

Package ‘sourceR’

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Type Package

Title Fits a Non-Parametric Bayesian Source Attribution Model

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Description Implements a non-parametric source attribution model to attribute cases of disease to sources in Bayesian framework with source and type effects. Type effects are clustered using a Dirichlet Process. Multiple times and locations are supported.

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Depends R (>= 3.4.0), dplyr, tensorA, assertthat

Imports gtools, hashmap, R6, cluster, stats, gplots, SPIn, grDevices, reshape2

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KeepSource TRUE

LazyData TRUE

Suggests testthat, R.rsp

VignetteBuilder R.rsp

NeedsCompilation no

Author Poppy Miller [aut, cre, cph],
Chris Jewell [aut],
Jonathan Marshall [ctb],
Nigel French [ctb]

Maintainer Poppy Miller <p.miller@lancaster.ac.uk>

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R topics documented:

AdaptiveDirMRW	2
AdaptiveLogDirMRW	3

AdaptiveMultiMRW	4
Alpha	5
Alpha_	6
campy	6
DataNode	7
DirichletNode	8
DirichletProcessNode	8
DPModel_impl	9
FormulaNode	10
GammaNode	11
HaldDP	11
Node	17
PoisGammaDPUpdate	18
PoissonNode	19
Prev	20
Q	20
sim_SA	21
sliceTensor	22
sourceR	22
StochasticNode	23
X	23
Y	24

Index	25
--------------	-----------

AdaptiveDirMRW	<i>AdaptiveDirMRW</i>
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Description

This class implements an Adaptive Multi-site Metropolis random walk algorithm, constrained so the parameter vector sums to 1.

Usage

AdaptiveDirMRW

Format

Object of [R6Class](#) with methods for updating a [DirichletNode](#) instance.

Details

An adaptive multivariate Gaussian proposal is used for $d-1$ elements of a d -dimensional parameter vector contained in node, with the d th element updated to ensure that the vector sums to 1. This makes the updater useful for Dirichlet distributed random variables.

For details of the adaptive scheme, see Roberts and Rosenthal (2012) Examples of Adaptive MCMC. *Journal of Computational and Graphical Statistics*. **18**:349–367.

Please note that no checks are performed as to the suitability of this algorithm for a particular [StochasticNode](#). It is up to the user to use the correct update algorithm for the appropriate nodes.

Value

Object of [AdaptiveDirMRW](#)

Fields

cov the current covariance
 burnin the number of updates to burn in
 tune the current tuning matrix
 naccept the number of accepted proposals
 ncalls the number of times update has been called
 node the node to which the updater is attached

Methods

`new(node, toupdate = function() 1:length(node$getData()), tune = rep(0.1, length(node$getData()))`, constructor takes an instance of a [StochasticNode](#) node, function to choose the indices of the elements to update (by default all elements), initial tuning vector (diagonal of adaptive tuning matrix), and the number of calls between adaptations.
`update()` when called, updates node
`acceptance()` return the acceptance rate

<code>AdaptiveLogDirMRW</code>	<i>AdaptiveLogDirMRW</i>
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Description

This class implements an Adaptive Multi-site logarithmic Metropolis-Hastings random walk algorithm, constrained so the parameter vector sums to 1.

Usage

`AdaptiveLogDirMRW`

Format

Object of [R6Class](#) with methods for updating a [DirichletNode](#) instance.

Details

An adaptive multivariate log-Gaussian proposal is used for $d-1$ elements of a d -dimensional parameter vector contained in node, with the d th element updated to ensure that the vector sums to 1. This makes the updater useful for Dirichlet distributed random variables, improving on [AdaptiveDirMRW](#) by ensuring proposals do not go negative.

For details of the adaptive scheme, see Roberts and Rosenthal (2012) Examples of Adaptive MCMC. *Journal of Computational and Graphical Statistics*. **18**:349–367.

Please note that no checks are performed as to the suitability of this algorithm for a particular [StochasticNode](#). It is up to the user to use the correct update algorithm for the appropriate nodes.

Value

Object of [AdaptiveLogDirMRW](#)

Fields

cov the current covariance
 burnin the number of updates to burn in
 tune the current tuning matrix
 naccept the number of accepted proposals
 ncalls the number of times update has been called
 node the node to which the updater is attached

Methods

`new(node, touupdate = function() 1:length(node$getData()), tune = rep(0.1, length(node$getData()))`, constructor takes an instance of a [StochasticNode](#) node, function to choose the indices of the elements to update (by default all elements), initial tuning vector (diagonal of adaptive tuning matrix), and number of calls between adaptations.
`update()` when called, updates node
`acceptance()` return the acceptance rate

 AdaptiveMultiMRW

AdaptiveMultiMRW

Description

This class implements an Adaptive Multi-site Metropolis random walk algorithm.

Usage

AdaptiveMultiMRW

Format

Object of [R6Class](#) with methods for updating a [Node](#) instance.

Details

A multivariate Gaussian proposal is used, for which the proposal variance is a scaled version of the evolving empirical posterior covariance matrix. See Roberts and Rosenthal (2012) Examples of Adaptive MCMC. *Journal of Computational and Graphical Statistics*. **18**:349–367.

Please note that no checks are performed as to the suitability of this algorithm for a particular [StochasticNode](#). It is up to the user to use the correct update algorithm for the appropriate nodes.

Value

Object of [AdaptiveMultiMRW](#)

Fields

cov the current covariance
 burnin the number of updates to burn in
 tune the current tuning matrix
 naccept the number of accepted proposals
 ncalls the number of times update has been called
 node the node to which the updater is attached

Methods

new(node, tune = rep(0.1, length(node\$getData()))), burning = 100) constructor takes an instance of a [StochasticNode](#) node, initial tuning vector (diagonal of adaptive tuning matrix), and number of burnin calls.
 update() when called, updates node
 acceptance() return the acceptance rate

Alpha

Constructs alpha prior

Description

The Alpha constructure function returns an R6 Alpha_ class which feeds sanitised prior or initialisation values for alpha into the model.

Usage

```
Alpha(data, alpha, source, time = NULL, location = NULL)
```

Arguments

data	long-format data.frame containing Dirichlet prior hyperparameter, source, time, and location columns
alpha	name of hyperparameter column
source	name of source column
time	name of optional time column
location	name of optional location column

Value

An Alpha_ data structure for use in sourceR models

Alpha_	<i>Alpha prior hyperparameter class</i>
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Description

Alpha prior hyperparameter class

Usage

Alpha_

Format

An object of class R6ClassGenerator of length 24.

campy	<i>Human cases of campylobacteriosis and numbers of source samples positive for Campylobacter.</i>
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Description

A dataset containing the number of human cases of campylobacteriosis and numbers of source samples positive for *Campylobacter* for each bacterial subtype.

Usage

campy

Format

A list containing the human cases ('cases'), source samples ('sources'0 and, prevalences ('prev').

cases: data frame with 91 rows and 2 variables:

Human number of human cases of campylobacteriosis between 2005-2008 in the Manawatu region of New Zealand

Type MLST type id for the samples

sources: data frame with 690 rows and 3 variables

Count number of source samples positive for campylobacteriosis

Source Source id for the samples

Type MLST type id for the samples

prev: data frame with 6 rows and 4 variables

Value Prevalence value (number of positive samples divided by total number of samples)

Source Source id for the samples

n_positive number of positive source samples for campylobacter (PCR)

n_total total number of source samples tested for campylobacter

DataNode

DataNode

Description

Represents a static data node in a DAG.

Usage

DataNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [Node](#), please see base class documentation.

Value

Object of [DataNode](#)

Fields

data the data

Methods

`getData()` returns the node's data.

DirichletNode *DirichletNode*

Description

Represents a d-dimensional Dirichlet distribution node in a DAG.

Usage

DirichletNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [StochasticNode](#), please see base class documentation.

Value

Object of [DirichletNode](#)

Methods

`new(data, alpha)` Create a DirichletNode with data vector data (length > 1) and parameter vector alpha.

DirichletProcessNode *DirichletProcessNode*

Description

Represents a Dirichlet process as a single node in a DAG.

Usage

DirichletProcessNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [StochasticNode](#), please see base class documentation.

Value

Object of [DirichletProcessNode](#)

Methods

`new(theta, s, alpha, base, ...)` Create a [DirichletProcessNode](#) with value vector `theta` (length > 1), initial grouping vector `s`, concentration parameter `alpha`, and base distribution `base`. `Base` should be a distribution function (`dnorm`, `dgamma`, etc) whose parameters are specified in ...

DPMoDel_impl

Builds the source attribution model. Is not intended to be used by a regular user. Developers only here!

Description

Builds the source attribution model. Is not intended to be used by a regular user. Developers only here!

Usage

DPMoDel_impl

Format

[R6Class](#) object.

Value

Object of [R6Class](#).

Fields

`y` 3D array of [type, time, location] of the number of human cases
`X` 3D array of the number of positive samples for each type, source and time [type, source, time]
`R` 3D array of normalised relative prevalences for each timepoint [type, source, time]
`Time` a character vector of timepoint ids matching time dimension in `y` and `R`
`Location` a character vector of location ids matching location dimension in `y`
`Sources` a character vector of source ids matching the source dimension in `X`
`Type` a character vector of type ids matching the type dimension in `X`
`prev` a 2D array (matrix) of [source, time].

q concentration parameter for the DP
theta shape parameter for the Gamma base distribution for the DP
theta rate parameter for the Gamma base distribution for the DP
r 3D array of [type, src, time] for the hyperprior on the relative prevalences R
alpha 3D array of [source, time, location] for the prior on the alpha parameters
s vector giving initial group allocation for each type for the DP
theta vector giving initial values for each group in the DP
alpha 3D array of [source, time, location] giving initial values for the alpha parameters

 FormulaNode

FormulaNode

Description

Represents a formula node in a DAG. Inherit from this node to specify some kind of formula within the DAG, e.g. a linear predictor and/or link function. Override the `FormulaNode$getData()` method to apply your own function.

Usage

FormulaNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [Node](#), please see base class documentation.

Value

Object of [FormulaNode](#)

Methods

`getData()` returns the node's transformed data.

GammaNode

GammaNode

Description

Represents a Gamma distribution node in a DAG. Requires parent nodes for shape and rate respectively as specified in [dgamma](#).

Usage

GammaNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [StochasticNode](#), please see base class documentation.

Value

Object of [GammaNode](#)

Methods

`new(data, shape=1, rate=1)` Create a Gamma node with data data and Nodes shape and rate as specified in [dgamma](#).

Halddp

Builds a Halddp source attribution model

Description

Builds a Halddp source attribution model

Usage

Halddp(y, x, k, priors, a_q, inits = NULL)

Arguments

y	a Y object containing case observations
x	an X object containing source observations
k	a Prev object containing source prevalences
priors	priors list with elements named a_r, a_alpha, a_theta and b_theta, corresponding to the prior parameters for the r, alpha, and base distribution for the DP parameters respectively.

<i>Parameter</i>	<i>Prior Distribution</i>	<i>Prior Parameters</i>
a_r	Dirichlet(concentration)	A single positive number or an X object containing the prior values for each source, time and type. If a single number is supplied, it will be used for all times, sources and types.
a_alpha	Dirichlet(concentration)	A single positive number or an Alpha object containing the prior values for each source, time and location. If a single number is supplied, it will be used for all times, sources and locations.
Type effects base distribution parameters	Gamma(shape, rate)	Single number for each of the shape (a_theta) and rate (b_theta) of the Gamma base distribution.
a_q	the Dirichlet Process concentration parameter.	
inits	initial values for the mcmc algorithm. This is an optional list that may contain any of the following items: alpha,q, and r.	
<i>Parameter</i>	<i>Description</i>	
r	An object of type X giving the initial values for R matrix, If not specified defaults to the element-wise maximum likelihood estimates of r from the source matrix.	
Source effects (alpha)	An object of type Alpha specifying alpha value for each source/time/location. If not specified, default initial values for the source effects are drawn from the prior distribution.	
Type effects (q)	An object of type Q giving the initial clustering and values for q If not specified, defaults to a single group with a theta value calculated as $\theta = \text{sum}(y_{i,t,l}) / \text{sum}_l = 1^L (\text{sum}_t = 1^T (\text{sum}_i = 1^n (\text{sum}_j = 1^m (\text{alpha}_{j,t,l} * r_{i,j,t} * k_{j,t}))))$. i.e. $\theta = \text{sum}(y_{i,t,l}) / \text{sum}(\text{lambda}_{i,j,t,l} / \theta)$	

Format

Object of [R6Class](#) with methods for creating a HaldDP model, running the model, and accessing and plotting the results.

Value

Object of [HaldDP](#) with methods for creating a HaldDP model, running the model, and accessing and plotting the results.

Description

This function fits a non-parametric Poisson source attribution model for human cases of disease. It supports multiple types, sources, times and locations. The number of human cases for each type, time and location follow a Poisson likelihood.

HaldDP Object Methods

`mcmc_params(n_iter = 1000, burn_in = 0, thin = 1, n_r = ceiling(private$nTypes * 0.2), update_schem` when called, sets the mcmc parameters.

`n_iter` sets the number of iterations returned (after removing `burn_in` and thinning results by `thin` i.e. a total of $(n_iter * thin) + burn_in$ iterations are run)

`n_r` is a positive integer that sets the total number of `r_{ijt1}` parameters to be updated at each time-location-source combination (the default is 20 percent updated per iteration)

`update_schema` a character vector containing the parameters to update (any of 'q', 'alpha', 'r').

`update(n_iter, append = TRUE)` when called, updates the HaldDP model by running `n_iter` iterations.

If missing `n_iter`, the `n_iter` last set using `mcmc_params()` or `update()` is used.

`append` is a logical value which determines whether the next `n_iter` iterations are appended to any previous iterations, or overwrites them. When `append = TRUE`, the starting values are the last iteration and no `burn_in` is removed. Running the model for the first time, or changing any model or fitting parameters will set `append = FALSE`.

`get_data` returns a list containing the human data `y` (an array `y[types, times, locations]`), the source data `X` (an array `X[types, sources, times]`), the prevalence data (an array `k[sources, times]`), the type names, source names, time names, location names and number of different types, sources, times and locations.

`get_priors` returns a list containing the DP concentration parameter `a_q`, and the priors (R6 class with members named `a_alpha` (members are array `a_alpha[sources, times, locations]`), `a_r` (an array `a_r[types, sources, times]`), `a_theta` and `b_theta`).

`get_inits` returns an R6 class holding the initial values (members are `alpha` (an array `alpha[sources, times, locations]`), `theta` (an array `theta[types, iters]`), `s` (an array `s[types, iters]`), and `r` (an array `r[types, sources, times]`)).

`get_mcmc_params` returns a list of fitting parameters (`n_iter`, `append`, `burn_in`, `thin`, `update_schema` (R6 class with members `alpha`, `q`, `r`)).

`get_acceptance` returns an R6 class containing the acceptance rates for each parameter (members are `alpha` (an array `alpha[sources, times, locations]`), and `r` (an array `r[types, sources, times]`)).

`extract(params = c("alpha", "q", "s", "r", "lambda_i", "xi", "xi_prop"), times = NULL, locations = NULL)` returns a list containing a subset of the parameters (determined by the `params` vector, `times`, `locations`, `sources`, `types` and `iters`).

If `flatten` is set to `TRUE`, it returns a dataframe with 1 column per parameter, otherwise it returns a list containing `params` containing a subset of the following arrays: `alpha[Sources, Times, Locations, iters]`, `q[Types, iters]`, `s[Types, iters]`, `r[Types, Sources, Times, iters]`, `lambda_i[Types, Times, Locations, iters]`, `xi[Sources, Times, Locations, iters]`.

`drop` determines whether to delete the dimensions of an array which have only one level when `flatten = FALSE`.

`summary(alpha = 0.05, params = c("alpha", "q", "s", "r", "lambda_i", "xi", "xi_prop"), times = NULL)` returns a list containing the median and credible intervals for a subset of the parameters. The default credible interval type are Chen-Shao ("chen-shao") highest posterior density intervals (alternatives are "percentiles" and "spin"). See `extract` for details on the subsetting. `xi_prop` returns the proportion of cases attributed to each source `j` and is calculated by dividing each iteration of `lambda_{jt1}` values by their sum within each time `t` and location `l`.

`plot_heatmap(iters, cols = c("blue", "white"), hclust_method = "complete")` Creates a dendrogram and heatmap for the type effect groupings (`s` parameter in the model). This uses the `heatmap.2` function from `gplots`.

iters is a vector containing the iterations to be used in constructing the graph. Default is all iterations in posterior.

hclust_method allows the user to select the method used by stats::hclust to cluster the type effect groupings s.

cols gives the colours for completely dissimilar (dissimilarity value of 1), and identical (dissimilarity value of 0). All other values will be in between the two chosen colours. See ?colorRampPalette for more details..

Details

This function fits a source attribution model for human cases of disease. It supports multiple types, sources, times and locations. The number of human cases for each type, time and location follows a Poisson or Negative Binomial likelihood. *Model*

$$y_{itl} \sim \text{Poisson}(\lambda_{itl})$$

where

$$\lambda_{itl} = \sum_{j=1}^m \lambda_{ijtl} = q_{k(i)} \sum_{j=1}^m (r_{ijt} \cdot k_j \cdot \text{alpha}_{jtl})$$

The parameters are defined as follows:

$$a_{jtl}$$

is the unknown source effect for source j , time t , location l

$$q_{s(i)}$$

is the unknown type effect for type i in group s .

$$x_{ij}$$

is the known number of positive samples for each source j type i combination

$$n_{ij}$$

is the known total number of samples for each source j type i combination

$$k_j$$

is the fixed prevalence in source (i.e. the number of positive samples divided by the number of negative samples) j

$$r_{ijt}$$

is the unknown relative occurrence of type i on source j .

Priors

$$r_{.jt} \sim \text{Dirichlet}(a_{r_{1jt}}, \dots, a_{r_{njt}})$$

$$a_{tl} \sim \text{Dirichlet}(a_{\text{alpha}_{1tl}}, \dots, a_{\text{alpha}_{mtl}})$$

$$q \sim \text{DP}(a_q, \text{Gamma}(a_{\text{theta}}, b_{\text{theta}}))$$

Author(s)

Chris Jewell and Poppy Miller <p.miller at lancaster.ac.uk>

References

Chen, M.-H. and Shao, Q.-M. (1998). Monte Carlo estimation of Bayesian credible and HPD intervals, *Journal of Computational and Graphical Statistics*, 7.

Liu Y, Gelman A, Zheng T (2015). Simulation-efficient shortest probability intervals, *Statistics and Computing*.

Examples

```
## Not run:
#### Format data using Y, X, and Prev functions #####
## Input data must be in long format
y <- Y(
  # Cases
  data = sim_SA$cases,
  y = "Human",
  type = "Type",
  time = "Time",
  location = "Location"
)

x <- X(
  # Sources
  data = sim_SA$sources,
  x = "Count",
  type = "Type",
  time = "Time",
  source = "Source"
)

k <- Prev(
  # Prevalences
  data = sim_SA$prev,
  prev = "Value",
  time = "Time",
  source = "Source"
)

#### Create Dirichlet(1) priors #####

## Create alpha prior data frame
prior_alpha_long <- expand.grid(
  Source = unique(sim_SA$sources$Source),
  Time = unique(sim_SA$sources$Time),
  Location = unique(sim_SA$cases$Location),
  Alpha = 1
)
# Use the Alpha() constructor to specify alpha prior
prior_alpha <- Alpha(
  data = prior_alpha_long,
  alpha = 'Alpha',
  source = 'Source',
  time = 'Time',
  location = 'Location'
)
```

```

## Create r prior data frame
prior_r_long <- expand.grid(
  Type   = unique(sim_SA$sources$Type),
  Source = unique(sim_SA$sources$Source),
  Time   = unique(sim_SA$sources$Time),
  Value  = 0.1
)
# Use X() constructor to specify r prior
prior_r <- X(
  data   = prior_r_long,
  x      = 'Value',
  type   = 'Type',
  time   = 'Time',
  source = 'Source'
)

## Pack all priors into a list
priors <- list(
  a_theta = 0.01,
  b_theta = 0.00001,
  a_alpha = prior_alpha,
  a_r      = prior_r
)

## If all prior values are the same, they can be specified in shorthand
## Equivalent result to the longform priors specified above
priors <- list(
  a_theta = 0.01,
  b_theta = 0.00001,
  a_alpha = 1,
  a_r      = 0.1
)

##### Set initial values (optional) #####
types <- unique(sim_SA$cases$Type)
q_long <- data.frame(q=rep(15, length(types)), Type=types)
init_q <- Q(q_long, q = 'q', type = 'Type')
inits <- list(q = init_q) # Pack starting values into a list

##### Construct model #####
my_model <- HaldDP(y = y, x = x, k = k, priors = priors, inits = inits, a_q = 0.1)

##### Set mcmc parameters #####
my_model$mcmc_params(n_iter = 30, burn_in = 2, thin = 1)

##### Update model #####
my_model$update()
## Add an additional 10 iterations
my_model$update(n_iter = 10, append = TRUE)

##### Extract posterior #####
## returns the posterior for the r, alpha, q, c,

```



```

## lambda_i, xi and xi_prop parameters,
## for all times, locations, sources and types
## the posterior is returned as a list or arrays
my_model$extract()

## returns the posterior for the r and alpha parameters,
## for time 1, location B, sources Source3, and Source4,
## types 5, 25, and 50, and iterations 200:300
## the posterior is returned as a list of dataframes
my_model$extract(params = c("r", "alpha"),
                 times = "1", location = "B",
                 sources = c("Source3", "Source4"),
                 types = c("5", "25", "50"),
                 iters = 20:30,
                 flatten = TRUE)

#### Calculate medians and credible intervals #####
my_model$summary(alpha = 0.05, CI_type = "percentiles")
## subsetting is done in the same way as extract()
my_model$summary(alpha = 0.05, CI_type = "chen-shao",
                 params = c("r", "alpha"),
                 times = "1", location = "B",
                 sources = c("Source3", "Source4"),
                 types = c("5", "25", "50"),
                 iters = 20:30,
                 flatten = TRUE)

#### Plot heatmap and dendrogram of the type effect grouping #####
my_model$plot_heatmap()

#### Extract data, initial values, prior values, acceptance
## rates for the mcmc algorithm and mcmc parameters
my_model$get_data()
my_model$get_inits()
my_model$get_priors()
my_model$get_acceptance()
my_model$get_mcmc_params()

## End(Not run)

```

Node

Node

Description

This is a base class representing a node in a DAG. Is not intended to be used by a regular user. Developers only here!

Usage

Node

Format

Object of [R6Class](#) with methods for constructing a DAG.

Value

Object of [Node](#)

Fields

parents a list of parent nodes
 children a list of child nodes
 name a tag name applied to the node

Methods

`new(parents = list(), children = list(), name)` creates a new [Node](#) with parent nodes, child nodes, and a name.
`logDensity()` calculate the log probability density/mass function evaluated at the current node value.
`addChild(node)` add node as a child. Returns node.
`addParent(node)` add node as a parent. Returns node.
`removeParent(name)` remove the parent node named name. Returns node.
`removeChild(name)` remove the child node named name. Returns node.

<code>PoisGammaDPUpdate</code>	<i>PoisGammaDPUpdate</i>
--------------------------------	--------------------------

Description

This class implements a marginal Gibbs sampler for a Dirichlet process prior on the mean of a Poisson distributed random variable, with a Gamma-distributed base function.

Usage

`PoisGammaDPUpdate`

Format

Object of [R6Class](#) with methods for updating a [DirichletProcessNode](#) instance.

Details

The marginal Gibbs sampler is based on the description in Gelman et al. Bayesian Data Analysis, 3rd edition, Chapter 23.

Please note that no checks are performed as to the suitability of this algorithm for a particular [StochasticNode](#). It is up to the user to use the correct update algorithm for the appropriate nodes.

Value

Object of [PoisGammaDPUpdate](#)

Methods

`new(node)` constructor takes an instance of a [DirichletProcessNode](#) node

`update()` when called, updates node

PoissonNode

PoissonNode

Description

Represents a Poisson distribution node in a DAG

Usage

PoissonNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [StochasticNode](#), please see base class documentation.

Value

Object of [PoissonNode](#)

Methods

`new(data, lambda, offset)` create a PoissonNode, with mean [Node](#) lambda, and offset [Node](#) offset.

Prev	<i>Constructs prevalence data</i>
------	-----------------------------------

Description

The Prev constructor function returns an R6 Prevalence class which feeds data into sourceR models.

Usage

```
Prev(data, prev, source, time = NULL)
```

Arguments

data	long-format data.frame containing prevalence data by source and time.
prev	character string giving name of prevalence column in data
source	character string giving name of source column in data
time	optional column denoting times of prevalence observation

Value

A Prev data structure for use in sourceR models

Q	<i>Constructs initial values for q</i>
---	--

Description

The Q constructor returns a R6 Q_ class which feeds sanitised initial values for q into the model.

Usage

```
Q(data, q, type)
```

Arguments

data	long-format data.frame containing type-name and q-value for each observation.
q	name of 'q' column
type	name of type column

Value

A Q_ data structure for use in sourceR models

sim_SA	<i>Simulated data: Human cases of campylobacteriosis and numbers of source samples positive for Campylobacter.</i>
--------	--

Description

A simulated dataset containing the number of human cases of campylobacteriosis, the numbers of source samples positive for *Campylobacter* for each bacterial subtype, and the overall source prevalence.

Usage

```
sim_SA
```

Format

A list containing the human cases ('cases'), source samples ('sources'), prevalences ('prev') and true values ('truevals').

cases: data frame with 364 rows and 4 variables:

Human number of human cases of campylobacteriosis

Time Time id for the samples

Location Location id for the samples

Type MLST type id for the samples

sources: data frame with 1092 rows and 4 variables

Count number of source samples positive for campylobacteriosis

Time Time id for the samples

Source Source id for the samples

Type MLST type id for the samples

prev: data frame with 12 rows and 3 variables

Value Prevalence value (number of positive samples divided by total number of samples)

Time Time id for the samples

Source Source id for the samples

truevals: list containing a long format data frame for each model parameter giving the true value of that parameter.

alpha A dataframe with 24 rows and 4 variables: Value contains the true alpha values, Time, Location and Source contain the time, location and source id's respectively.

q A dataframe with 91 rows and 2 variables: Value contains the true q values, and Type contains the type id's.

- lambda_i** A dataframe with 364 rows and 4 variables: Value contains the true lambda_i values, Time, Location and Type contain the time, location and type id's respectively.
- xi** A dataframe with 24 rows and 4 variables: Value contains the true xi values, Time, Location and Source contain the time, location and source id's respectively.
- r** A dataframe with 2184 rows and 5 variables: Value contains the true r values, Time, Type, Location and Source contain the time, type, location and source id's respectively.

<code>sliceTensor</code>	<i>Slices a tensorA::tensor</i>
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Description

Slices a `tensorA::tensor`, preserving the dimnames. This is a workaround for a buggy implementation of `[.tensor` as of `tensorA v0.36`.

Usage

```
sliceTensor(x, ...)
```

Arguments

<code>x</code>	tensor array to be sliced
<code>...</code>	arguments used to subset the sensor array

<code>sourceR</code>	<i>sourceR: A package for fitting Bayesian non-parametric source attribution models.</i>
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Description

The `sourceR` package currently provides an S6 class called `HaldDP` to fit the HaldDP model. Further source attribution models are planned to be added to the package.

sourceR methods

The main `sourceR` method is [HaldDP](#).

 StochasticNode

StochasticNode

Description

Represents a stochastic node in a DAG

Usage

StochasticNode

Format

Object of [R6Class](#) with methods for constructing a DAG.

Details

Derived from [Node](#), please see base class documentation.

Value

Object of [StochasticNode](#)

Fields

data contains the node's data

Methods

logPosterior() return the value of the log posterior distribution of the node.

getData() returns the node's data.

 X

Constructs source data

Description

The X constructor function returns an R6 X class which feeds source data into sourceR models.

Usage

X(data, x, type, source, time = NULL)

Arguments

data	long-format data.frame containing source data
x	character string giving name of source counts column in data
type	character string giving name of type column in data
source	character string giving name of source column in data
time	optional column denoting times of source observation

Value

A X source data structure for use in sourceR models

Y	<i>Constructs disease count data</i>
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Description

The Y constructor function returns an R6 Y class which feeds disease count data into sourceR models.

Usage

```
Y(data, y, type, time = NULL, location = NULL)
```

Arguments

data	long-format data.frame containing source data
y	character string giving name of disease counts column in data
type	character string giving name of type column in data
time	optional column denoting times of disease count observations
location	optional column denoting location of disease count observations

Value

A Y disease count data structure for use in sourceR models

Index

*Topic **DAG**

- AdaptiveDirMRW, [2](#)
- AdaptiveLogDirMRW, [3](#)
- AdaptiveMultiMRW, [4](#)
- DataNode, [7](#)
- DirichletNode, [8](#)
- DirichletProcessNode, [8](#)
- FormulaNode, [10](#)
- GammaNode, [11](#)
- Node, [17](#)
- PoisGammaDPUpdate, [18](#)
- PoissonNode, [19](#)
- StochasticNode, [23](#)

*Topic **Dirichlet-process**

- PoisGammaDPUpdate, [18](#)

*Topic **MCMC**

- AdaptiveDirMRW, [2](#)
- AdaptiveLogDirMRW, [3](#)
- AdaptiveMultiMRW, [4](#)

*Topic **datasets**

- Alpha_, [6](#)
- campy, [6](#)
- DPMoDel_impl, [9](#)
- sim_SA, [21](#)

*Topic **node**

- AdaptiveDirMRW, [2](#)
- AdaptiveLogDirMRW, [3](#)
- AdaptiveMultiMRW, [4](#)
- DataNode, [7](#)
- DirichletNode, [8](#)
- DirichletProcessNode, [8](#)
- FormulaNode, [10](#)
- GammaNode, [11](#)
- Node, [17](#)
- PoisGammaDPUpdate, [18](#)
- PoissonNode, [19](#)
- StochasticNode, [23](#)

AdaptiveDirMRW, [2](#), [3](#), [4](#)

AdaptiveLogDirMRW, [3](#), [4](#)

AdaptiveMultiMRW, [4](#), [5](#)

Alpha, [5](#), [12](#)

Alpha_, [6](#)

campy, [6](#)

DataNode, [7](#), [7](#)

dgamma, [11](#)

DirichletNode, [2](#), [3](#), [8](#), [8](#)

DirichletProcessNode, [8](#), [9](#), [18](#), [19](#)

DPMoDel_impl, [9](#)

FormulaNode, [10](#), [10](#)

GammaNode, [11](#), [11](#)

HaldDP, [11](#), [12](#), [22](#)

Node, [5](#), [7](#), [10](#), [17](#), [18](#), [19](#), [23](#)

PoisGammaDPUpdate, [18](#), [19](#)

PoissonNode, [19](#), [19](#)

Prev, [11](#), [20](#)

Q, [12](#), [20](#)

R6Class, [2](#), [3](#), [5](#), [7–12](#), [18](#), [19](#), [23](#)

sim_SA, [21](#)

sliceTensor, [22](#)

sourceR, [22](#)

sourceR-package (sourceR), [22](#)

StochasticNode, [3–5](#), [8](#), [9](#), [11](#), [19](#), [23](#), [23](#)

X, [11](#), [12](#), [23](#)

Y, [11](#), [24](#)