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BeviMed-package

Bayesian Evaluation of Variant Involvement in Mendelian Disease

Description

A fast integrative genetic association test for rare diseases.

Details

BeviMed estimates a probability of association between a case/control label and allele counts at rare variant sites in a genomic locus and also, given that there is an association, the probabilities that each variant is involved in the disease. It does so by estimating the evidence for a model where the case/control label is independent of the allele configurations, and a model in which the probability of the case/control label depends on the corresponding allele configuration and a latent partition of variants into pathogenic and non-pathogenic groups.

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Author(s)

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References

Greene et al., A Fast Association Test for Identifying Pathogenic Variants Involved in Rare Diseases, The American Journal of Human Genetics (2017), http://dx.doi.org/10.1016/j.ajhg.2017.05.015.

See Also

bevimed

bevimed

Bayesian Evaluation of Variant Involvement in Mendelian Disease

Description

Infer probabilities of association between disease label and locus and posterior parameter values under BeviMed model.

Usage

```
bevimed(y, G, ploidy = rep(2L, length(y)),
   prior_prob_association = 0.01, prior_prob_dominant = 0.5,
   dominant_args = NULL, recessive_args = NULL, ...)
```

Arguments

y Logical vector of case (TRUE) control (FALSE) status.

G Integer matrix of variant counts per individual, one row per individual and one

column per variant.

ploidy Integer vector giving ploidy of samples.

prior_prob_association

The prior probability of association.

prior_prob_dominant

The prior probability of dominant inheritance given that there is an association.

dominant_args Arguments to pass to bevimed_m conditioning on dominant inheritance.

recessive_args Arguments to pass to bevimed_m conditioning on recessive inheritance.

... Arguments to be passed to bevimed_m for both modes of inheritance.

Value

BeviMed object containing results of inference.

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References

Greene et al., A Fast Association Test for Identifying Pathogenic Variants Involved in Rare Diseases, The American Journal of Human Genetics (2017), http://dx.doi.org/10.1016/j.ajhg.2017.05.015.

See Also

prob_association, bevimed_m, summary.BeviMed, bevimed_polytomous

Perform inference under model gamma = 1 conditional on mode of bevimed_m inheritance

Description

Sample from posterior distribution of parameters under model gamma = 1 and conditional on mode of inheritance, set via the min_ac argument.

Usage

```
bevimed_m(y, G, min_ac = 1L, tau_shape = c(1, 1), pi_shape = c(6, 1),
 omega_shape = if (max(min_ac) == 1L) c(2, 8) else c(2, 2),
 samples_per_chain = 1000, stop_early = FALSE, blocks = 5,
 burn = as.integer(samples_per_chain/10), temperatures = (0:6/6)^2,
 tune_temps = 0, return_z_trace = TRUE, return_x_trace = TRUE,
 raw_only = FALSE, swaps = as.integer(length(temperatures)/2),
 optimise_z0 = FALSE, tune_omega_and_phi_proposal_sd = FALSE,
 tune_block_size = 100, variant_weights = NULL,
 standardise_weights = TRUE, log_phi_mean = -0.15,
 log_phi_sd = sqrt(0.3), tandem_variant_updates = if (max(min_ac) ==
 1) 0 else min(sum(y), ncol(G)), ...)
```

Arguments

у	Logical	vector	of case	(TRUE)) control	(FALSE)	status.
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G Integer matrix of variant counts per individual, one row per individual and one

column per variant.

min_ac Integer vector with a length equalling the number of individuals or length 1 (in

> which case the given value is used for all individuals) giving the minimum number of alleles at pathogenic variant sites each individual requires in order to classify as having a 'pathogenic allele configuration'. Thus, this parameter encodes the mode of inheritance. For instance, setting this parameter to 1 corresponds to dominant inheritance. If there are differences in ploidy between individuals in the locus, it is necessary to set it on an sample level basis - e.g. to ensure sex is

accounted for if the locus lies on the X chromosome.

tau_shape Beta shape hyper-priors for prior on rate of affection (i.e. being a case) amongst

individuals with non-pathogenic variant combinations (i.e. they have less than

min_ac variants.

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pi_shape Beta shape hyper-priors for prior on rate of affection (i.e. being a case) amongst

individuals with pathogenic variant combinations (i.e. they have at least min_ac

variants.

omega_shape Beta shape hyper-priors for prior on rate of pathogenicity amongst variants.

samples_per_chain

Number of samples to draw from each chain.

stop_early Logical value determining whether to attempt to stop the sampling as soon as

certain conditions are met (i.e. either the estimated marginal log likelihood lies within a certain confidence interval, or we are sufficiently confidence that the log Bayes factor against of model gamma = 1 over model gamma = 0 is sufficiently

low).

blocks Maximum number of blocks of samples_per_chain samples to draw before ei-

ther the confidence interval for the marginal likelihood under the model gamma = 1 is sufficiently small or terminating the sampling. This parameter is ignored

if unless stop_early==TRUE.

burn Number of samples to drop from the start of the chain.

temperatures Numeric vector of temperatures of power posteriors. One chain will be created

for each element of the vector at the corresponding temperature.

tune_temps Integer value - if greater than 0, the temperatures argument is ignored, and

instead tune_temps tuned temperatures are used instead.

return_z_trace Logical value determining whether to store the z-vectors for each chain, which

uses alot of memory, particularly if samples_per_chain, k and length(temperatures)

are large.

return_x_trace Logical value determining whether to store the x variable determined by success

 $samples \ of \ z. \ Potentially \ uses \ alot \ of \ memory, \ particularly \ if \ samples_per_chain,$

k and length(temperatures) are large.

raw_only Logical value determining whether to return raw output of MCMC routine only.

swaps Number of swaps between adjacent tempered chains to perform per update cy-

cle.

optimise_z0 Logical value determining whether to use a simulated annealing optimisation

run to tune the initial values of z.

tune_omega_and_phi_proposal_sd

Logical value determining whether the proposal SDs of the Metropolis-Hastings

estimated parameters should be tuned for a target acceptance range.

tune_block_size

Integer value giving number of samples to draw when estimatating the accep-

tance rate of the omega/phi proposals.

variant_weights

Vector of log-odds off-sets for rates of pathogenicity of individual variants rela-

tive to the global rate, omega.

standardise_weights

Boolean value determining whether weights should be standardised by subtracting their mean and dividing by their sample standard deviation. If FALSE,

weights are untransformed.

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log_phi_mean Mean for normal prior on scaling factor phi.

log_phi_sd SD for normal prior on scaling factor phi. Setting to 0 causes the weights to be

fixed and not estimated.

tandem_variant_updates

Number of tandem variant updates to make per update cycle.

.. Other arguments to be passed to stop_chain and/or tune_proposal_sds.

Details

A BeviMed_m object is a list containing elements:

- 'parameters': a list containing arguments used in the function call, including the adjusted weights used in the inference in the 'c_weights' slot,
- 'traces': a list of traces of model parameters from all MCMC chains for each parameter. Parameters sampled are z, omega, phi and x (the indicator of having a pathogenic configuration of alleles). The list of traces is named by parameter name, and each is a matrix where the rows correspond to samples. \$z has k columns for each temperature, with the samples from the true posterior (i.e. with temperature equal to 1) of z corresponding to the final k columns. Likewise, the true posterior is given by the final column for the traces of phi and omega. The trace of x is only given for temperature equal to 1 to reduce memory usage.
- 'final': a list named by model parameter giving the final sample of each,
- 'swaps': a list with an element named 'accept' which is a logical vector whose ith element indicates whether the ith swap between adjacent tempered chains was accepted or not, and an element named 'at_temperature', an integer vector whose ith element indicates which pair of consecutive temperatures was the ith to be proposed for swapping (giving the lowest one).

Value

An object of class BeviMed_m.

References

Greene et al., A Fast Association Test for Identifying Pathogenic Variants Involved in Rare Diseases, The American Journal of Human Genetics (2017), http://dx.doi.org/10.1016/j.ajhg.2017.05.015.

See Also

bevimed_m, prob_association_m

bevimed_polytomous 7

bevimed_polytomous

Model selection for multiple association models

Description

Apply bevimed to the no association model (gamma = 0) and multiple association models for different sets of variants, for instance, corresponding to different functional consequences.

Usage

```
bevimed_polytomous(y, G, ploidy = rep(2L, length(y)), variant_sets,
  prior_prob_association = rep(0.01/length(variant_sets),
  length(variant_sets)), tau0_shape = c(1, 1), moi = rep("dominant",
  length(variant_sets)), model_specific_args = vector(mode = "list",
  length = length(variant_sets)), ...)
```

Arguments

У	Logical vector of case (TRUE) control (FALSE) status.
G	Integer matrix of variant counts per individual, one row per individual and one

column per variant.

ploidy Integer vector giving ploidy of samples.

variant_sets List of integer vectors corresponding to sets of indices of G, each of which is to

be considered in a model explaining the phenotype, y.

prior_prob_association

The prior probability of association.

tau0_shape Beta shape hyper-priors for prior on rate of case labels.

moi Character vector giving mode of inheritance for each model.

model_specific_args

List of named lists of parameters to use in bevimed_m applications for specific

models.

... Other arguments to pass to bevimed_m.

References

Greene et al., A Fast Association Test for Identifying Pathogenic Variants Involved in Rare Diseases, The American Journal of Human Genetics (2017), http://dx.doi.org/10.1016/j.ajhg.2017.05.015.

See Also

bevimed_m, bevimed

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call_cpp R interface to BeviMed $c++$ MCMC procedure	
--	--

Description

Allows other functions in the package to call the c++ function passing arguments more succinctly and by name.

Usage

```
call_cpp(samples_per_chain, y, block_starts, block_ends, cases, counts,
  min_ac, tau_shape, pi_shape, omega_shape, temperatures, z0_matrix,
  estimate_omega, logit_omegas, logit_omega_proposal_sds, variant_weights,
  estimate_phi, log_phis, log_phi_mean, log_phi_sd, log_phi_proposal_sds,
  chain_swaps_per_cycle, annealing, tandem_variant_updates,
  comphet_variant_block_starts, comphet_variant_block_ends,
  comphet_variants, return_z_trace, return_x_trace, burn = 0,
  check = TRUE)
```

Arguments

samples_per_chain

Number of samples to draw from each chain.

y Logical vector of subject affectedness status.

block_starts Integer vector of k 0-indexed start positions (with respect to cases and counts)

for contiguous blocks relating to the k variants.

block_ends Integer vector of (exclusive) k 0-indexed end positions.

cases 0 based vector of case indices with respect to y.

counts Vector of variant counts.

min_ac Integer vector with a length equalling the number of individuals or length 1 (in

which case the given value is used for all individuals) giving the minimum number of alleles at pathogenic variant sites each individual requires in order to classify as having a 'pathogenic allele configuration'. Thus, this parameter encodes the mode of inheritance. For instance, setting this parameter to 1 corresponds to dominant inheritance. If there are differences in ploidy between individuals in the locus, it is necessary to set it on an sample level basis - e.g. to ensure sex is

accounted for if the locus lies on the X chromosome.

tau_shape Beta distribution parameterisation of benign variant configuration rate of affec-

tion, q.

pi_shape Beta distribution parameterisation of pathogenic variant configuration rate of

affection, p.

omega_shape Beta distribution of global rate of pathogenicty of variants in gene given pathogenic-

ity of gene, omega.

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temperatures Numeric vector of temperatures of power posteriors. One chain will be created

for each element of the vector at the corresponding temperature.

z0_matrix Matrix of logicals, where the rows are used as an initial zs for the chains.

estimate_omega Logical value determining whether to estimate the parameter omega.

logit_omegas Numeric vector of logit omega values, one value per chain.

logit_omega_proposal_sds

Numeric vector of proposal standard deviations for Metropolis-Hastings sampling of logit omega parameter, one value per chain.

variant_weights

Vector of log-odds off-sets for rates of pathogenicity of individual variants rela-

tive to the global rate, omega.

estimate_phi Logical value determining whether to estimate a scaling factor of variant_weights.

log_phis Numeric vector of log phi values, one value per chain.

log_phi_mean Mean for normal prior on scaling factor phi.

log_phi_sd SD for normal prior on scaling factor phi.

log_phi_proposal_sds

Numeric vector of proposal standard deviations for Metropolis-Hastings sampling of log phi parameter, one value per chain.

chain_swaps_per_cycle

Number of chain swaps to propose per update cycle.

annealing Logical value determining whether to anneal the chains, e.g. for optimisation.

tandem_variant_updates

Number of tandem variant updates to make per update cycle.

comphet_variant_block_starts

0-indexed start positions for contiguous blocks of variants in comphet_variants.

comphet_variant_block_ends

As comphet_variant_block_starts for (exclusive) stop positions.

comphet_variants

Integer vector giving variant numbers (0-based, i.e. between 0 and k-1). Used to pick pairs of variants for tandem updates from.

return_z_trace Logical value determining whether to store the z-vectors for each chain, which

 $uses\ a lot\ of\ memory,\ particularly\ if\ samples_per_chain,\ k\ and\ length (temperatures)$

are large.

return_x_trace Logical value determining whether to store the x variable determined by success

samples of z. Potentially uses alot of memory, particularly if samples_per_chain,

k and length(temperatures) are large.

burn Number of samples to drop from the start of the chain.

check Logical value indicating whether to perform validation on the arguments before

calling the c++ function.

Value

Object of class BeviMed_raw, containing the output of the MCMC sampling.

CI_gamma1_evidence

Estimate confidence interval for estimated marginal likelihood

Description

Central limit theorem not applicable so use simulation to estimate confidence interval for evidence.

Usage

```
CI_gamma1_evidence(temperatures, y_log_lik_t_equals_1_traces,
  confidence = 0.95, simulations = 1000)
```

Arguments

temperatures Numeric vector of temperatures of power posteriors. One chain will be created

for each element of the vector at the corresponding temperature.

y_log_lik_t_equals_1_traces

Numeric matrix of log probabilities of y at different temperatures (columns) in

different iterations (rows).

confidence Numeric value of statistical confidence with which returning interval should

contain the true value.

simulations Integer value of number of simulations to use in estimation of the confidence

interval.

Value

Confidence interval as numeric vector of length 2.

```
conditional_prob_pathogenic
```

Calculate probability of pathogencity for variants conditional on mode of inheritance.

Description

Calls bevimed_m and extract_conditional_prob_pathogenic to obtain probabilities of pathogenicity.

Usage

```
conditional_prob_pathogenic(...)
```

Arguments

.. Arguments to pass to bevimed_m.

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Value

Probabilities of pathogenicity.

See Also

```
extract_conditional_prob_pathogenic, bevimed_m
```

expected_explained

Calculate expected number of explained cases

Description

Use bevimed_m to perform inference under model gamma = 1 and return only the expected number of cases explained by pathogenic allele configurations.

Usage

```
expected_explained(...)
```

Arguments

... Arguments to pass to bevimed_m.

Value

Numeric value.

See Also

bevimed_m, extract_expected_explained

explaining_variants

Calculate expected number of pathogenic variants in cases

Description

Use bevimed_m to perform inference under model gamma = 1 and return only the expected number of pathogenic variants in cases.

Usage

```
explaining_variants(...)
```

Arguments

.. Arguments to pass to bevimed_m.

Value

Numeric value.

See Also

```
extract_explaining_variants, bevimed_m
```

```
extract_conditional_prob_pathogenic
```

Extract probability of pathogenicity for variant conditional on a given association model

Description

Extract the probability of pathogenicity for individual variants from a BeviMed_m object.

Usage

```
extract_conditional_prob_pathogenic(x)
```

Arguments

Х

Object of class x_BeviMed_m. See function bevimed_m.

Value

Vector of probabilities of pathogenicity for individual variants.

See Also

```
conditional_prob_pathogenic, bevimed_m
```

```
extract_expected_explained
```

Extract expected number of explained cases

Description

Extract expected number of cases explained by pathogenic configurations of alleles from BeviMed_m object.

Usage

```
extract_expected_explained(x)
```

Arguments

Χ

Object of class x_BeviMed_m. See function bevimed_m.

Value

Numeric value.

See Also

```
expected_explained, bevimed_m
```

```
extract_explaining_variants
```

Extract expected number of pathogenic variants in cases

Description

Extract expected number of variants involved in cases explained by pathogenic conigurations of alleles from $BeviMed_m object$.

Usage

```
extract_explaining_variants(x)
```

Arguments

Х

Object of class x_BeviMed_m. See function bevimed_m.

Value

Numeric value.

See Also

```
explaining_variants, bevimed_m
```

extract_gamma1_evidence

Extract evidence for model gamma = 1

Description

Extract evidence from BeviMed_m object.

Usage

```
extract_gamma1_evidence(x)
```

Arguments

Χ

Object of class x_BeviMed_m. See function bevimed_m.

Value

Log marginal likelihood.

See Also

```
gamma1_evidence, bevimed_m
```

extract_prob_association

Extract the posterior probability of association

Description

Get posterior probability of association as numeric value, or optionally as numeric vector of length two with probabilities broken down by mode of inheritance (by passing by_model=TRUE), from a BeviMed object.

Usage

```
extract_prob_association(x, by_model = FALSE)
```

Arguments

x Object of class BeviMed.

by_model Logical value determining whether to return probabilities broken down by mode

of inheritance.

Value

Probability values.

See Also

prob_association, bevimed

extract_prob_pathogenic

Extract variant marginal probabilities of pathogenicity

Description

Extract the marginal probability of pathogenicity for individual variants from BeviMed object, optionally broken down by mode of inheritance/model.

Usage

```
extract_prob_pathogenic(x, by_model = TRUE)
```

Arguments

x Object of class BeviMed.

by_model Logical value determining whether to return probabilities broken down by mode

of inheritance.

Value

A vector of probabilities of pathogenicity for individual variants, or if by_model is TRUE, then a matrix of probabilities, with rows corresponding to modes of inheritance and columns to variants.

See Also

prob_pathogenic, bevimed

gamma1_evidence

_	Calculate marginal probability of observed case-control status y under model gamma = 0
---	--

Description

Marginal probability calculated exactly by integration.

Usage

```
gamma0_evidence(y, tau0_shape = c(1, 1))
```

Arguments

y Logical vector of case (TRUE) control (FALSE) status. tau0_shape Beta shape hyper-priors for prior on rate of case labels

Value

Log marginal likelihood.

See Also

bevimed, gamma1_evidence

gamma1_evidence

 $Calculate\ evidence\ under\ model\ gamma=1$

Description

Use bevimed_m to perform inference under model gamma = 1 and return only the log evidence/integrated likelihood.

Usage

```
gamma1_evidence(...)
```

Arguments

... Arguments to pass to bevimed_m.

Value

Log marginal likelihood.

See Also

```
bevimed_m, extract_gamma1_evidence
```

log_BF

log_BF	Calculate log Bayes factor between an association model with a given
	$mode\ of\ inheritance\ and\ model\ gamma=0$

Description

Compute log Bayes factor of an association model and model gamma = 0.

Usage

```
log_BF(y, tau0\_shape = c(1, 1), ...)
```

Arguments

y Logical vector of case (TRUE) control (FALSE) status.
tau@_shape Beta shape hyper-priors for prior on rate of case labels.
... Arguments to pass to bevimed_m.

Value

Log Bayes factor.

See Also

```
bevimed_m, prob_association_m
```

print.BeviMed

Print readable summary of BeviMed object

Description

Print summary statistics of BeviMed inference, including probability of association, probability of dominant inheritance given association and probability of pathogenicity of each variant under dominant and recessive inheritance.

Usage

```
## S3 method for class 'BeviMed'
print(x, ...)
```

Arguments

x BeviMed object.

... Arguments passed to summary.BeviMed

Value

Prints a summary.

See Also

```
summary.BeviMed
```

print.BeviMed_m

Print BeviMed_m object

Description

Print summary statistics for BeviMed_m object.

Usage

```
## S3 method for class 'BeviMed_m'
print(x, ...)
```

Arguments

x Object of class x_BeviMed_m. See function bevimed_m.... Unused arguments.

Value

Prints a summary.

See Also

```
summary.BeviMed_m
```

print.BeviMed_summary Print readable summary of BeviMed_summary object.

Description

Print summary statistics of BeviMed inference, including probability of association, probability of dominant inheritance given association and probability of pathogenicity of each variant under dominant and recessive inheritance.

Usage

```
## S3 method for class 'BeviMed_summary'
print(x, print_prob_pathogenic = TRUE, ...)
```

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Arguments

```
x BeviMed_summary object.
```

print_prob_pathogenic

Logical value indicating whether to print list of marginal probabilities of $z_j = 1$ for all variants j under each mode of inheritance.

... Unused arguments

Value

Prints a summary

prob_association

Calculate probability of association

Description

Calculate probability of an association between case/control label and allele configuration, optionally broken down by mode of inheritance/model.

Usage

```
prob_association(by_model = FALSE, ...)
```

Arguments

by_model Logical value determining whether to return probabilities broken down by mode

of inheritance.

... Arguments to pass to bevimed.

Value

Probability of association.

See Also

bevimed, extract_prob_association

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prob_association_m

Calculate probability of association for one mode of inheritance

Description

Equivalent to prob_association where the prior probability of one mode of inheritance is 1. This function is faster, as it only calls bevimed_m once.

Usage

```
prob_association_m(y, min_ac = 1L, prior_prob_association = 0.01, ...)
```

Arguments

у

Logical vector of case (TRUE) control (FALSE) status.

min_ac

Integer vector with a length equalling the number of individuals or length 1 (in which case the given value is used for all individuals) giving the minimum number of alleles at pathogenic variant sites each individual requires in order to classify as having a 'pathogenic allele configuration'. Thus, this parameter encodes the mode of inheritance. For instance, setting this parameter to 1 corresponds to dominant inheritance. If there are differences in ploidy between individuals in the locus, it is necessary to set it on an sample level basis - e.g. to ensure sex is

accounted for if the locus lies on the X chromosome.

prior_prob_association

The prior probability of association.

... Other arguments to pass to log_BF.

Value

Probability value.

See Also

log_BF, prob_association, bevimed_m

prob_pathogenic

Calculate variant marginal probabilities of pathogencity

Description

Calls bevimed and extract_prob_pathogenic to obtain marginal probabilities of pathogenicity.

Usage

```
prob_pathogenic(by_model = FALSE, ...)
```

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Arguments

by_model Logical value determining whether to return probabilities broken down by mode

of inheritance.

... Arguments to pass to bevimed.

Value

If by_model is FALSE, a vector of probabilities of pathogenicity for each variant, otherwise a list of vectors of probabilities of pathogenicity conditional on each compared association model.

See Also

```
extract_prob_pathogenic, bevimed
```

stack_BeviMeds

Concatenate objects of class BeviMed_raw

Description

This function could be used to stitch together consecutive chains to create one larger sampled set of states from the MCMC procedure.

Usage

```
stack_BeviMeds(objects)
```

Arguments

objects

list of BeviMed_raw objects.

Value

BeviMed object.

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stop_chain

Apply the MCMC algorithm in blocks until conditions are met

Description

Sample blocks of a given size until either the estimated log marginal likelihood falls within a given confidence interval, there is sufficient confidence that the evidence model gamma = 1 is at most a certain quantity, or a certain number of blocks have been sampled.

Usage

```
stop_chain(y, blocks_remaining, start_zs, start_logit_omegas,
   start_log_phis, temperatures, tolerance = 1, confidence = 0.95,
   simulations = 1000, log_evidence_threshold = -Inf,
   y_log_lik_t_equals_1_traces = matrix(ncol = length(temperatures), nrow
   = 0), full_block_traces = list(), verbose = FALSE, ...)
```

Arguments

y Logical vector of case (TRUE) control (FALSE) status.

blocks_remaining

Maximum number of blocks left before termination.

start_zs Initial (logical) z-matrix.

start_logit_omegas

Initial values of logit_omega (numeric vector - one value per chain).

start_log_phis Initial values of log_phi (numeric vector - one value per chain).

temperatures Numeric vector of temperatures of power posteriors. One chain will be created

for each element of the vector at the corresponding temperature.

tolerance Maximum width for confidence_interval of log marginal likelihood to allow be-

fore stopping the chain.

confidence Numeric value of statistical confidence with which returning interval should

contain the true value.

simulations Integer value of number of simulations to use in estimation of the confidence

interval.

log_evidence_threshold

Numeric value used to determine whether to stop the sampling procedure after successive blocks. If we are confident (to the level of confidence) that the evidence for model gamma = 1 is under this value, sampling is halted.

y_log_lik_t_equals_1_traces

Numeric matrix of log probabilities of y at different temperatures (columns) in different iterations (rows).

full_block_traces

List of outputs of calls to MCMC routine.

verbose To print execution progress or not.

... Other arguments passed to call_cpp

23 subset_variants

Value

An object of class BeviMed.

subset_variants

Remove variants with no data for pathogenicity

Description

Subset an allele count matrix given a minimum allele count threshold for pathogenicity per individual so that only variants for which data relevant to pathogencity are retained. This is useful to apply before running bevimed as it reduces the size of the parameter space used in the inference.

Usage

```
subset_variants(G, min_ac = 1L, return_variants = FALSE)
```

Arguments

G

Integer matrix of variant counts per individual, one row per individual and one column per variant.

min_ac

Integer vector with a length equalling the number of individuals or length 1 (in which case the given value is used for all individuals) giving the minimum number of alleles at pathogenic variant sites each individual requires in order to classify as having a 'pathogenic allele configuration'. Thus, this parameter encodes the mode of inheritance. For instance, setting this parameter to 1 corresponds to dominant inheritance. If there are differences in ploidy between individuals in the locus, it is necessary to set it on an sample level basis - e.g. to ensure sex is

accounted for if the locus lies on the X chromosome.

return_variants

Logical value determining whether to return an integer vector of indices of retained variants or the subsetted allele count matrix

summary.BeviMed

Summarise a BeviMed object

Description

Create a summary of inference over model gamma = 0 and association models.

Usage

```
## S3 method for class 'BeviMed'
summary(object, ...)
```

Arguments

object Object of class BeviMed.

... Arguments passed to summary.BeviMed_m.

Details

Returns a BeviMed_summary object, which is a list containing elements:

- 'prob_association': the probability of association under each association model,
- 'prior_prob_association': the prior probability of association for each association model,
- 'gamma0_evidence': the log evidence under model gamma = 0,
- 'models': a list of summaries of model conditional inferences, i.e. objects of class BeviMed_m_summary. See summary.BeviMed_m for more details.

Value

Object of class BeviMed_summary.

See Also

```
summary.BeviMed_m
```

summary.BeviMed_m

Summarise a BeviMed_m object

Description

Create a summary of inference conditional on mode of inheritance.

Usage

```
## S3 method for class 'BeviMed_m'
summary(object, confidence = 0.95,
    simulations = 1000, ...)
```

Arguments

object Object of class BeviMed_m. See function bevimed_m.

confidence Numeric value of statistical confidence with which returning interval should

contain the true value.

simulations Integer value of number of simulations to use in estimation of the confidence

interval.

... Unused arguments.

summary.BeviMed_m 25

Details

Returns a BeviMed_m_summary object, which is a list containing elements:

- 'gamma1_evidence': the log evidence under model gamma = 1,
- 'gamma1_evidence_confidence_interval': a confidence interval for the log evidence under model gamma = 1,
- 'conditional_prob_pathogenic': vector of marginal probabilities of pathogenicity for individual variants,
- 'expected_explained': the expected number of cases with a pathogenic configuration of alleles,
- 'explaining_variants': the expected number of variants present for which cases harbour a rare allele,
- 'number_of_posterior_samples': the number of samples from the posterior distribution of the model parameters which upon which the summary is based,
- 'omega_estimated': logical value indicating whether the parameter omega was estimated,
- 'omega': the posterior mean of omega,
- 'omega_acceptance_rate': if omega was estimated, the rate of acceptance of proposed omega values in the Metropolis-Hastings sampling routine,
- 'phi_estimated': logical value indicating whether the parameter phi was estimated,
- 'phi': the posterior mean of phi,
- 'phi_acceptance_rate': if phi was estimated, the rate of acceptance of proposed phi values in the Metropolis-Hastings sampling routine,
- 'N': number of samples in the analysis,
- 'k': number of variants in the analysis,
- 'variant_counts': list of counts of each variant for cases and controls,
- 'temperatures': numeric vector of temperatures used as temperatures for tempered MCMC chains

Value

Object of class BeviMed_m_summary.

See Also

summary.BeviMed

26 tune_proposal_sds

sum_ML_over_PP

Calculate marginal likelihood from power posteriors output

Description

Calculate the Marginal Likelihood by summation over power posterior likelihood exptectances

Usage

```
sum_ML_over_PP(y_log_lik_t_equals_1_traces, temperatures)
```

Arguments

```
y_log_lik_t_equals_1_traces
```

Numeric matrix of log probabilities of y at different temperatures (columns) in different iterations (rows).

temperatures Numeric vector of temperatures used to produce y_log_lik_t_equals_1_traces.

Value

Numeric value of estimated log marginal likelihood.

tune_proposal_sds

Tune proposal standard deviation for MH sampled parameters

Description

Tune the proposal standard deviations for the Metropolis-Hastings updates of either phi or omega

Usage

```
tune_proposal_sds(tune_for = c("logit_omega"), initial_proposal_sds,
  target_acceptance_range = c(0.3, 0.7), other_param_proposal_sd = 0.7,
  max_tuning_cycles = 10, initial_rate = 1, rate_decay = 1.2,
  verbose = FALSE, ...)
```

Arguments

tune_for

Character vector of length one, naming which variable to tune the proposal SDs for: either "logit_omega" or "log_phi".

initial_proposal_sds

Numeric vector with the initial values of the proposal SDs.

target_acceptance_range

Numeric vector of length 2 where the first element is the lower bound for the acceptance interval and the second is the upper bound.

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other_param_proposal_sd

The proposal SD to use for log_phi when tuning logit_omega or vice versa.

max_tuning_cycles

Maximum number of tuning cycles to perform before returning the proposal

SDs as they are.

initial_rate Initial rate at which to mutate the proposal SDs.

rate_decay Geometric rate of decay for size of proposal SD mutation with each successive

tuning cycle.

verbose To print execution progress or not.

... Other arguments to be passed to call_cpp.

Value

Numeric vector of proposal SDs for the different temperature chains.

tune_temperatures

Tune temperatures

Description

Tune temperatures using interval bisection to minimimise Kullback-Liebler divergence between adjacent power posteriors

Usage

```
tune_temperatures(number_of_temperatures, return_temperatures = FALSE,
...)
```

Arguments

number_of_temperatures

 $\label{thm:continuous} Integer \ value \ giving \ number \ of \ tuned \ temperatures \ (including \ 0 \ and \ 1) \ to \ obtain.$ $\ return_temperatures$

Logical value determining whether to return just the numeric vector of tuned temperatures or to return the BeviMed_m-classed object containing the output of the MCMC sampling.

... Other arguments to pass to call_cpp.

Value

If return_temperatures == TRUE, a numeric vector of tuned temperatures, otherwise an object of class BeviMed_m.

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