

Author(s)

Quanli Wang

UpdateBeta	<i>Update beta.</i>
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Description

Update beta – the concentration parameter in the Dirichlet process for the individual-level latent classes. Currently, this is assumed to be the same within all group-level classes.

Usage

```
UpdateBeta(ba, bb, v)
```

Arguments

ba	Hyper-parameter a for beta.
bb	Hyper-parameter b for beta.
v	Matrix of the beta-distributed variables in the stick breaking representation of the individual-level latent classes by the group-level latent classes.

Value

Updated (posterior) value for beta based on the corresponding full conditional distribution..

Author(s)

Quanli Wang

UpdateLambda

Update lambda.

Description

Update lambda – the list of matrices of the group-level probabilities for each group-level variable by the group-level latent classes when the weighting/capping option is not used. Each matrix in the list is for each group-level variable.

Usage

```
UpdateLambda(dHH, FF, G_all, HHdata_all)
```

Arguments

dHH	A vector containing the number of levels for each household-level variable.
FF	Maximum number of household-level latent classes allowed.
G_all	A vector of the household-level latent class indexes for all households both in the original data and the sampled impossible households.
HHdata_all	Data matrix for the household-level data from both the original data and the sampled impossible households.

Details

Function for obtaining a posterior sample of lambda when the weighting/capping option is not used.

Value

Updated (posterior) value for lambda based on the corresponding full conditional distribution.

Author(s)

Quanli Wang

UpdateLambdaWeighted *Update lambda.*

Description

Update lambda – the list of matrices of the group-level probabilities for each group-level variable by the group-level latent classes – when the weighting/capping option is used. The weighting options allows capping the number of impossible households to sample and re-weight the multinomial counts within each latent class back to the expected truth. Each matrix in the list is for each group-level variable.

Usage

```
UpdateLambdaWeighted(dHH, FF, G_all, HHdata_all, struc_weight)
```

Arguments

dHH	A vector containing the number of levels for each household-level variable.
FF	Maximum number of household-level latent classes allowed.
G_all	A vector of the household-level latent class indexes for all households both in the original data and the sampled impossible households.
HHdata_all	Data matrix for the household-level data from both the original data and the sampled impossible households.
struc_weight	A vector of weights by household sizes used in capping the number of sampled impossible households.

Details

Function for obtaining a posterior sample of lambda when the weighting/capping option is used.

Value

Updated (posterior) value for lambda based on the corresponding full conditional distribution.

Author(s)

Quanli Wang, Olanrewaju Akande

UpdateOmega *Update omega and v.*

Description

Update omega – the matrix of the probabilities for the individual-level latent classes by the group-level latent classes – and v – the matrix of the beta-distributed variables in the stick breaking representation of the individual-level latent classes by the group-level latent classes – when the weighting/capping option is not used.

Usage

```
UpdateOmega(beta, M_all, FF, SS)
```

Arguments

beta	Concentration parameter in the Dirichlet process for the individual-level latent classes. Currently, this is assumed to be the same within all group-level classes.
M_all	A vector of both the household-level and individual-level latent class indexes for all households both in the original data and the sampled impossible households.
FF	Maximum number of household-level latent classes allowed.
SS	Maximum number of individual-level latent classes allowed.

Value

A list containing the updated (posterior) values for omega and v based on the corresponding full conditional distributions.

Author(s)

Quanli Wang

UpdateOmegaWeighted	<i>Update omega and v.</i>
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Description

Update omega – the matrix of the probabilities for the individual-level latent classes by the group-level latent classes – and v – the matrix of the beta-distributed variables in the stick breaking representation of the individual-level latent classes by the group-level latent classes – when the weighting/capping option is used. The weighting options allows capping the number of impossible households to sample and re-weight the multinomial counts within each latent class back to the expected truth.

Usage

```
UpdateOmegaWeighted(beta, M_all, FF, SS, struc_weight)
```

Arguments

beta	Concentration parameter in the Dirichlet process for the individual-level latent classes. Currently, this is assumed to be the same within all group-level classes.
M_all	A vector of both the household-level and individual-level latent class indexes for all households both in the original data and the sampled impossible households.
FF	Maximum number of household-level latent classes allowed.
SS	Maximum number of individual-level latent classes allowed.
struc_weight	A vector of weights by household sizes used in capping the number of sampled impossible households.

Value

A list containing the updated (posterior) values for omega and v based on the corresponding full conditional distributions.

Author(s)

Quanli Wang, Olanrewaju Akande

UpdatePhi	<i>Update phi.</i>
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Description

Update phi – the matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes – when the weighting/capping option is not used.

Usage

```
UpdatePhi(IndividualData_all, M_all, FF, SS, p, d, maxd, individual_variable_index)
```

Arguments

IndividualData_all	Data matrix for the individual-level data from both the original data and the sampled impossible households.
M_all	A vector of both the household-level and individual-level latent class indexes for all households both in the original data and the sampled impossible households.
FF	Maximum number of household-level latent classes allowed.
SS	Maximum number of individual-level latent classes allowed.
p	Number of individual-level variables.
d	A vector for the number of levels of each individual-level variable.
maxd	Maximum value in d.
individual_variable_index	Vector of column indexes for the individual-level variables.

Details

Function for obtaining a posterior sample of phi when the weighting/capping option is not used.

Value

Updated (posterior) value for phi based on the corresponding full conditional distribution.

Author(s)

Quanli Wang

UpdatePhiWeighted	<i>Update phi.</i>
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Description

Update phi – the matrix of posterior samples for the individual-level probabilities for each individual-level variable by each pair of group-level and individual-level latent classes – when the weighting/capping option is used. The weighting options allows capping the number of impossible households to sample and re-weight the multinomial counts within each latent class back to the expected truth.

Usage

```
UpdatePhiWeighted(IndividualData_all, M_all, FF, SS, p, d, maxd,
                  individual_variable_index, struc_weight)
```

Arguments

IndividualData_all	Data matrix for the individual-level data from both the original data and the sampled impossible households.
M_all	A vector of both the household-level and individual-level latent class indexes for all households both in the original data and the sampled impossible households.
FF	Maximum number of household-level latent classes allowed.
SS	Maximum number of individual-level latent classes allowed.
p	Number of individual-level variables.
d	A vector for the number of levels of each individual-level variable.
maxd	Maximum value in d.
individual_variable_index	Vector of column indexes for the individual-level variables.
struc_weight	A vector of weights by household sizes used in capping the number of sampled impossible households.

Details

Function for obtaining a posterior sample of phi when the weighting/capping option is used.

Value

Updated (posterior) value for phi based on the corresponding full conditional distribution.

Author(s)

Quanli Wang, Olanrewaju Akande

UpdatePi	<i>Update pi and u.</i>
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Description

Update pi – the vector of the probabilities for the group-level latent classes – and u – the vector of the beta-distributed variables in the stick breaking representation of the group-level latent classes – when the weighting/capping option is not used.

Usage

```
UpdatePi(alpha, G_all, FF)
```

Arguments

alpha	Concentration parameter in the Dirichlet process for the group-level latent classes
G_all	A vector of the household-level latent class indexes for all households both in the original data and the sampled impossible households.
FF	Maximum number of household-level latent classes allowed.

Details

Function for obtaining a posterior sample of pi when the weighting/capping option is not used.

Value

A list containing the updated (posterior) values for pi and u based on the corresponding full conditional distributions.

Author(s)

Quanli wang

UpdatePiWeighted	<i>Update pi and u.</i>
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Description

Update pi – the vector of the probabilities for the group-level latent classes – and u – the vector of the beta-distributed variables in the stick breaking representation of the group-level latent classes when the weighting/capping option is used. The weighting options allows capping the number of impossible households to sample and re-weight the multinomial counts within each latent class back to the expected truth.

Usage

```
UpdatePiWeighted(alpha, G_all, FF, struc_weight)
```

Arguments

alpha	Concentration parameter in the Dirichlet process for the group-level latent classes
G_all	A vector of the household-level latent class indexes for all households both in the original data and the sampled impossible households.
FF	Maximum number of household-level latent classes allowed.
struc_weight	A vector of weights by household sizes used in capping the number of sampled impossible households.

Details

Function for obtaining a posterior sample of π when the weighting/capping option is used.

Value

A list containing the updated (posterior) values for π and u based on the corresponding full conditional distributions.

Author(s)

Quanli wang, Olanrewaju Akande

Index

*Topic **constraints**

checkconstraints, [2](#)
checkconstraints_HHhead_at_group_level,
[3](#)

*Topic **household data without constraint**

samplehouseholds, [14](#)
samplehouseholds_HHhead_at_group_level,
[15](#)

*Topic **household level**

sampleG, [13](#)

*Topic **impossible household**

checkconstraints, [2](#)
checkconstraints_HHhead_at_group_level,
[3](#)
GetImpossibleHouseholds, [4](#)

*Topic **individual level**

sampleM, [16](#)

*Topic **mcmc**

RunModel, [10](#)

*Topic **model**

RunModel, [10](#)

*Topic **possible household**

checkconstraints, [2](#)
checkconstraints_HHhead_at_group_level,
[3](#)

*Topic **sampler**

GetImpossibleHouseholds, [4](#)
sampleG, [13](#)
samplehouseholds, [14](#)
samplehouseholds_HHhead_at_group_level,
[15](#)
sampleM, [16](#)
UpdateAlpha, [17](#)
UpdateBeta, [17](#)
UpdateLambda, [18](#)
UpdateLambdaWeighted, [19](#)
UpdateOmega, [19](#)
UpdateOmegaWeighted, [20](#)
UpdatePhi, [21](#)
UpdatePhiWeighted, [22](#)
UpdatePi, [23](#)
UpdatePiWeighted, [23](#)

*Topic **synthetic data**

GetImpossibleHouseholds, [4](#)

*Topic **utility function**

households2individuals, [7](#)

checkconstraints, [2](#)
checkconstraints_HHhead_at_group_level,
[3](#)

GetImpossibleHouseholds, [4](#)
groupcount, [5](#)
groupcount1D, [6](#)

households2individuals, [7](#)

initData, [7](#)
initOutput, [8](#)
initParameters, [9](#)

RunModel, [10](#)

sampleG, [13](#)
samplehouseholds, [14](#)
samplehouseholds_HHhead_at_group_level,
[15](#)
sampleM, [16](#)

UpdateAlpha, [17](#)
UpdateBeta, [17](#)
UpdateLambda, [18](#)
UpdateLambdaWeighted, [19](#)
UpdateOmega, [19](#)
UpdateOmegaWeighted, [20](#)
UpdatePhi, [21](#)
UpdatePhiWeighted, [22](#)
UpdatePi, [23](#)
UpdatePiWeighted, [23](#)