Package 'methyvim'

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Title Targeted Variable Importance for Differential Methylation Analysis

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Description This package provides facilities for differential methylation analysis based on variable importance measures (VIMs), a class of statistical target parameters that arise in causal inference. The estimation and inference procedures provided are nonparametric, relying on ensemble machine learning to flexibly assess functional relationships among covariates and the outcome of interest. These tools can be applied to differential methylation at the level of CpG sites, to obtain valid statistical inference even after corrections for multiple hypothesis testing.

Depends R (>= 3.4.0)

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URL https://github.com/nhejazi/methyvim

BugReports https://github.com/nhejazi/methyvim/issues

Encoding UTF-8

- Imports stats, cluster, methods, ggplot2, ggsci, gridExtra, superheat, dplyr, gtools, tmle, future, doFuture, BiocGenerics, BiocParallel, SummarizedExperiment, GenomeInfoDb, bumphunter, IRanges, limma, minfi
- Suggests testthat, knitr, rmarkdown, BiocStyle, SuperLearner, earth, nnet, gam, arm, snow, parallel, BatchJobs, minfiData, methyvimData

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fdr_msa

FDR-MSA correction

Description

Modified FDR Controlling Procedure for Multi-Stage Analyses (MJ van der Laan and C Tuglus, 2009, <doi:10.2202/1544-6115.1397>)

Usage

fdr_msa(pvals, total_obs)

Arguments

pvals	Numeric vector containing the p-values that result from any chosen statistical hypothesis testing procedure.
total_obs	Numeric indicating the total number of observations that would have been avail- able for testing prior to the selection procedure employed in the multi-stage analysis performed.

Value

A numeric vector of corrected p-values, controlling the False Discovery Rate, using the method of Tuglus and van der Laan.

Examples

```
g <- 1e4
n <- 1e2
p <- abs(rnorm(n, mean = 1e-8, sd = 1e-2))
# treating the vector p as one of p-values, FDR-MSA may be applied
fdr_p <- fdr_msa(pvals = p, total_obs = g)</pre>
```

methyheat

Description

Heatmap for methytmle objects

Usage

methyheat(x, ..., n_sites = 25, type = "raw")

Arguments

х	Object of class methytmle as produced by an appropriate call to methyvim.
	Additional arguments passed to superheat. Consult the documentation of the superheat package for a list of options.
n_sites	Numeric indicating the number of CpG sites to be shown in the plot. If the number of sites analyzed is greater than this cutoff, sites to be displayed are chosen by ranking sites based on their raw (marginal) p-values.
type	Whether to plot the original data (M-values or Beta-values) for the set of top CpG sites or to plot the measurements after applying a transformation into influence curve space (with respect to the target parameter of interest). The latter uses the fact that the parameters have asymptotically linear representations to obtain a rotation of the raw data into an alternative space; moreover, in this setting, the heatmap reduces to visualizing a supervised clustering procedure.

Value

Nothing. This function is called for its side-effect of outputting a heatmap to the graphics device. The heatmap is constructed using the superheat package.

Examples

methytmle-class

Description

Constructor for class methytmle

Value

methytmle object, subclassed from GenomicRatioSet.

Examples

```
library(methyvimData)
suppressMessages(library(SummarizedExperiment))
data(grsExample)
# cast the GenomicRatioSet to class methytmle
methy_tmle <- .methytmle(grsExample)</pre>
```

methyvim

Differential Methylation Statistics with Variable Importance Measures

Description

Computes the Targeted Minimum Loss-Based Estimate of a specified statistical target parameter, formally defined within models from causal inference. The variable importance measures currently supported are the Average Treatment Effect (ATE) and a Nonparametric Variable Importance Measure (NPVI, formally defined by Chambaz, Neuvial, and van der Laan <doi:10.1214/12-EJS703>).

Usage

```
methyvim(data_grs, var_int, vim = c("ate", "rr", "npvi"), type = c("Beta",
  "Mval"), filter = c("limma"), filter_cutoff = 0.05, window_bp = 1000,
  corr_max = 0.75, obs_per_covar = 20, sites_comp = NULL,
  parallel = TRUE, future_param = NULL, bppar_type = NULL,
  return_ic = FALSE, shrink_ic = FALSE, tmle_type = c("glm", "sl"),
  tmle_args = list(family = "binomial", g_lib = c("SL.mean", "SL.glm",
  "SL.bayesglm", "SL.gam"), Q_lib = c("SL.mean", "SL.glm", "SL.gam",
  "SL.earth"), npvi_cutoff = 0.25, npvi_descr = NULL),
  tmle_backend = c("tmle", "drtmle", "tmle.npvi"))
```

Arguments

data_grs An object of class minfi::GenomicRatioSet, containing standard data structures associated with DNA Methylation experiments. Consult the documentation for minfi to construct such objects.

A numeric vector containing subject-level measurements of the variable of in- terest. The length of this vector must match the number of subjects exactly. If argument vim is set to "ate" or "rr", then the variable of interest is treated as an exposure, and the variable must be binary in such cases. If setting vim to target parameters assessing continuous treatment effects, then the variable need not be binary of course.
Character indicating the variable importance measure to be used in the estima- tion procedure. Currently supported options are the ATE for discretized expo- sures and NPVI for continuous exposures. ATE and RR are the appropriate choices when the underlying scientific question is of the effect of an exposure on methylation, while NPVI (and other continuous treatment parameters) ought to be used when the effect of methylation on an outcome is sought.
Character indicating the particular measure of DNA methylation to be used as the observed data in the estimation procedure, either Beta values or M-values. The data are accessed via minfi::getBeta or minfi::getM.
Character indicating the model to be implemented when screening the data_grs object for CpG sites. The only currently supported option is "limma". Contributions for other methods are welcome.
Numeric indicating the p-value cutoff that defines which sites pass through the filter.
Numeric indicating the maximum genomic distance (in base pairs) between two sites for them to be considered neighboring sites.
Numeric indicating the maximum correlation that a neighboring site can have with the target site.
Numeric indicating the number of observations needed for for covariate included in W for downstream analysis. This ensures the data is sufficient to control for the covariates.
A numeric indicating the maximum number of sites for which a variable impor- tance measure is to be estimated post-screening. This is not typically useful in scientific settings, but may be useful when a large number of CpG sites pass the initial screening phase.
Logical indicating whether parallelization ought to be used. See the documenta- tion of set_parallel for more information, as this argument is passed directly to that internal function.
Character indicating the type of parallelization to be used from the list available via the future package. See the documentation for set_parallel for more information, as this argument is passed directly to that internal function.
Character specifying the type of backend to be used for parallelization via BiocParallel. See the documentation for set_parallel for more information, as this argument is passed directly to that internal function.
Logical indicating whether an influence curve estimate should be returned for each site that passed through the filter.
Logical indicating whether limma should be applied to reduce the variance in the ic based estimates in return_ic.
Character indicating the general class of regression models to be used in fitting the propensity score and outcome regressions. This is generally a shorthand and is overridden by tmle_args if that argument is changed from its default values.

tmle_args	List giving several key arguments to be passed to one of tmle::tmle or tmle.npvi::tmle.npvi									
	depending on the particular variable importance measure specified. This over-									
	rides tmle_type, which itself provides sensible defaults. Consider changing									
	this away from default settings only if you have sufficient experience with soft-									
	ware and the underlying theory for Targeted Learning. For more information, consider consulting the documentation of the tmle and tmle.npvi packages.									
tmle_backend	A character indicating the package to be used in the estimation procedure. The user should only set this parameter if they have sufficient familiarity with the backend packages used for estimation. Current choices include tmle, drtmle, and tmle.npvi.									

Value

An object of class methytmle, with all unique slots filled in, in particular, including indices of CpG sites that pass screening, cluster of neighboring CpG sites, and a matrix of the results of the estimation procedure performed for the given variable importance measure. Optionally, estimates of the propensity score and outcome regressions, as well as the original data rotated into influence curve space may be returned, if so requested.

Examples

methyvolc

Volcano plot for methytmle objects

Description

Volcano plot for methytmle objects

Usage

```
methyvolc(x, param_bound = 2, pval_bound = 0.2)
```

Arguments

х	Object of class methytmle as produced by an appropriate call to methyvim.
param_bound	Numeric for a threshold indicating the magnitude of the size of the effect considered to be interesting. This is used to assign groupings and colors to individual CpG sites.
pval_bound	Numeric for a threshold indicating the magnitude of p-values deemed to be in- teresting. This is used to assign groupings and colors to individual CpG sites.

plot.methytmle

Value

Object of class ggplot containing a volcano plot of the estimated effect size on the x-axis and the $-\log 10$ (p-value) on the y-axis. The volcano plot is used to detect possibly false positive cases, where a test statistic is significant due to low variance.

Examples

plot.methytmle *Plot p-values of methytmle objects*

Description

Plot p-values of methytmle objects

Usage

```
## S3 method for class 'methytmle'
plot(x, ..., type = "both")
```

Arguments

х	Object of class methytmle as produced by an appropriate call to methyvim.
•••	Additional arguments passed plot as necessary.
type	The type of plot to build: one of side-by-side histograms (type "both") com- paring raw p-values to FDR-adjusted p-values (using the FDR-MSA correction) or either of these two histogram separately. Set this argument to "raw_pvals" for a histogram of the raw p-values, and to "fdr_pvals" for a histogram of the FDR-corrected p-values.

Value

Object of class ggplot containing a histogram or side-by-side histograms of the raw (marginal) and corrected p-values, with the latter computed automatically using the method of Tuglus and van der Laan.

Examples

wrap_in_try Wrap a Function in a Try Statement

Description

Function factory that generates versions of functions wrapped in try. Originally fround in and borrowed from package origami.

Usage

wrap_in_try(fun, ...)

Arguments

fun	A function to be wrapped in a try statement.
	Additional arguments passed to the previous argument fun

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