# Package 'dlmtree'

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Type Package

Title Bayesian Treed Distributed Lag Models

Version 1.1.0

Description Estimation of distributed lag models (DLMs) based on a Bayesian additive regression trees framework. Includes several extensions of DLMs: treed DLMs and distributed lag mixture models (Mork and Wilson, 2023) <doi:10.1111/biom.13568>; treed distributed lag nonlinear models (Mork and Wilson, 2022) <doi:10.1093/biostatistics/kxaa051>; heterogeneous DLMs (Mork, et. al., 2024) <doi:10.1080/01621459.2023.2258595>; monotone DLMs (Mork and Wilson, 2024) <doi:10.1214/23-BA1412>. The package also includes visualization tools and a 'shiny' interface to check model convergence and to help interpret results.

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https://danielmork.github.io/dlmtree/

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## adj\_coexposure

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adj\_coexposure Adjusting for expected changes in co-exposure (TDLMM)

## Description

Estimates the marginal effects of an exposure while accounting for expected changes in co-occurring exposures at the same time point. Values of co-occurring exposures are modeled nonlinearly using a spline model with predictions made at the lower an upper values for the exposure of interest.

#### Usage

```
adj_coexposure(
  exposure.data,
  object,
  contrast_perc = c(0.25, 0.75),
  contrast_exp = list(),
  conf.level = 0.95,
  keep.mcmc = FALSE,
  verbose = TRUE
)
```

#### Arguments

exposure.data	Named list of exposure matrices used as input to TDLMM.
object	Model output for TDLMM from dlmtree() function.
contrast_perc	2-length vector of percentiles or named list corresponding to lower and upper exposure percentiles of interest. Names must equal list names in 'exposure.data'.
contrast_exp	Named list consisting lower and upper exposure values. This takes precedence over contrast_perc if both inputs are used.
conf.level	Confidence level used for estimating credible intervals. Default is 0.95.
keep.mcmc	If TRUE, return posterior samples.
verbose	TRUE (default) or FALSE: print output

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## Details

adj\_coexposure

#### Value

A list with the following components (or posterior samples if keep.mcmc = TRUE):

Name	vector of exposure names
Time	integer vector of lags
Effect	posterior mean of marginal effects
SE	standard error of the estimate
Lower	lower bound of credible interval of the marginal effect estimate
Upper	upper bound of credible interval of the marginal effect estimate
cEffect	cumulative marginal effects
cLower	lower bound of credible interval of the cumulative marginal effect
cUpper	upper bound of credible interval of the cumulative marginal effect
CW	boolean vector indicating critical window

coExp

Randomly sampled exposure from Colorado counties

## Description

Matrix of five different exposures, each measured over 40 weeks.

## Usage

data(coExp)

## Format

matrix

## Source

https://aqs.epa.gov/aqsweb/airdata/download\_files.html

## References

https://www.epa.gov/outdoor-air-quality-data

combine.models

## Description

Method for combining information from DLMs of single exposure

#### Usage

```
combine.models(mlist)
```

## Arguments

mlist a list of models

## Details

combine.models

#### Value

A data frame with model fit information of the models included in the list

combine.models.tdlmm Combines information from DLMs of mixture exposures.

## Description

Method for combining information from DLMs of mixture exposures.

#### Usage

combine.models.tdlmm(mlist)

## Arguments

mlist a list of models

## Details

combine.models.tdlmm

## Value

A data frame with model fit information of the models included in the list

cppIntersection

#### Description

fast set intersection tool assumes sorted vectors A and B

## Usage

```
cppIntersection(A, B)
```

## Arguments

Α	sorted integer vector A
В	sorted integer vector B

## Value

vector of resulting intersection

diagnose.summary.hdlm diagnose

#### Description

diagnose generic function for S3method

## Usage

```
## S3 method for class 'summary.hdlm'
diagnose(x, ...)
## S3 method for class 'summary.hdlmm'
diagnose(x, ...)
## S3 method for class 'summary.monotone'
diagnose(x, ...)
## S3 method for class 'summary.tdlm'
diagnose(x, ...)
## S3 method for class 'summary.tdlmm'
diagnose(x, ...)
## S3 method for class 'summary.tdlmm'
```

## dlmEst

diagnose(x, ...)
diagnose(x, ...)

## Arguments

x	a summary object resulting from summary() applied to an object of class 'tdlm', 'tdlmm', 'tdlnm', 'hdlm', 'hdlmm', 'monotone'
	not used.

## Value

shiny interface for assessing model convergence. The interface includes tabs for MCMC diagnostics such as trace plots, density plots, and convergence measures for distributed lag effects, DLM tree sizes, and hyperparameters.

dlmEst	Calculates the distributed lag effect with DLM matrix for linear mod-
	els.

## Description

Calculates the distributed lag effect with DLM matrix for linear models.

#### Usage

```
dlmEst(dlm, nlags, nsamp)
```

## Arguments

dlm	A numeric matrix containing the model fit information
nlags	total number of lags
nsamp	number of mcmc iterations

#### Value

A cube object of lag effect x lag x mcmc

#### dlmtree

#### Description

The 'dlmtree' function accommodates various response variable types, including continuous, binary, and zero-inflated count values. The function is designed to handle both single exposure and exposure mixtures. For a single exposure, users are offered options to model non-linear effects (tdlnm), linear effects (tdlm), or heterogeneous subgroup/individualized effects (hdlm). In the case of exposure mixtures, the function supports lagged interactions (tdlnm), and heterogeneous subgroup/individualized effects (hdlmm) allowing for a comprehensive exploration of mixture exposure heterogeneity. Additionally, users can fine-tune parameters to impose effect shrinkage and perform exposure selection, enhancing the adaptability and precision of the modeling process. For more detailed documentation, visit: dlmtree website.

#### Usage

```
dlmtree(
  formula,
  data,
  exposure.data,
  dlm.type = "linear",
  family = "gaussian",
 mixture = FALSE,
 het = FALSE,
  control.mcmc = list(),
  control.hyper = list(),
  control.family = list(),
  control.tdlnm = list(),
  control.het = list(),
  control.mix = list().
  control.monotone = list(),
  control.diagnose = list()
)
```

#### Arguments

formula	object of class formula, a symbolic description of the fixed effect model to be fitted, e.g. $y \sim a + b$ .
data	data frame containing variables used in the formula.
exposure.data	numerical matrix of exposure data with same length as data, for a mixture setting (tdlmm, hdlmm): named list containing equally sized numerical matrices of exposure data having same length as data.
dlm.type	dlm model specification: "linear" (default), "nonlinear", "monotone".
family	'gaussian' for continuous response, 'logit' for binomial, 'zinb' for zero-inflated negative binomial.

#### dlmtree

mixture	flag for mixture, set to TRUE for tdlmm and hdlmm (default: FALSE).
het	flag for heterogeneity, set to TRUE for hdlm and hdlmm (default: FALSE).
control.mcmc	list of MCMC control parameters. This is passed to dlmtree.control.mcmc.
control.hyper	list of hyperparameter control parameters. This is passed to dlmtree.control.hyper
control.family	list of family control parameters. This is passed to dlmtree.control.family
control.tdlnm	list of TDLNM control parameters. This is passed to dlmtree.control.tdlnm
control.het	list of control parameters for heterogeneous models. This is passed to dlmtree.control.het
control.mix	list of mixture control parameters. This is passed to dlmtree.control.mix
control.monotor	ne
	list of control parameters for monotone model. This is passed to dlmtree.control.monotone
control.diagnos	se
	list of control parameters for diagnostics. This is passed to dlmtree.control.diagnose

#### Details

#### dlmtree

Model is recommended to be run for at minimum 5000 burn-in iterations followed by 15000 sampling iterations with a thinning factor of 5. Convergence can be checked by re-running the model and validating consistency of results. Examples are provided below for the syntax for running different types of models. For more examples, visit: dlmtree website.

#### Value

object of one of the classes: tdlm, tdlmm, tdlnm, hdlm, hdlmm, monotone

## Examples

# The first three examples are for one lagged exposure

```
# Treed distributed lag nonlinear model (TDLNM)
# Gaussian regression model
D <- sim.tdlnm(sim = "A", error.to.signal = 1)</pre>
tdlnm.fit <- dlmtree(formula = y ~ .,</pre>
                      data = D$dat,
                      exposure.data = D$exposures,
                      dlm.type = "nonlinear",
                      family = "gaussian")
# summarize results
tdlnm.sum <- summary(tdlnm.fit)</pre>
tdlnm.sum
# plot results
plot(tdlnm.sum)
# Heterogeneous TDLM (HDLM), similar to first example but with heterogeneous exposure response
D <- sim.hdlmm(sim = "B", n = 1000)
hdlm.fit <- dlmtree(y ~ .,</pre>
                     data = D$dat,
                     exposure.data = D$exposures,
                     dlm.type = "linear",
                     family = "gaussian",
                     het = TRUE)
# summarize results
hdlm.sum <- summary(hdlm.fit)</pre>
hdlm.sum
# shiny app for HDLM
if (interactive()) {
  shiny(hdlm.fit)
}
# The next two examples are for a mixture (or multivariate) exposure
# Treed distributed lag mixture model (TDLMM)
# Model for mixutre (or multivariate) lagged exposures
# with a homogenious exposure-time-response function
D <- sim.tdlmm(sim = "B", error = 25, n = 1000)</pre>
tdlmm.fit <- dlmtree(y ~ .,</pre>
                      data = D$dat, exposure.data = D$exposures,
                      control.mix = list(interactions = "noself"),
                      dlm.type = "linear", family = "gaussian",
                      mixture = TRUE)
```

```
# summarize results
tdlmm.sum <- summary(tdlmm.fit)</pre>
# plot the marginal exposure-response for one exposure
plot(tdlmm.sum, exposure1 = "e1")
# plot exposure-response surface
plot(tdlmm.sum, exposure1 = "e1", exposure2 = "e2")
# heterogeneous version of TDLMM
D <- sim.hdlmm(sim = "D", n = 1000)
hdlmm.fit <- dlmtree(y ~ .,</pre>
                     data = D$dat,
                      exposure.data = D$exposures,
                     dlm.type = "linear",
                      family = "gaussian",
                     mixture = TRUE,
                      het = TRUE)
# summarize results
hdlmm.sum <- summary(hdlmm.fit)</pre>
hdlmm.sum
# summarize results
if (interactive()) {
 shiny(hdlmm.fit)
}
```

dlmtree.control.diagnose

Diagnostic control settings for dlmtree model fitting

## Description

Diagnostic control settings for dlmtree model fitting

## Usage

```
dlmtree.control.diagnose(
   subset = NULL,
   lowmem = FALSE,
   verbose = TRUE,
   save.data = TRUE,
   diagnostics = FALSE,
```

```
initial.params = NULL
)
```

## Arguments

subset	integer vector to analyze only a subset of data and exposures.
lowmem	TRUE or FALSE (default): turn on memory saver for DLNM, slower computation time.
verbose	TRUE (default) or FALSE: print output
save.data	TRUE (default) or FALSE: save data used for model fitting. This must be set to TRUE to use shiny() function on hdlm or hdlmm
diagnostics	TRUE or FALSE (default) keep model diagnostic such as the number of terminal nodes and acceptance ratio.
initial.params	initial parameters for fixed effects model, FALSE = none (default), "glm" = generate using GLM, or user defined, length must equal number of parameters in fixed effects model.

## Value

list of control parameters for diagnostics.

dlmtree.control.family

Family control settings for dlmtree model fitting

## Description

Family control settings for dlmtree model fitting

## Usage

```
dlmtree.control.family(binomial.size = 1, formula.zi = NULL)
```

## Arguments

binomial.size	integer type scalar (if all equal, default: 1) or vector defining binomial size for 'logit' family.
formula.zi	(only applies to family = 'zinb') object of class formula, a symbolic description of the fixed effect of zero-inflated (ZI) model to be fitted, e.g. $y \sim a + b$ . This only applies to ZINB where covariates for ZI model are different from NB model. This is set to the argument 'formula' by default.

## Value

list of family control parameters.

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dlmtree.control.het Control settings for dlmtree model fitting, when used for heterogeneous models

## Description

Control settings for dlmtree model fitting, when used for heterogeneous models

## Usage

```
dlmtree.control.het(
  modifiers = "all",
  modifier.splits = 20,
  modtree.params = c(0.95, 2),
  modtree.step.prob = c(0.25, 0.25, 0.25),
  dlmtree.type = "shared",
  selection.prior = 0.5
)
```

## Arguments

modifiers	string vector containing desired modifiers to be included in a modifier tree. The	
	strings in the vector must match the names of the columns of the data. By	
	default, a modifier tree considers all covariates in the formula as modifiers unless stated otherwise.	
modifier.splits		
	integer value to determine the possible number of splitting points that will be used for a modifier tree.	
modtree.params	numerical vector of alpha and beta hyperparameters controlling modifier tree depth. (default: $alpha = 0.95$ , $beta = 2$ )	
modtree.step.prob		
	numerical vector for probability of each step for modifier tree updates: 1) grow, 2) prune, 3) change. (default: $c(0.25, 0.25, 0.25))$	
dlmtree.type	specification of dlmtree type for HDLM: shared (default) or nested.	
selection.prior		
	scalar hyperparameter for sparsity of modifiers. Must be between 0.5 and 1. Smaller value corresponds to increased sparsity of modifiers.	

## Value

list of control parameters for heterogeneous models.

dlmtree.control.hyper Hyperparameter control settings for dlmtree model fitting

#### Description

Hyperparameter control settings for dlmtree model fitting

## Usage

```
dlmtree.control.hyper(
    shrinkage = "all",
    params = c(0.95, 2),
    step.prob = c(0.25, 0.25)
)
```

## Arguments

shrinkage	character "all" (default), "trees", "exposures", "none", turns on horseshoe-like shrinkage priors for different parts of model.
params	numerical vector of alpha and beta hyperparameters controlling dlm tree depth. (default: $alpha = 0.95$ , $beta = 2$ )
step.prob	numerical vector for probability of each step for dlm tree updates: 1) grow/prune, 2) change, 3) switch exposure. (default: c(0.25, 0.25, 0.25))

## Value

list of hyperparameter control parameters.

dlmtree.control.mcmc MCMC control settings for dlmtree model fitting

## Description

MCMC control settings for dlmtree model fitting

## Usage

```
dlmtree.control.mcmc(n.trees = 20, n.burn = 1000, n.iter = 2000, n.thin = 10)
```

#### Arguments

n.trees	integer for number of trees in ensemble.
n.burn	integer for length of MCMC burn-in.
n.iter	integer for number of MCMC iterations to run model after burn-in.
n.thin	integer MCMC thinning factor, i.e. keep every tenth iteration.

#### dlmtree.control.mix

#### Value

list of MCMC control parameters.

dlmtree.control.mix Control settings for dlmtree model fitting, when used for mixture models

## Description

Control settings for dlmtree model fitting, when used for mixture models

#### Usage

```
dlmtree.control.mix(interactions = "noself", sparsity.prior = 1)
```

#### Arguments

interactions	'noself' (default) which estimates interactions only between two different expo-
	sures, 'all' which also allows interactions within the same exposure, or 'none'
	which eliminates all interactions and estimates only main effects of each expo-
	sure.

sparsity.prior positive scalar hyperparameter for sparsity of exposures. (default: 1)

#### Value

list of mixture control parameters.

dlmtree.control.monotone

Control settings for dlmtree model fitting, when used for monotone model

#### Description

Control settings for dlmtree model fitting, when used for monotone model

#### Usage

```
dlmtree.control.monotone(
  gamma0 = NULL,
  sigma = NULL,
  tree.time.params = c(0.95, 2),
  tree.exp.params = c(0.95, 2),
  time.kappa = NULL
)
```

#### Arguments

gamma0	vector (with length equal to number of lags) of means for logit-transformed prior probability of split at each lag; e.g., gamma_0l = 0 implies mean prior probability of split at lag $l = 0.5$ .	
sigma	symmetric matrix (usually with only diagonal elements) corresponding to gamma_0 to define variances on prior probability of split; e.g., gamma_0l = 0 with lth diagonal element of sigma=2.701 implies that 95% of the time the prior probability of split is between 0.005 and 0.995, as a second example setting gamma_0l=4.119 and the corresponding diagonal element of sigma=0.599 implies that 95% of the time the prior probability of a split is between 0.8 and 0.99.	
tree.time.params		
	numerical vector of hyperparameters for monotone time tree.	
tree.exp.params		
	numerical vector of hyperparameters for monotone exposure tree.	
time.kappa	scaling factor in dirichlet prior that goes alongside 'time.split.prob' to control the amount of prior information given to the model for deciding probabilities of splits between adjacent lags.	

#### Value

list of control parameters for monotone model.

dlmtree.control.tdlnm Control settings for dlmtree model fitting, when used for TDLNM

## Description

Control settings for dlmtree model fitting, when used for TDLNM

#### Usage

```
dlmtree.control.tdlnm(
    exposure.splits = 20,
    time.split.prob = NULL,
    exposure.se = NULL
)
```

## Arguments

```
exposure.splits
```

scalar indicating the number of splits (divided evenly across quantiles of the exposure data) or list with two components: 'type' = 'values' or 'quantiles', and 'split.vals' = a numerical vector indicating the corresponding exposure values or quantiles for splits.

time.split.prob	
	probability vector of a spliting probabilities for time lags. (default: uniform probabilities)
exposure.se	numerical matrix of exposure standard errors with same size as exposure.data or a scalar smoothing factor representing a uniform smoothing factor applied to each exposure measurement. (default: sd(exposure.data)/2)

#### Value

list of TDLNM control parameters.

dlmtreeGPFixedGaussian

dlmtree model with fixed Gaussian process approach

#### Description

dlmtree model with fixed Gaussian process approach

#### Usage

```
dlmtreeGPFixedGaussian(model)
```

#### Arguments

model

A list of parameter and data contained for the model fitting

#### Value

A list of dlmtree model fit, mainly posterior mcmc samples

dlmtreeGPGaussian dlmtree model with Gaussian process approach

## Description

dlmtree model with Gaussian process approach

#### Usage

```
dlmtreeGPGaussian(model)
```

#### Arguments

model A list of parameter and data contained for the model fitting

## Value

dlmtreeHDLMGaussian dlmtree model with shared HDLM approach

#### Description

dlmtree model with shared HDLM approach

## Usage

```
dlmtreeHDLMGaussian(model)
```

## Arguments

model A list of parameter and data contained for the model fitting

#### Value

A list of dlmtree model fit, mainly posterior mcmc samples

dlmtreeHDLMMGaussian dlmtree model with HDLMM approach

## Description

dlmtree model with HDLMM approach

## Usage

```
dlmtreeHDLMMGaussian(model)
```

## Arguments

model A list of parameter and data contained for the model fitting

## Value

dlmtreeTDLMFixedGaussian

dlmtree model with fixed Gaussian approach

#### Description

dlmtree model with fixed Gaussian approach

## Usage

dlmtreeTDLMFixedGaussian(model)

#### Arguments

model A list of parameter and data contained for the model fitting

## Value

A list of dlmtree model fit, mainly posterior mcmc samples

```
dlmtreeTDLMNestedGaussian
```

dlmtree model with nested Gaussian approach

## Description

dlmtree model with nested Gaussian approach

## Usage

dlmtreeTDLMNestedGaussian(model)

## Arguments

model A list of parameter and data contained for the model fitting

## Value

dlmtreeTDLM\_cpp

## Description

dlmtree model with nested HDLM approach

## Usage

dlmtreeTDLM\_cpp(model)

## Arguments

model

A list of parameter and data contained for the model fitting

## Value

A list of dlmtree model fit, mainly posterior mcmc samples

dlnmEst	Calculates the distributed lag effect with DLM matrix for non-linear
	models.

## Description

Calculates the distributed lag effect with DLM matrix for non-linear models.

## Usage

dlnmEst(dlnm, predAt, nlags, nsamp, center, se)

## Arguments

dlnm	A numeric matrix containing the model fit information
predAt	Number of splits in the model
nlags	total number of lags
nsamp	number of mcmc iterations
center	center parameter
se	Standard error parameter

## Value

A cube object of lag effect x lag x mcmc

dlnmPLEst

*Calculates the distributed lag effect with DLM matrix for non-linear models.* 

## Description

Calculates the distributed lag effect with DLM matrix for non-linear models.

## Usage

dlnmPLEst(dlnm, predAt, nlags, nsamp, center)

## Arguments

dlnm	A numeric matrix containing the model fit information
predAt	Number of splits in the model
nlags	total number of lags
nsamp	number of mcmc iterations
center	center parameter

## Value

A cube object of lag effect x lag x mcmc

drawTree Draws a new tree structure
-------------------------------------

## Description

A recursive method for drawing a new tree structure

## Usage

```
drawTree(depth, alpha, beta)
```

## Arguments

depth	depth of a tree
alpha	tree shape parameter, $0 < alpha < 1$
beta	tree size parameter, beta $> 0$

## Details

drawTree

## Value

integer value of number of terminal nodes

estDLM

Calculates subgroup-specific lag effects for heterogeneous models

## Description

Method for calculating subgroup-specific lag effects for heterogeneous models: HDLM, HDLMM

## Usage

```
estDLM(
   object,
   new.data,
   group.index,
   conf.level = 0.95,
   exposure = NULL,
   keep.mcmc = FALSE,
   mem.safe = FALSE,
   verbose = TRUE
)
```

## Arguments

object	object of a model fit. Must be 'hdlm' or 'hdlmm'
new.data	data frame with new observations with the same number of modifiers
group.index	list of index (row numbers) for subgroup specification
conf.level	confidence level for credible interval of effects
exposure	exposure of interest for 'hdlmm' method
keep.mcmc	store mcmc in the output
mem.safe	boolean memory parameter for rule index
verbose	TRUE (default) or FALSE: print output

#### Details

estDLM

#### Value

A list with the following components:

conf.level	Specified confidence level
mod	a list of modifers with a vector of values from the model
n	Number of observation per specified subgroup

## exposureCov

groupIndex	list of index (row numbers) for specified subgroup
dlmMean	distributed lag effects per subgroups
dlmCI	credible intervals for distributed lag effects per subgroups
dlmCum	cumulative effects per subgroups
dlFunction	type of DLM class
plotData	data frame built for easier visualization of distributed lag effects for each sub- group (facet)

exposureCov	Exposure covariance structure	
-------------	-------------------------------	--

## Description

Matrix containing pairwise covariances for real exposure data consisting of five different exposures, each measured over 37 weeks.

#### Usage

data(exposureCov)

#### Format

matrix

#### Source

https://aqs.epa.gov/aqsweb/airdata/download\_files.html

## References

https://www.epa.gov/outdoor-air-quality-data

get_sbd_dlmtree	Download simulated data for dlmtree articles
-----------------	--

## Description

Download simulated data for dlmtree articles

## Usage

get\_sbd\_dlmtree()

A data frame with 10000 rows (observations) and 202 variables. All data is simulated. The variables are:

bwgaz	Outcome to be used. Simulated birth weight for gestational age z-score.	
ChildSex	Binary sex of child.	
MomAge	Continuous age in years.	
GestAge	Continuous estimated gestational age at birth in weeks.	
MomHeightIn	Continuous maternal height in inches.	
MomPriorWeight	_bs	
	Continuous mothers pre-pregnancy weight in pounds.	
MomPriorBMI	Continuous mothers pre-pregnancy BMI.	
race	Categorical race.	
Hispanic	Binary indicator of Hispanic.	
MomEdu	Categorical maternal highest educational attainment.	
SmkAny	Binary indicator of any smoking during pregnancy.	
Marital	Categorical maternal marital status.	
Income	Categorical income.	
EstDateConcept	Estimated date of conception.	
EstMonthConcept	t	
	Estimated month of conception.	
EstYearConcept	Estimated year of conception.	
pm25_1 - pm25_37		
	Weekly average exposure to PM2.5 for weeks 1 to 37. The columns are already scaled by the exposure IQR of 0.35.	
no2_1 - no2_37	Weekly average exposure to NO2 for weeks 1 to 37. The columns are already scaled by the exposure IQR of 9.13.	
so2_1 - so2_37	Weekly average exposure to SO2 for weeks 1 to 37. The columns are already scaled by the exposure IQR of 0.96.	
co2_1 - co2_37	Weekly average exposure to CO for weeks 1 to 37. The columns are already scaled by the exposure IQR of 0.15.	
temp_1 - temp_37	7	
	Weekly average exposure to temperature for weeks 1 to 37. The columns are already scaled by the exposure IQR of 27.93	
source	Variable indicating that the data came from the bdlim package.	

## Examples

sbd\_dlmtree <- get\_sbd\_dlmtree()</pre>

mixEst

## Description

Calculates the lagged interaction effects with MIX matrix for linear models.

## Usage

mixEst(dlm, nlags, nsamp)

## Arguments

dlm	A numeric matrix containing the model fit information
nlags	total number of lags
nsamp	number of mcmc iterations

#### Value

A cube object of interaction effect x lag x mcmc

monotdlnm\_Cpp *dlmtree model with monotone tdlnm approach* 

## Description

dlmtree model with monotone tdlnm approach

## Usage

```
monotdlnm_Cpp(model)
```

## Arguments

model A list of parameter and data contained for the model fitting

## Value

pip

## Description

Method for calculating posterior inclusion probabilities (PIPs) for modifiers in HDLM & HDLMM

## Usage

pip(object, type = 1)

## Arguments

object	An object of class dlmtree.
type	Type=1 indicates single modifier PIPs. Type=2 indicates joint modifier PIPs for two modifiers.

## Details

pip

## Value

numeric vector of PIPs named with modifiers (type=1) or data.frame of PIPs with the following columns (type=2):

var1	first modifier of joint modifiers
var2	second modifier of joint modifiers
pip	joint PIPs for the two modifiers

## Examples

plot.summary.monotone Returns variety of plots for model summary of class 'monotone'

#### Description

Method for returning variety of plots for model summary of class 'monotone'

## Usage

```
## S3 method for class 'summary.monotone'
plot(x, plot.type = "mean", val = c(), time = c(), ...)
```

## Arguments

x	object of class 'summary.monotone', output of summary of 'monotone'
plot.type	string indicating plot type, options are 'mean' (default) which shows mean exposure- time response surface, 'se', 'ci-min', 'ci-max', 'slice' which takes a slice of the plot at a given 'val' or 'time', 'animate' which creates a animation of slices of the surface plot across exposure values (requires package gganimate)
val	exposure value for slice plot
time	time value for slice plot
	additional parameters to alter plots: 'main', 'xlab', 'ylab', 'flab' which sets the effect label for surface plots, 'start.time' which sets the first time value

## Details

plot.summary.monotone

## Value

A plot of distributed lag effect estimated with monotone-TDLNM

plot.summary.tdlm Plots a distributed lag function for model summary of 'tdlm'

## Description

Method for plotting a distributed lag function for model summary of 'tdlm'

#### Usage

## S3 method for class 'summary.tdlm'
plot(x, ...)

## Arguments

х	object of class 'summary.tdlm', output of summary of 'tdlm'
	additional plotting parameters for title and labels 'start.time' which sets the first time value

#### Details

plot.summary.tdlm

#### Value

A plot of distributed lag effect estimated with tdlm

plot.summary.tdlmm Plots DLMMs for model summary of class 'tdlmm'

## Description

Method for plotting DLMMs for model summary of class 'tdlmm'. Includes plots for marginal exposure effects as well as interactions between two exposures.

#### Usage

```
## S3 method for class 'summary.tdlmm'
plot(
    x,
    type = "marginal",
    exposure1 = NULL,
    exposure2 = NULL,
    time1 = c(),
    time2 = c(),
    show.cw = TRUE,
    cw.plots.only = TRUE,
    trueDLM = NULL,
    scale = NULL,
    ...
)
```

## Arguments

х	an object of type 'summary.tdlmm' from summary.tdlmm() output
type	plot type, 'marginal' (default)
exposure1	exposure for plotting DLM
exposure2	exposure paired with 'exposure1' for plotting interaction
time1	plot a cross section from an interaction plot at specific time for 'exposure1'

## plot.summary.tdlnm

time2	plot a cross section from an interaction plot at specific time for 'exposure2'
show.cw	indicate location of critical windows in interaction plot with red points
cw.plots.only	show only plots with critical windows
trueDLM	A vector of true effects that can be obtained from the simulated data. Only applicable for simulation studies
scale	default = NULL, if scale is not NULL, the effects are exponentiated
	additional plotting parameters for title and labels

## Details

plot.summary.tdlmm

#### Value

A plot of distributed lag effect or interaction surface estimated with tdlmm

plot.summary.tdlnm *Returns variety of plots for model summary of class 'tdlnm'* 

## Description

Method for returning variety of plots for model summary of class 'tdlnm'

#### Usage

## S3 method for class 'summary.tdlnm'
plot(x, plot.type = "mean", val = c(), time = c(), ...)

## Arguments

х	object of class 'summary.tdlnm', output of summary of 'tdlnm'
plot.type	string indicating plot type, options are 'mean' (default) which shows mean exposure- time response surface, 'cumulative' which shows the cumulative effects per exposure-concentration level, 'effect' which returns a grid of exposure concen- tration and lag to determine if the credible interval contains zero, with the direc- tion of the effect indicated, 'se', 'ci-min', 'ci-max', 'slice' which takes a slice of the plot at a given 'val' or 'time', 'animate' which creates a animation of slices of the surface plot across exposure values (requires package gganimate)
val	exposure value for slice plot
time	time value for slice plot
	additional plotting parameters for title and labels 'flab' which sets the effect label for surface plots, 'start.time' which sets the first time value

#### Details

plot.summary.tdlnm

#### Value

A plot of distributed lag effect estimated with tdlnm

pm25Exposures PM2.5 Exposure data

#### Description

Data.frame containing a sample of weekly average PM2.5 exposures across a range of states/counties. The PM2.5 data was downloaded from US EPA (https://aqs.epa.gov/aqsweb/airdata/download\_files.html) daily data summaries and averaged by week. Forty-week ranges were assess for non-missingness and grouped for this dataset.

#### Usage

data(pm25Exposures)

## Format

data.frame; columns: S = state, C = city, 1-40 = weekly exposure data

#### Source

https://aqs.epa.gov/aqsweb/airdata/download\_files.html

#### References

https://www.epa.gov/outdoor-air-quality-data

ppRange

Makes a 'pretty' output of a group of numbers

#### Description

Method for making a 'pretty' output of a group of numbers. For example: 2,3,4,5,8,9,12,15,16 becomes 2-5,8-9,12,15-16

#### Usage

ppRange(r)

#### Arguments

r set of integers to make 'pretty'

## predict

## Details

ppRange

## Value

character string of values representing 'r'

predict

predict

## Description

predict generic function for S3method

## Usage

```
predict(
 х,
  new.data,
 new.exposure.data,
  ci.level = 0.95,
  type = "response",
  outcome = NULL,
  fixed.idx = list(),
 est.dlm = FALSE,
  verbose = TRUE,
  • • •
)
## S3 method for class 'hdlm'
predict(
  х,
 new.data,
 new.exposure.data,
 ci.level = 0.95,
  type = "response",
  outcome = NULL,
  fixed.idx = list(),
 est.dlm = FALSE,
  verbose = TRUE,
  . . .
)
## S3 method for class 'hdlmm'
predict(
  х,
 new.data,
```

predict

```
new.exposure.data,
ci.level = 0.95,
type = "response",
outcome = NULL,
fixed.idx = list(),
est.dlm = FALSE,
verbose = TRUE,
...
```

## Arguments

x	fitted dlmtree model with class 'hdlm', 'hdlmm'
new.data	new data frame which contains the same covariates and modifiers used to fit the model
new.exposure.d	ata
	new data frame/list which contains the same length of exposure lags used to fit the model
ci.level	credible interval level for posterior predictive distribution
type	type of prediction: "response" (default) or "waic". "waic" must be specified with 'outcome' parameter
outcome	outcome required for WAIC calculation
fixed.idx	fixed index
est.dlm	flag for estimating dlm effect
verbose	TRUE (default) or FALSE: print output
	not used

## Value

list with the following elements:

ztg posterior predictive mean of fixed effect

ztg.lims lower/upper bound of posterior predictive distribution of fixed effect

dlmest estimated exposure effect

dlmest.lower lower bound of estimated exposure effect

dlmest.upper upper bound of estimated exposure effect

fhat posterior predictive mean of exposure effect

fhat.lims lower/upper bound of posterior predictive distribution of exposure effect

y posterior predictive mean

y.lims lower/upper bound of posterior predictive distribution

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print

#### Description

print generic function for S3method

#### Usage

```
print(x, ...)
## S3 method for class 'tdlnm'
print(x, ...)
## S3 method for class 'tdlm'
print(x, ...)
## S3 method for class 'tdlmm'
print(x, ...)
## S3 method for class 'hdlm'
print(x, ...)
## S3 method for class 'hdlmm'
print(x, ...)
## S3 method for class 'monotone'
print(x, ...)
## S3 method for class 'summary.hdlm'
print(x, digits = 3, ...)
## S3 method for class 'summary.hdlmm'
print(x, digits = 3, ...)
## S3 method for class 'summary.monotone'
print(x, digits = 3, ...)
## S3 method for class 'summary.tdlm'
print(x, digits = 3, ...)
## S3 method for class 'summary.tdlmm'
print(x, digits = 3, ...)
## S3 method for class 'summary.tdlnm'
print(x, digits = 3, ...)
```

## Arguments

х	An object of class 'tdlm', 'tdlnm', 'tdlnm', 'hdlm', 'hdlmm', 'monotone', representing a fitted model using dlmtree(); or a summary object produced by applying summary() to one of these model objects.
	additional parameters
digits	number of decimal places to round the numeric values to

## Value

For a fitted model object, prints an assorted model output including model formula call and available methods. For a summary object, prints a summary output of a model fit in the R console.

rcpp\_pgdraw Multiple draw polya gamma latent variable for var c[i] with size b[i]

#### Description

Multiple draw polya gamma latent variable for var c[i] with size b[i]

## Usage

rcpp\_pgdraw(b, z)

## Arguments

b	vector of binomial sizes
z	vector of parameters

## Value

Eigen::VectorXd

rtmvnorm	Truncated multivariate normal sampler, mean mu, cov sigma, trun-
	cated (0, Inf)

## Description

Truncated multivariate normal sampler, mean mu, cov sigma, truncated (0, Inf)

## Usage

rtmvnorm(mu, sigma, iter)

## ruleIdx

## Arguments

mu	vector of mean parameters
sigma	covariance matrix
iter	number of iterations

## Value

VectorXd

ruleIdx Calculates the weights for each modifier rule	
---	--

## Description

Method for calculating the weights for each modifier rule

## Usage

ruleIdx(mod, mem.safe = FALSE)

## Arguments

mod	a list of modifier splitting rules
mem.safe	boolean memory parameter

## Value

A list of weights per rule with modifiers

scaleModelMatrix Centers and scales a matrix

## Description

Method for centering and scaling a matrix

## Usage

```
scaleModelMatrix(M)
```

## Arguments

M a matrix to center and scale

## Details

scaleModelMatrix

#### Value

a scaled matrix

shiny

shiny

## Description

shiny generic function for S3method

## Usage

shiny(fit)
## S3 method for class 'hdlm'

shiny(fit)

## S3 method for class 'hdlmm'
shiny(fit)

## Arguments

fit object of class 'hdlm', 'hdlmm' to which S3method is applied

## Value

shiny interface for further analysis on heterogeneous analyses. The interface includes tabs for modifier selection, personalized exposure effects and subgroup-specific effects.

sim.hdlmm

Creates simulated data for HDLM & HDLMM

## Description

Method for creating simulated data for HDLM & HDLMM

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#### sim.hdlmm

## Usage

```
sim.hdlmm(
   sim = "A",
   n = 1000,
   error = 1,
   effect.size = 1,
   exposure.data = NULL
)
```

## Arguments

sim	character (A - E) specifying simulation scenario
n	sample size
error	positive scalar specifying error variance for Gaussian response
effect.size	the effect size of the window of susceptibility
exposure.data	exposure data. A matrix of exposure data for simulation A, B, C and a named list of exposure data for simulation D, E

## Details

sim.hdlmm

Simulation scenarios:

- Scenario A: Two subgroups with early/late windows determined by continuous and binary modifiers
- Scenario B: Two subgroups with scaled effect determined by a continuous modifier
- Scenario C: No heterogeneity i.e., same effect on all individuals
- Scenario D: Three subgroups with three corresponding exposures. Subgroups are determined by continuous and binary modifiers
- Scenario E: Two subgroups with two exposures. First group is associated with the scaled main effect and lagged interaction while the second group is only associated with the scaled main effect, no interaction.

#### Value

Simulated data and true parameters

## Examples

sim.hdlmm(sim = "A", n = 1000)

sim.tdlmm

#### Description

Method for creating simulated data for TDLM & TDLMM

## Usage

```
sim.tdlmm(
    sim = "A",
    n = 5000,
    error = 10,
    mean.p = 0.5,
    prop.active = 0.05,
    expList = NULL,
    r = 1
)
```

#### Arguments

sim	character (A - F) specifying simulation scenario
n	sample size for simulation
error	positive scalar specifying error variance for Gaussian response
mean.p	scalar between zero and one specifying mean probability for simulation scenario A
prop.active	proportion of active exposures for simulation scenario C
expList	named list of exposure data
r	dispersion parameter of negative binomial distribution

## Details

sim.tdlmm

Simulation scenarios:

- Scenario A: Binary response with single exposure effect
- Scenario B: Continuous response with main effect of PM2.5 and interaction
- Scenario C: Continuous response to test exposure selection using exposure
- Scenario D: Continuous response to test exposure selection using one exposure of main effect and two interaction effects among four exposures
- Scenario E: Zero-inflated count response with single exposure effect
- Scenario F: Zero-inflated count response with single exposure effect with main effect of PM2.5 and interaction

## sim.tdlnm

## Value

Simulated data and true parameters

#### Examples

sim.tdlmm(sim = "A", mean.p = 0.5, n = 1000)

sim.tdlnm

Creates simulated data for TDLNM

## Description

Method for creating simulated data for TDLNM

## Usage

sim.tdlnm(sim = "A", error.to.signal = 1)

#### Arguments

sim character (A - D) specifying simulation scenario error.to.signal scalar value setting error: sigma^2/var(f)

## Details

sim.tdlnm

Simulation scenarios:

- Scenario A: Piecewise constant effect
- Scenario B: Linear effect
- Scenario C: Logistic effect, piecewise in time
- Scenario D: Logistic effect, smooth in time

## Value

Simulated data and true parameters

## Examples

sim.tdlnm(sim = "A", error.to.signal = 1)

splitPIP

## Description

Calculates the posterior inclusion probability (PIP).

## Usage

```
splitPIP(dlnm, nlags, niter)
```

## Arguments

dlnm	A numeric matrix containing the model fit information
nlags	total number of lags
niter	number of mcmc iterations

## Value

A matrix of split counts per mcmc

splitpoints	Determines split points for continuous modifiers
-------------	--

## Description

Method for determining split points for continuous modifiers

## Usage

```
splitpoints(object, var, round = NULL)
```

## Arguments

object	An object of class 'hdlm', 'hdlmm'	
var	The name of a continuous variable for which the split points will be reported	
round	The number of decimal places to round the variable (var) to. No rounding occurs if round=NULL (default) For positive integer values of round, the variable will be rounded and split points will be reported at the resulting level	

## Details

splitpoints

#### summary

#### Value

A data frame with split points and the probability that a split point was >= that split point value

#### Examples

summary

summary

#### Description

summary generic function for S3method

#### Usage

```
summary(x, conf.level = 0.95, ...)
## S3 method for class 'hdlm'
summary(x, conf.level = 0.95, mcmc = FALSE, ...)
## S3 method for class 'hdlmm'
summary(x, conf.level = 0.95, mcmc = FALSE, ...)
## S3 method for class 'monotone'
summary(
 х,
  conf.level = 0.95,
  pred.at = NULL,
  cenval = 0,
  exposure.se = NULL,
 mcmc = FALSE,
  verbose = TRUE,
  . . .
)
## S3 method for class 'tdlm'
```

## summary

```
summary(x, conf.level = 0.95, mcmc = FALSE, ...)
## S3 method for class 'tdlmm'
summary(
 х,
 conf.level = 0.95,
 marginalize = "mean",
 log10BF.crit = 0.5,
 mcmc = FALSE,
 verbose = TRUE,
  . . .
)
## S3 method for class 'tdlnm'
summary(
 х,
 conf.level = 0.95,
 pred.at = NULL,
 cenval = 0,
 exposure.se = NULL,
 mcmc = FALSE,
 verbose = TRUE,
  . . .
)
```

## Arguments

х	an object of class 'tdlm', 'tdlmm', 'tdlnm', 'hdlm', 'hdlmm', 'monotone'	
conf.level	confidence level for computation of credible intervals	
	additional parameters	
mcmc	keep all mcmc iterations (large memory requirement)	
pred.at	numerical vector of exposure values to make predictions for at each time period	
cenval	scalar exposure value that acts as a reference point for predictions at all other exposure values	
exposure.se	scalar smoothing factor, if different from model	
verbose	show progress in console	
marginalize	value(s) for calculating marginal DLMs, defaults to "mean", can also specify a percentile from 1-99 for all other exposures, or a named vector with specific values for each exposure	
log10BF.crit	Bayes Factor criteria for selecting exposures and interactions, such that $log10$ (BayesFactor) > x. Default = 0.5.	

## Value

list of summary outputs of the model fit

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tdlmm\_Cpp

## Description

dlmtree model with tdlmm approach

## Usage

tdlmm\_Cpp(model)

## Arguments

model A list of parameter and data contained for the model fitting

#### Value

A list of dlmtree model fit, mainly posterior mcmc samples

tdlnm\_Cpp dlmtree model with tdlnm approach

## Description

dlmtree model with tdlnm approach

#### Usage

tdlnm\_Cpp(model)

## Arguments

model A list of parameter and data contained for the model fitting

## Value

zeroToInfNormCDF

#### Description

Integrates (0,inf) over multivariate normal

## Usage

zeroToInfNormCDF(mu, sigma)

## Arguments

mu	vector of mean parameters
sigma	covariance matrix

## Value

double

zinbCo

Time-series exposure data for ZINB simulated data

#### Description

Data.frame containing a sample of weekly average wildfire PM, PM2.5, O3 across a range of counties of Colorado. The exposure data was downloaded from US EPA (https://aqs.epa.gov/aqsweb/airdata/download\_files.html) daily data summaries and averaged by week.

## Usage

data(zinbCo)

#### Format

data.frame;

#### Source

https://aqs.epa.gov/aqsweb/airdata/download\_files.html

#### References

https://www.epa.gov/outdoor-air-quality-data

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