# Package 'bipd'

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Title Bayesian Individual Patient Data Meta-Analysis using 'JAGS'

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Description  We use a Bayesian approach to run individual patient data meta-analysis and network meta-analysis using 'JAGS'. The methods incorporate shrinkage methods and calculate patient-specific treatment effects as described in Seo et al. (2021) <doi:10.1002 sim.8859="">. This package also includes user-friendly functions that impute missing data in an individual patient data using mice-related packages.</doi:10.1002>
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## **Description**

A package for individual patient data meta-analysis using 'JAGS'

#### **Details**

We use a Bayesian approach to run individual patient data meta-analysis and network meta-analysis using 'JAGS'. The methods incorporate shrinkage methods and calculate patient-specific treatment effects as described in Seo et al. (2021) <DOI:10.1002/sim.8859>. This package also includes user-friendly functions that impute missing data in an individual patient data using mice-related packages.

#### References

Audigier V, White I, Jolani S, et al. Multiple Imputation for Multilevel Data with Continuous and Binary Variables. *Statistical Science*. 2018;33(2):160-183. doi: 10.1214/18STS646

Debray TPA, Moons KGM, Valkenhoef G, et al. Get real in individual participant data (IPD) meta-analysis: a review of the methodology. *Res Synth Methods*. 2015;6(4):293-309. doi: 10.1002/jrsm.1160

Dias S, Sutton AJ, Ades AE, et al. A Generalized Linear Modeling Framework for Pairwise and Network Meta-analysis of Randomized Controlled Trials. *Medical Decision Making*. 2013;33(5):607-617. doi: 10.1177/0272989X12458724

Fisher DJ, Carpenter JR, Morris TP, et al. Meta-analytical methods to identify who benefits most from treatments: daft, deluded, or deft approach? *BMJ*. 2017;356:j573. doi: 10.1136/bmj.j573

O'Hara RB, Sillanpaa MJ. A review of Bayesian variable selection methods: what, how and which. *Bayesian Anal.* 2009;4(1):85-117. doi: 10.1214/09BA403

Riley RD, Debray TP, Fisher D, et al. Individual participant data meta-analysis to examine interactions between treatment effect and participant-level covariates: Statistical recommendations for conduct and planning. *Stat Med.* 2020:39(15):2115-2137. doi: 10.1002/sim.8516

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Seo M, White IR, Furukawa TA, et al. Comparing methods for estimating patient-specific treatment effects in individual patient data meta-analysis. *Stat Med.* 2021;40(6):1553-1573. doi: 10.1002/sim.8859

add.mcmc

Convenient function to add results (i.e. combine mcmc.list)

## **Description**

This is a convenient function to add results (i.e. combine mcmc.list). This can be useful when combining results obtained from multiple imputation

## Usage

```
add.mcmc(x, y)
```

#### **Arguments**

x first result in a format of mcmc.list

y second result in a format of mcmc.list

```
ds <- generate_ipdma_example(type = "continuous")
ds2 <- generate_ipdma_example(type = "continuous")
ipd <- with(ds, ipdma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))
ipd2 <- with(ds2, ipdma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))

samples <- ipd.run(ipd, pars.save = c("beta", "gamma", "delta"), n.chains = 3, n.burnin = 500,
n.iter = 5000)
samples2 <- ipd.run(ipd2, pars.save = c("beta", "gamma", "delta"), n.chains = 3, n.burnin = 500,
n.iter = 5000)
combined <- add.mcmc(samples, samples2)</pre>
```

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findMissingPattern

Find missing data pattern in a given data

#### **Description**

Find missing data pattern in a given data i.e. whether variables are systematically missing or sporadically missing. Also calculates missing count and percentage for exploratory purposes.

## Usage

```
findMissingPattern(
  dataset = NULL,
  covariates = NULL,
  typeofvar = NULL,
  studyname = NULL,
  treatmentname = NULL,
  outcomename = NULL)
```

## **Arguments**

dataset data which contains variables of interests

covariates vector of variable names that the user is interested in finding a missing data

pattern

type of covariate variables; should be a vector of these values: "continuous",

"binary", or "count". Order should follow that of covariates parameter.

studyname study name in the data specified treatment name in the data specified outcomename outcome name in the data specified

## Value

missing number of patients for each study and covariate
missingpercent missing percentage of patients for each study and covariate

sys\_missing a vector indicating whether each covariate is systematically missing spor\_missing a vector indicating whether each covariate is sporadically missing

sys\_covariates a vector of systematically missing covariates

spor\_covariates

a vector of sporadically missing covariates

without\_sys\_covariates

a vector of covariates that are not systematically missing

covariates vector of variable names that the user is interested in finding a missing data

pattern

study name in the data specified
treatment name in the data specified
outcomename outcome name in the data specified

## **Examples**

```
simulated_dataset <- generate_sysmiss_ipdma_example(Nstudies = 10, Ncov = 5, sys_missing_prob = 0.3,
magnitude = 0.2, heterogeneity = 0.1)

missP <- findMissingPattern(simulated_dataset, covariates = c("x1", "x2", "x3", "x4", "x5"),
typeofvar = c("continuous", "binary", "continuous", "continuous"), studyname = "study",
treatmentname = "treat", outcomename = "y")
missP</pre>
```

generate\_ipdma\_example

Generate a simulated IPD-MA data for demonstration

## **Description**

Generate a simulated IPD-MA data for demonstration

## Usage

```
generate_ipdma_example(type = "continuous")
```

#### **Arguments**

type

"continuous" for continuous outcome and "binary" for binary outcome

#### Value

returns simulated IPD-MA data

#### **Examples**

```
ds <- generate_ipdma_example(type = "continuous")
head(ds)</pre>
```

generate\_ipdnma\_example

Generate a simualted IPD-NMA data for demonstration

## **Description**

Generate a simulated IPD-NMA data for demonstration

```
generate_ipdnma_example(type = "continuous")
```

#### **Arguments**

type

"continuous" for continuous outcome and "binary" for binary outcome

#### Value

return simulated IPD-NMA data ds <- generate\_ipdnma\_example(type = "continuous") head(ds)

```
generate_sysmiss_ipdma_example
```

Generate a simulated IPD-MA data with systematically missing covariates

## **Description**

Generate a simulated IPD-MA data with systematically missing covariates

## Usage

```
generate_sysmiss_ipdma_example(
  Nstudies = 10,
  Ncov = 5,
  sys_missing_prob = 0.1,
  magnitude = 0.3,
  heterogeneity = 0.1,
  interaction = TRUE
)
```

#### **Arguments**

Nstudies number of studies. Default is 10.

Ncov number of covariates in total. Options are 5 or 10 studies. Default is set to 5.

sys\_missing\_prob

probability of systematically missing studies for each covariates. Default is set

to 0.3.

magnitude magnitude of the regression estimates (the mean). Default is set to 0.2. heterogeneity heterogeneity of regression estimates across studies. Default is set to 0.1.

interaction whether to include treatment indicator and treatment

## Value

returns simulated IPD-MA data with systematically missing covariates

```
simulated_dataset <- generate_sysmiss_ipdma_example(Nstudies = 10, Ncov = 5, sys_missing_prob = 0.3,
magnitude = 0.2, heterogeneity = 0.1)
head(simulated_dataset)</pre>
```

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ipd.run

Run the model using the ipd object

## **Description**

This is the core function that runs the model in our program. Before running this function, we need to specify data, prior, JAGS code, etc. using ipd.model type function.

## Usage

```
ipd.run(
   ipd,
   pars.save = NULL,
   inits = NULL,
   n.chains = 3,
   n.adapt = 1000,
   n.burnin = 1000,
   n.iter = 10000
)
```

## **Arguments**

ipd	ipd object created from ipd.model type function
pars.save	parameters to save. For instance, "beta" - coefficients for main effects; "gamma" - coefficients for effect modifiers; "delta" - average treatment effect
inits	initial values specified for the parameters to save
n.chains	number of MCMC chains to sample
n.adapt	number of iterations for adaptation (Note that the samples from adaptation phase is non-Markovian and do not constitute a Markov chain)
n.burnin	number of iterations for burn-in
n.iter	number of iterations to run after the adaptation

## Value

MCMC samples stored using JAGS. The returned samples have the form of mcmc.list and coda functions can be directly applied.

```
ds <- generate_ipdma_example(type = "continuous")
ipd <- with(ds, ipdma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))
samples <- ipd.run(ipd, n.chains = 3, n.burnin = 500, n.iter = 5000)</pre>
```

8 ipd.run.parallel

ipd.run.parallel

Run the model using the ipd object with parallel computation

## **Description**

This function runs the model through parallel computation using dclone R package. Before running this function, we need to specify data, prior, JAGS code, etc. using ipd.model type function.

## Usage

```
ipd.run.parallel(
   ipd,
   pars.save = NULL,
   inits = NULL,
   n.chains = 2,
   n.adapt = 1000,
   n.burnin = 1000,
   n.iter = 10000
)
```

## **Arguments**

ipd	ipd object created from ipd.model type function
pars.save	parameters to save. For instance, "beta" - coefficients for main effects; "gamma" - coefficients for effect modifiers; "delta" - average treatment effect
inits	initial values specified for the parameters to save
n.chains	number of MCMC chains to sample
n.adapt	number of iterations for adaptation (Note that the samples from adaptation phase is non-Markovian and do not constitute a Markov chain)
n.burnin	number of iterations for burn-in
n.iter	number of iterations to run after the adaptation

#### Value

MCMC samples stored using JAGS. The returned samples have the form of mcmc.list and coda functions can be directly applied.

```
ds <- generate_ipdma_example(type = "continuous")
ipd <- with(ds, ipdma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))
samples <- ipd.run.parallel(ipd, n.chains = 2, n.burnin = 500, n.iter = 5000)</pre>
```

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ipdma.impute Impute missing data in individual participant data with two treatments (i.e. placebo and a treatment).

## **Description**

Impute missing data in individual participant data with two treatments. Data is clustered by different studies. In the presence of systematically missing variables, the function defaults to 21.2stage.norm, 21.2stage.bin, and 21.2stage.pois methods in micemd package. If there are no systematically missing variables, the function defaults to use 21.pmm in miceadds package which generalizes predictive mean matching using linear mixed model. If there is only one study available, the function defaults to use pmm in mice package.

## Usage

```
ipdma.impute(
  dataset = NULL,
  covariates = NULL,
  typeofvar = NULL,
  sys_impute_method = "21.2stage",
  interaction = NULL,
  meth = NULL,
  pred = NULL,
  studyname = NULL,
  treatmentname = NULL,
  outcomename = NULL,
  m = 5
)
```

#### **Arguments**

dataset data which contains variables of interests

covariates vector of variable names to find missing data pattern

typeofvar type of covariate variables; should be a vector of these values: "continuous",

"binary", or "count". Order should follow that of covariates parameter specified.

Covariates that are specified "binary" are automatically factored.

sys\_impute\_method

method used for systematically missing studies. Options are "21.glm", "21.2stage", or "21.jomo". Default is set to "21.2stage". There is also an option to ignore all the clustering level and impute using predictive mean matching by setting this

parameter to "pmm".

interaction indicator denoting whether treatment-covariate interactions should be included.

Default is set to true.

meth imputation method to be used in the mice package. If left unspecified, function

picks a reasonable one.

pred correct prediction matrix to be used in the mice package. If left unspecified,

function picks a reasonable one.

study name in the data specified.

treatment name in the data specified.

outcome name outcome name in the data specified.

m number of imputed datasets. Default is set to 5.

#### Value

missingPattern missing pattern object returned by running findMissingPattern function

meth imputation method used with the mice function pred prediction matrix used with the mice function

imp imputed datasets that is returned from the mice function

imp.list imputed datasets in a list format

#### **Examples**

```
simulated_dataset <- generate_sysmiss_ipdma_example(Nstudies = 10, Ncov = 5, sys_missing_prob = 0.3,
magnitude = 0.2, heterogeneity = 0.1)

# load in mice packages
library(mice) #for datasets with only one study level
library(miceadds) #for multilevel datasets without systematically missing predictors
library(micemd) #for multilevel datasets with systematically missing predictors.
imputation <- ipdma.impute(simulated_dataset, covariates = c("x1", "x2", "x3", "x4", "x5"),
typeofvar = c("continuous", "binary", "binary", "continuous", "continuous"), interaction = TRUE,
studyname = "study", treatmentname = "treat", outcomename = "y", m = 5)</pre>
```

ipdma.model.deft.onestage

Make a (deft-approach) one-stage individual patient data metaanalysis object containing data, priors, and a JAGS model code

## **Description**

This function sets up data and JAGS code that is needed to run (deft-approach) one-stage IPD-MA models in JAGS.

```
ipdma.model.deft.onestage(
  y = NULL,
  study = NULL,
  treat = NULL,
  X = NULL,
```

```
response = "normal",
  type = "random",
  mean.alpha = 0,
  prec.alpha = 0.001,
  mean.beta = 0,
  prec.beta = 0.001,
  mean.gamma.within = 0,
  prec.gamma.within = 0.001,
  mean.gamma.across = 0,
  prec.gamma.across = 0,
  prec.gamma.across = 0.001,
  mean.delta = 0,
  prec.delta = 0.001,
  hy.prior = list("dhnorm", 0, 1)
)
```

## Arguments

у	outcome of the study. Can be continuous or binary.	
study	vector indicating which study the patient belongs to. Please change the study names into numbers (i.e. 1, 2, 3, etc)	
treat	vector indicating which treatment the patient was assigned to (i.e. $1$ for treatment, $0$ for placebo)	
X	matrix of covariate values for each patient. Dimension would be number of patients x number of covariates.	
response	specification of the outcome type. Must specify either "normal" or "binomial".	
type	assumption on the treatment effect: either "random" for random effects model or "fixed" for fixed effects model. Default is "random".	
mean.alpha	prior mean for the study intercept	
prec.alpha	prior precision for the study intercept	
mean.beta	prior mean for the regression coefficients of the main effects of the covariates; main effects are assumed to have common effect.	
prec.beta	prior precision for the regression coefficients of the main effects of the covariates	
mean.gamma.with	nin	
	prior mean for effect modifiers of within study information.	
prec.gamma.with	nin	
	prior precision for the effect modifiers of within study information.	
mean.gamma.across		
	prior mean for the effect modifiers of across study information; effect modification is assumed to have common effect.	
prec.gamma.across		
	prior precision for the effect modifiers of across study information	
mean.delta	prior mean for the average treatment effect	
prec.delta	prior precision for the average treatment effect	

		٠		
hy	nr	1	O	r

prior for the heterogeneity parameter. Supports uniform, gamma, and half normal for normal and binomial response It should be a list of length 3, where first element should be the distribution (one of dunif, dgamma, dhnorm) and the next two are the parameters associated with the distribution. For example, list("dunif", 0, 5) gives uniform prior with lower bound 0 and upper bound 5 for the heterogeneity parameter.

#### Value

data. JAGS data organized in a list so that it can be used when running code in JAGS

code JAGS code that is used to run the model. Use cat(code) to see the code in a readable format

model. JAGS JAGS code in a function. This is used when running model in parallel study specific averages of covariates

## References

Fisher DJ, Carpenter JR, Morris TP, et al. Meta-analytical methods to identify who benefits most from treatments: daft, deluded, or deft approach?. *BMJ*. 2017;356:j573 doi: 10.1136/bmj.j573

## **Examples**

```
ds <- generate_ipdma_example(type = "continuous")
ipd <- with(ds, ipdma.model.deft.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal"))
samples <- ipd.run(ipd)
treatment.effect(ipd, samples, newpatient= c(1,0.5), reference = c(0, 0))</pre>
```

ipdma.model.onestage Make an one-stage individual patient data meta-analysis object containing data, priors, and a JAGS model code

## **Description**

This function sets up data and JAGS code that is needed to run one-stage IPD-MA models in JAGS.

```
ipdma.model.onestage(
  y = NULL,
  study = NULL,
  treat = NULL,
  X = NULL,
  response = "normal",
  type = "random",
```

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```
shrinkage = "none",
scale = TRUE,
mean.alpha = 0,
prec.alpha = 0.001,
mean.beta = 0,
prec.beta = 0.001,
mean.gamma = 0,
prec.gamma = 0.001,
mean.delta = 0,
prec.delta = 0.001,
hy.prior = list("dhnorm", 0, 1),
lambda.prior = NULL,
p.ind = NULL,
hy.prior.eta = NULL
```

#### **Arguments**

prec.gamma

у	outcome of the study. Can be continuous or binary.
study	vector indicating which study the patient belongs to. Please change the study names into numbers (i.e. 1, 2, 3, etc)
treat	vector indicating which treatment the patient was assigned to (i.e. 1 for treatment, 0 for placebo)
X	matrix of covariate values for each patient. Dimension would be number of patients x number of covariates.
response	specification of the outcome type. Must specify either "normal" or "binomial".
type	assumption on the treatment effect: either "random" for random effects model or "fixed" for fixed effects model. Default is "random".
shrinkage	shrinkage method applied to the effect modifiers. "none" correspond to no shrinkage. "laplace" corresponds to a adaptive shrinkage with a Laplacian prior (ie often known as Bayesian LASSO). "SSVS" corresponds to the Stochastic Search Variable Selection method. SSVS is not strictly a shrinkage method, but pulls the estimated coefficient toward zero through variable selection in each iteration of the MCMC. See O'hara et al (2009) for more details.
scale	indicator for scaling the covariates by the overall average; default is TRUE.
mean.alpha	prior mean for the study intercept
prec.alpha	prior precision for the study intercept
mean.beta	prior mean for the regression coefficients of the main effects of the covariates; main effects are assumed to have common effect.
prec.beta	prior precision for the regression coefficients of the main effects of the covariates
mean.gamma	prior mean for the effect modifiers. This parameter is not used if penalization is

prior precision for the effect modifiers. This parameter is not used if penalization

placed on effect modifiers.

is placed on effect modifiers.

mean.delta	prior mean for the average treatment effect
prec.delta	prior precision for the average treatment effect
hy.prior	prior for the heterogeneity parameter. Supports uniform, gamma, and half normal for normal and binomial response It should be a list of length 3, where first element should be the distribution (one of dunif, dgamma, dhnorm) and the next two are the parameters associated with the distribution. For example, list("dunif", 0, 5) gives uniform prior with lower bound 0 and upper bound 5 for the heterogeneity parameter.
lambda.prior	(only for shrinkage = "laplace") two options for laplace shrinkage. We can put a gamma prior on the lambda (i.e. list("dgamma",2,0.1)) or put a uniform prior on the inverse of lambda (i.e. list("dunif",0,5))
p.ind	(only for shrinkage = "SSVS") prior probability of including each of the effect modifiers. Length should be same as the total length of the covariates.
g	(only for shrinkage = "SSVS") multiplier for the precision of spike. Default is $g = 1000$ .
hy.prior.eta	(only for shrinkage = "SSVS") standard deviation of the slab prior. Currently only support uniform distribution. Default is list("dunif", $0, 5$ )

#### Value

data.JAGS	data organized in a list so that it can be used when running code in JAGS
code	JAGS code that is used to run the model. Use cat(code) to see the code in a readable format
model.JAGS	JAGS code in a function. This is used when running model in parallel
scale.mean	mean used in scaling covariates
scale.sd	standard deviation used in scaling covariates

## References

O'Hara RB, Sillanpaa MJ. A review of Bayesian variable selection methods: what, how and which. *Bayesian Anal.* 2009;4(1):85-117. doi: 10.1214/09BA403

Seo M, White IR, Furukawa TA, et al. Comparing methods for estimating patient-specific treatment effects in individual patient data meta-analysis. *Stat Med.* 2021;40(6):1553-1573. doi: 10.1002/sim.8859

```
ds <- generate_ipdma_example(type = "continuous")
ipd <- with(ds, ipdma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))
samples <- ipd.run(ipd)
treatment.effect(ipd, samples, newpatient= c(1,0.5))</pre>
```

ipdnma.model.onestage Make an one-stage individual patient data network meta-analysis object containing data, priors, and a JAGS model code

## **Description**

This function sets up data and JAGS code that is needed to run one-stage IPD-NMA models in

#### Usage

```
ipdnma.model.onestage(
  y = NULL,
  study = NULL,
  treat = NULL,
  X = NULL,
  response = "normal",
  type = "random",
  shrinkage = "none",
  scale = TRUE,
 mean.alpha = 0,
  prec.alpha = 0.001,
 mean.beta = 0,
 prec.beta = 0.001,
 mean.gamma = 0,
  prec.gamma = 0.001,
 mean.delta = 0,
  prec.delta = 0.001,
  hy.prior = list("dhnorm", 0, 1),
  lambda.prior = NULL,
  p.ind = NULL,
  g = NULL,
 hy.prior.eta = NULL
)
```

#### **Arguments**

treat

outcome of the study. Can be continuous or binary. У

vector indicating which study the patient belongs to. Please change the study study

names into numbers (i.e. 1, 2, 3, etc)

vector indicating which treatment the patient was assigned to. Since this is a network meta-analysis and there would be more than 2 treatments, careful naming of treatment is needed. This vector needs to be a sequence from 1:NT where NT is the total number of treatments. Treatment that is assigned 1 would be the

baseline treatment.

Χ matrix of covariate values for each patient. Dimension would be number of patients x number of covariates. specification of the outcome type. Must specify either "normal" or "binomial". response assumption on the treatment effect: either "random" for random effects model type or "fixed" for fixed effects model. Default is "random". shrinkage method applied to the effect modifiers. "none" correspond to no shrinkage shrinkage. "laplace" corresponds to a adaptive shrinkage with a Laplacian prior (ie often known as Bayesian LASSO). "SSVS" corresponds to the Stochastic Search Variable Selection method. SSVS is not strictly a shrinkage method, but pulls the estimated coefficient toward zero through variable selection in each iteration of the MCMC. See O'hara et al (2009) for more details. scale indicator for scaling the covariates by the overall average; default is TRUE. mean.alpha prior mean for the study intercept prec.alpha prior precision for the study intercept mean.beta prior mean for the regression coefficients of the main effects of the covariates; main effects are assumed to have common effect. prior precision for the regression coefficients of the main effects of the covariates prec.beta prior mean for the effect modifiers. This parameter is not used if penalization is mean.gamma placed on effect modifiers. prec.gamma prior precision for the effect modifiers. This parameter is not used if penalization is placed on effect modifiers. mean.delta prior mean for the average treatment effect prior precision for the average treatment effect prec.delta hy.prior prior for the heterogeneity parameter. Supports uniform, gamma, and half normal for normal and binomial response It should be a list of length 3, where first element should be the distribution (one of dunif, dgamma, dhnorm) and the next two are the parameters associated with the distribution. For example, list("dunif", 0, 5) gives uniform prior with lower bound 0 and upper bound 5 for the heterogeneity parameter. (only for shrinkage = "laplace") two options for laplace shrinkage. We can put lambda.prior a gamma prior on the lambda (i.e. list("dgamma",2,0.1)) or put a uniform prior on the inverse of lambda (i.e. list("dunif",0,5)) p.ind (only for shrinkage = "SSVS") prior probability of including each of the effect modifiers. Length should be same as the total length of the covariates. (only for shrinkage = "SSVS") multiplier for the precision of spike. Default is g g = 1000.hy.prior.eta (only for shrinkage = "SSVS") standard deviation of the slab prior. Currently

#### Value

data. JAGS data organized in a list so that it can be used when running code in JAGS code

JAGS code that is used to run the model. Use cat(code) to see the code in a readable format

only support uniform distribution. Default is list("dunif", 0, 5)

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model.JAGS	JAGS code in a function. This is used when running model in parallel
scale.mean	mean used in scaling covariates
scale.sd	standard deviation used in scaling covariates

#### References

Dias S, Sutton AJ, Ades AE, et al. A Generalized Linear Modeling Framework for Pairwise and Network Meta-analysis of Randomized Controlled Trials. *Medical Decision Making*. 2013;33(5):607-617. doi: 10.1177/0272989X12458724

O'Hara RB, Sillanpaa MJ. A review of Bayesian variable selection methods: what, how and which. *Bayesian Anal.* 2009;4(1):85-117. doi: 10.1214/09BA403

Seo M, White IR, Furukawa TA, et al. Comparing methods for estimating patient-specific treatment effects in individual patient data meta-analysis. *Stat Med.* 2021;40(6):1553-1573. doi: 10.1002/sim.8859

#### **Examples**

```
ds <- generate_ipdnma_example(type = "continuous")
ipd <- with(ds, ipdnma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))
samples <- ipd.run(ipd)
treatment.effect(ipd, samples, newpatient= c(1,0.5))</pre>
```

treatment.effect

Calculate patient-specific treatment effect

## Description

Function for calculating the patient-specific treatment effect. Patient-specific treatment effect includes the main effect of treatment and treatment-covariate interaction effect (i.e. effect modification). Reports odds ratio for the binary outcome.

```
treatment.effect(
  ipd = NULL,
  samples = NULL,
  newpatient = NULL,
  scale_mean = NULL,
  scale_sd = NULL,
  reference = NULL,
  quantiles = c(0.025, 0.5, 0.975)
)
```

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## Arguments

ipd	IPD object created from running ipdma.model type function
samples	MCMC samples found from running ipd.run function
newpatient	covariate values of patients that you want to predict treatment effect on. Must have length equal to total number of covariates.
scale_mean	option to specify different overall mean compared to what was calculated in IPD object. can be useful when using multiple imputation.
scale_sd	option to specify different overall standard deviation compared to what was calculated in IPD object.
reference	reference group used for finding patient-specific treatment effect. This is only used for deft approach
quantiles	quantiles for credible interval of the patient-specific treatment effect

#### Value

patient-specific treatment effect with credible interval at specified quantiles

#### References

Seo M, White IR, Furukawa TA, et al. Comparing methods for estimating patient-specific treatment effects in individual patient data meta-analysis. *Stat Med.* 2021;40(6):1553-1573. doi: 10.1002/sim.8859

Riley RD, Debray TP, Fisher D, et al. Individual participant data meta-analysis to examine interactions between treatment effect and participant-level covariates: Statistical recommendations for conduct and planning. *Stat Med.* 2020:39(15):2115-2137. doi: 10.1002/sim.8516

```
ds <- generate_ipdma_example(type = "continuous")
ipd <- with(ds, ipdma.model.onestage(y = y, study = studyid, treat = treat, X = cbind(z1, z2),
response = "normal", shrinkage = "none"))
samples <- ipd.run(ipd, pars.save = c("beta", "gamma", "delta"), n.chains = 3, n.burnin = 500,
n.iter = 5000)
treatment.effect(ipd, samples, newpatient = c(1,0.5))</pre>
```

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