Package ‘adaptMT’

October 12, 2022

Type Package

Title Adaptive P-Value Thresholding for Multiple Hypothesis Testing with Side Information

Version 1.0.0

Maintainer Lihua Lei <lihua.lei@berkeley.edu>

Description Implementation of adaptive p-value thresholding (AdaPT), including both a framework that allows the user to specify any algorithm to learn local false discovery rate and a pool of convenient functions that implement specific algorithms. See Lei, Lihua and Fithian, William (2016) <arXiv:1609.06035>.

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Encoding UTF-8

LazyData true


BugReports https://github.com/lihualei71/adaptMT/issues

Suggests glmnet, HDtweedie, mgcv, splines, testthat, knitr, rmarkdown, dplyr

RoxygenNote 6.0.1

Imports methods

VignetteBuilder knitr

NeedsCompilation no

Author Lihua Lei [aut, cre]

Repository CRAN

Date/Publication 2018-07-31 12:00:03 UTC
R topics documented:

- `adapt` .......................................................... 2
- `adapt_gam` ......................................................... 5
- `adapt_glm` ........................................................ 6
- `adapt_glmnet` ...................................................... 8
- `corr_lfdr` .......................................................... 9
- `ctgm_lfdr` .......................................................... 10
- `estrogen` .......................................................... 12
- `gen_adapt_model` ................................................ 13
- `gen_exp_family` .................................................. 15
- `plot_1d` ........................................................... 16
- `plot_2d` ........................................................... 17

Index .......................................................... 19

adapt  Adaptive P-value Thresholding

Description

adapt is a framework allowing for arbitrary exponential families for computing E-steps and arbitrary algorithms for fitting M-steps.

Usage

adapt(x, pvals, models, dist = beta_family(), s0 = rep(0.45, length(pvals)),
alphas = seq(0.01, 1, 0.01), params0 = list(pix = NULL, mux = NULL),
nfits = 20, nms = 1, niter_fit = 10, tol = 1e-04, niter_ms = 20,
cr = "BIC", verbose = list(print = TRUE, fit = FALSE, ms = TRUE))

Arguments

- `x`            covariates (i.e. side-information). Should be compatible to `models`. See Details
- `pvals`        a vector of values in [0, 1]. P-values
- `models`       an object of class "adapt_model" or a list of objects of class "adapt_model". See Details
- `dist`         an object of class "gen_exp_family". `beta_family()` as default
- `s0`           a vector of values in [0, 0.5). Initial threshold.
- `alphas`       a vector of values in (0, 1). Target FDR levels.
- `params0`      a list in the form of list(pix = , mux = ). Initial guess of pi(x) and mu(x). NULL as default
- `nfits`        a positive integer. Number of model-fitting steps. See Details
- `nms`          a non-negative integer. Number of model selection steps. See Details
- `niter_fit`    a positive integer. Number of EM iterations in model fitting
tol a positive scalar. EM algorithm stops when pi(x) and mu(x) in consecutive steps
differ by at most 'tol' for each element

niter_ms a positive integer. Number of EM iterations in model selection

cr a string. The criterion for model selection with BIC as default. Also support
AIC, AICC and HIC

verbose a list of logical values in the form of list(print = , fit = , ms = ). Each ele-
ment indicates whether the relevant information is outputted to the console. See
Details

Details

x should have a type compatible to the fitting functions in models. For GLM and GAM, x should
be a data.frame. For glmnet, x should be a matrix.

models could either be an adapt_model object, if a single model is used, or a list of adapt_model
objects, each of which corresponding to a model. Each element should be generated by gen_adapt_model.
For glm/gam/glmnet, one can use the shortcut by running gen_adapt_model with name = "glm"
or "gam" or "glmnet" but without specifying pifun, mufun, pifun_init and mufun_init. See
examples below.

nfits is the number of model fitting steps plus nms, the model selection steps, if models contains
multiple adapt_model objects. Suppose M is the number of masked p-values at the initial step,
then the model is updated at the initial step and at every time when [M/nfits] more p-values are
revealed. If nms > 0, model selection is performed at the initial step an at every time when [M/nms]
more p-values are revealed. Between two consecutive model selection steps, the model selected
from the last step is used for model fitting. For example, when M = 10000, nfits = 10 and nms = 2,
model selection will be performed at the initial step and when 5000 p-values are revealed, while the
model fitting will be performed when 1000, 2000, 3000, 4000, 6000, 7000, 8000, 9000 p-values are
revealed.

verbose has three elements: print, fit and ms. If print = TRUE, the progress of the main pro-
cedure is outputted to the console, in the form of "alpha = 0.05: FDPhat 0.0333, Number of Rej.
30" (where the numbers are made up for illustration). If fit = TRUE, a progress bar for the model
fitting is outputted to the console. Similarly, if ms = TRUE, a progress bar for the model selection is
outputted to the console.

For ultra-large scale problems (n > 10^5), it is recommended to keep alphas short because the
output s is of size n x length(alphas). is length(alphas).

The output qvals gives the q-values of each hypothesis. qvals[i] is defined as the minimum target
FDR level such that pvals[i] is rejected. For hypotheses with p-values above s0, the q-values are
set to be Inf because they are never rejected by AdaPT for whatever alpha.

The output order gives the order of (the indices of) p-values being revealed, i.e. being in the region
(s, 1-s). The latter hypotheses appeared in order have smaller q-values (i.e. are more likely to be
rejected).

Value

nrejs a vector of integers. Number of rejections for each alpha

rejs a list of vector of integers. The set of indices of rejections for each alpha
adapt

s

a matrix of size length(pvals) X length(alphas). Threshold curves for each alpha

params

a list. Each element is a list in the form of list(pix = , mux = , alpha = , nmasks =), recording the parameter estimates, the achieved alpha and the number of masked p-values. To avoid massive storage cost, it only contains the information when a new target FDR level is achieved. As a result, it might be shorter than nfits.

qvals

a vector of values in [0, 1]UInf. Q-values. See Details

order

a permutation of 1:length(pvals). Indices of hypotheses arranged in the order of reveal. See Details

alphas

same as the input alphas

dist

same as the input dist

models

a list of adapt_model objects of length params. The model used in each fitting step. As in params, it only contains the model when a new target FDR level is achieved and each element corresponds to an element of params.

info

a list of length nfits. Each element is a list recording extra information in each fitting step, e.g. degree of freedom (df) and variable importance (vi). As in params, it only contains the model information when a new target FDR level is achieved and each element corresponds to an element of params.

args

a list including the other inputs nfits, nms, niter_fit, niter_ms, tol, cr

Examples

# Load estrogen data
data(estrogen)
pvals <- as.numeric(estrogen$pvals)
x <- data.frame(x = as.numeric(estrogen$ord_high))
dist <- beta_family()

# Subsample the data for convenience
inds <- (x$x <= 5000)
pvals <- pvals[inds]
x <- x[inds,,drop = FALSE]

dist <- beta_family()

# Generate models for function adapt
library("splines")
formulas <- paste0("ns(x, df = ", 6:10, ")")
models <- lapply(formulas, function(formula){
  piargs <- muargs <- list(formula = formula)
  gen_adapt_model(name = "glm", piargs = piargs, muargs = muargs)
})

# Run adapt
res <- adapt(x = x, pvals = pvals, models = models, dist = dist, nfits = 10)
**Description**

`adaptive_gam` is a wrapper of `adapt` that fits pi(x) and mu(x) by `gam` from `mgcv` package.

**Usage**

```r
adaptive_gam(x, pvals, pi_formulas, mu_formulas, piargs = list(),
             muargs = list(), dist = beta_family(), s0 = rep(0.45, length(pvals)),
             alphas = seq(0.01, 1, 0.01), ...)```

**Arguments**

- `x`: covariates (i.e. side-information). Should be compatible to `models`. See Details
- `pvals`: a vector of values in [0, 1]. P-values
- `pi_formulas`: a vector/list of strings/formulas. Formulas for fitting pi(x) by gam. See Details
- `mu_formulas`: a vector/list of strings/formulas. Formulas for fitting mu(x) by gam. See Details
- `piargs`: a list. Other arguments passed to `gam` for fitting pi(x)
- `muargs`: a list. Other arguments passed to `gam` for fitting mu(x)
- `dist`: an object of class "gen_exp_family". `beta_family()` as default
- `s0`: a vector of values in [0, 0.5). Initial threshold.
- `alphas`: a vector of values in (0, 1). Target FDR levels.
- `...`: other arguments passed to `adapt` (except `models`)

**Details**

`pi_formulas` and `mu_formulas` can either be a list or a vector with each element being a string or a formula. For instance, suppose `x` has a single column with name `x1`, the following five options are valid for the same inputs (`ns` forms a spline basis with `df` knots and `s` forms a spline basis with knots automatically selected by generalized cross-validation):

1. `c("x1", "ns(x1, df = 8)", "s(x1)")`
2. `c("~ x1", "~ ns(x1, df = 8)", "s(x1)")`
3. `list("x1", "ns(x1, df = 8)", "s(x1)")`
4. `list("~ x1", "~ ns(x1, df = 8)", "s(x1)")`
5. `list(~ x1, ~ ns(x1, df = 8), s(x1))`

There is no need to specify the name of the response variable, as this is handled in the function.

When `x` has a few variables, it is common to use non-parametric GLM by replacing `x` by a spline basis of `x`. In this case, `ns` from `library(splines)` package or `s` from `mgcv` package are suggested. When `s` (from `mgcv` package) is used, it is treated as a single model because the knots will be selected automatically.
adapt_glm

Adaptive P-value Thresholding with Generalized Linear Models

Description

adapt_glm is a wrapper of adapt that fits pi(x) and mu(x) by glm.

Usage

adapt_glm(x, pvals, pi_formulas, mu_formulas, dist = beta_family(),
          s0 = rep(0.45, length(pvals)), alphas = seq(0.01, 1, 0.01),
          piargs = list(), muargs = list(), ...)

Arguments

x          covariates (i.e. side-information). Should be compatible to models. See Details
pvals      a vector of values in [0, 1]. P-values

See Also

adapt, adapt_glm, adapt_glmnet, gam, ns, s

Examples

# Generate a 2-dim x
n <- 400
x1 <- x2 <- seq(-100, 100, length.out = 20)
x <- expand.grid(x1, x2)
colnames(x) <- c("x1", "x2")

# Generate p-values (one-sided z test)
# Set all hypotheses in the central circle with radius 30 to be
# non-nulls. For non-nulls, z~N(2,1) and for nulls, z~N(0,1).
H0 <- apply(x, 1, function(coord){sum(coord^2) < 900})
mu <- ifelse(H0, 2, 0)
set.seed(0)
zvals <- rnorm(n) + mu
pvals <- 1 - pnorm(zvals)

# Run adapt_gam with a 2d spline basis
library("mgcv")
formula <- "s(x1, x2)"
dist <- beta_family()
res <- adapt_gam(x = x, pvals = pvals, pi_formulas = formula,
                 mu_formulas = formula, dist = dist, nfits = 5)
adapt_glm

- `pi_formulas`: a vector/list of strings/formulas. Formulas for fitting \( \pi(x) \) by glm. See Details
- `mu_formulas`: a vector/list of strings/formulas. Formulas for fitting \( \mu(x) \) by glm. See Details
- `dist`: an object of class "gen_exp_family". `beta_family()` as default
- `s0`: a vector of values in \([0, 0.5)\). Initial threshold.
- `alphas`: a vector of values in \((0, 1)\). Target FDR levels.
- `piargs`: a list. Other arguments passed to glm for fitting \( \pi(x) \)
- `muargs`: a list. Other arguments passed to glm for fitting \( \mu(x) \)
- ... other arguments passed to adapt (except models)

**Details**

`pi_formulas` and `mu_formulas` can either be a list or a vector with each element being a string or a formula. For instance, suppose \( x \) has a single column with name \( x1 \), the following five options are valid for the same inputs (\( ns \) forms a spline basis with \( df \) knots):

1. \( c("x1", "ns(x1, df = 8)") \);
2. \( c("~ x1", "~ ns(x1, df = 8)") \);
3. \( list("x1", "ns(x1, df = 8)") \);
4. \( list("~ x1", "~ ns(x1, df = 8)") \);
5. \( list(~ x1, ~ ns(x1, df = 8)) \)

There is no need to specify the name of the response variable, as this is handled in the function.

When \( x \) has a few variables, it is common to use non-parametric GLM by replacing \( x \) by a spline basis of \( x \). In this case, \( ns \) from \texttt{library(splines)} package is suggested.

**See Also**

adapt, adapt_gam, adapt_glmnet, glm, ns

**Examples**

```r
# Load estrogen data
data(estrogen)
pvals <- as.numeric(estrogen$pvals)
x <- data.frame(x = as.numeric(estrogen$ord_high))
dist <- beta_family()

# Subsample the data for convenience
inds <- (x$x <= 5000)
pvals <- pvals[inds]
x <- x[inds,,drop = FALSE]

# Run adapt_glm
library("splines")
formulas <- paste0("ns(x, df = ", 6:10, ")")
res <- adapt_glm(x = x, pvals = pvals, pi_formulas = formulas,
                 mu_formulas = formulas, dist = dist, nfits = 10)
```
# Run adapt by manually setting models for glm
models <- lapply(formulas, function(formula){
  piargs <- muargs <- list(formula = formula)
  gen_adapt_model(name = "glm", piargs = piargs, muargs = muargs)
})
res2 <- adapt(x = x, pvals = pvals, models = models,
  dist = dist, nfits = 10)

# Check equivalence
identical(res, res2)

## adapt_glmnet

### Description

`adapt_glmnet` is a wrapper of `adapt` that fits pi(x) and mu(x) by `glmnet` from glmnet package.

### Usage

```r
adapt_glmnet(x, pvals, piargs = list(), muargs = list(),
  dist = beta_family(), s0 = rep(0.45, length(pvals)), alphas = seq(0.01, 1, 0.01), ...)
```

### Arguments

- **x**: covariates (i.e. side-information). Should be compatible to `models`. See Details
- **pvals**: a vector of values in [0, 1]. P-values
- **piargs**: a list. Other arguments passed to glmnet for fitting pi(x)
- **muargs**: a list. Other arguments passed to glmnet for fitting mu(x)
- **dist**: an object of class "gen_exp_family". `beta_family()` as default
- **s0**: a vector of values in [0, 0.5]. Initial threshold.
- **alphas**: a vector of values in (0, 1). Target FDR levels.
- **...**: other arguments passed to `adapt` (except `models`)

### Details

`adapt_glmnet` by default implements LASSO on x with lambda selected by cross-validation. Specify in `piargs` and `muargs` if ridge or elastic-net penalty is needed.

### See Also

`adapt, adapt_glm, adapt_gam, glmnet`
Examples

```r
# Generate a 100-dim covariate x
set.seed(0)
m <- 100
n <- 1000
x <- matrix(runif(n * m), n, m)

# Generate the parameters from a conditional two-group
# logistic-Gamma GLM where pi(x) and mu(x) are both
# linear in x. pi(x) has an intercept so that the average
# of pi(x) is 0.3
inv_logit <- function(x) {exp(x) / (1 + exp(x))}
pi1 <- 0.3
beta.pi <- c(3, 3, rep(0, m-2))
beta0.pi <- uniroot(function(b){
    mean(inv_logit(x %*% beta.pi + b)) - pi1
}, c(-100, 100))$root
pi <- inv_logit(x %*% beta.pi + beta0.pi)
beta.mu <- c(2, 2, rep(0, m-2))
beta0.mu <- 0
mu <- pmax(1, x %*% beta.mu + beta0.mu)

# Generate p-values
H0 <- as.logical(ifelse(runif(n) < pi, 1, 0))
y <- ifelse(H0, rexp(n, 1/mu), rexp(n, 1))
pvals <- exp(-y)

# Run adapt_glmnet
res <- adapt_glmnet(x, pvals, s0 = rep(0.15, n), nfits = 5)
```

corr_lfdr

Quantifying Information Loss of Adaptive P-Value Thresholding

Description
corr_lfdr computes the oracle local FDR estimate, by using revealing all p-values, and computes the Pearson correlation between it and the estimate within each step of adapt.

Usage
corr_lfdr(obj, x, pvals, model = NULL, niter_oracle = 100)

Arguments

- **obj**: an ‘adapt’ object. Output of `adapt` function
- **x**: covariates (i.e. side-information). Should be compatible to models.
- **pvals**: a vector of values in [0, 1]. P-values
model an optional argument. If model = NULL then the last model in obj$models is used for fitting the oracle model (i.e. with all p-values revealed). Otherwise it should be an ‘adapt_model’ object.

niter_oracle an positive integer. Number of iterations in EM algorithm.

Value

* corra vector of values in \([0, 1]\). Pearson correlation of oracle local FDR estimate and the estimates within each step. Each value corresponds to an entry of obj$params.

* oracle_lfdra vector of values in \([0, 1]\). Oracle local FDR estimate.

* lfdra matrix of values in \([0, 1]\). Local FDR estimates within each step.

* alphasa vector of values in \([0, 1]\). The target FDR levels corresponding to each local FDR estimate.

* nmasksa vector of integers. The number of masked p-values corresponding to each local FDR estimate.

Examples

```r
# Load estrogen data
data(estrogen)
pvals <- as.numeric(estrogen$pvals)
x <- data.frame(x = as.numeric(estrogen$order_high))
dist <- beta_family()

# Subsample the data for convenience
inds <- (x$x <= 5000)
pvals <- pvals[inds]
x <- x[inds,, drop = FALSE]

dist <- beta_family()
formulas <- paste0("ns(x, df = ", 6:10, ")")
res <- adapt_glm(x = x, pvals = pvals, pi_formulas = formulas,
                 mu_formulas = formulas, dist = dist, nfits = 10)

# Run corr_lfdr
obj <- corr_lfdr(res, x, pvals)
obj$corra
```

ctgm lfdr

Fitting Conditional Two-Groups Models on Unmasked P-Values

Description

ctgm lfdr computes the oracle local FDR estimate, by using all p-values without masking.
ctgm_lfdr

Usage

ctgm_lfdr(x, pvals, models, dist = beta_family(), type = c("over", "raw"),
       params0 = list(pix = NULL, mux = NULL), niter = 50, cr = "BIC",
       verbose = TRUE)

Arguments

x          covariates (i.e. side-information). Should be compatible to models. See Details
pvals      a vector of values in [0, 1]. P-values
models     an object of class "adapt_model" or a list of objects of class "adapt_model". See Details
dist       an object of class "gen_exp_family". beta_family() as default
type       a character. Either "over" or "raw" indicating the type of local FDR estimates. See Details
params0    a list in the form of list(pix = , mux = ). Initial values of pi(x) and mu(x). Both can be set as NULL
niter      a positive integer. Number of EM iterations.
cr         a string. The criterion for model selection with BIC as default. Also support AIC, AICC and HIC
verbose    a logical values in the form of list(fit = , ms = ). Indicate whether the progress of model fitting and model selection is displayed

Details

tctgm_lfdr implements the EM algorithm to fit pi(x) and mu(x) on unmasked p-values. Although it is not related to FDR control of AdaPT, it provides useful measures for post-hoc justification and other purposes. For instance, one can use these local FDR estimates for prioritizing the hypotheses if strict FDR control is not required.

In contrast to adapt, ctym_lfdr does not guarantee FDR control unless the model is correctly specified. It is recommended to use ctym_lfdr only when FDR control is not required.

x should have a type compatible to the fitting functions in models. For GLM and GAM, x should be a data.frame. For glmnet, x should be a matrix.

models could either be an adapt_model object, if a single model is used, or a list of adapt_model objects, each of which corresponding to a model. Each element should be generated by gen_adapt_model. For glm/gam/glmnet, one can use the shortcut by running gen_adapt_model with name = "glm" or "gam" or "glmnet" but without specifying pifun, mufun, pifun_init and mufun_init. See examples below.

When type = "over", it yields a conservative estimate of local FDR

\[ \text{lfdr}(p) = \frac{(1 - \pi_1 + \pi_1 f_1(1))(1 - \pi_1 + \pi_1 f_1(p))}{(1 - \pi_1 + \pi_1 f_1(1))}. \]

When type = "raw", it yields the original local FDR.

\[ \text{lfdr}(p) = \frac{(1 - \pi_1)}{(1 - \pi_1 + \pi_1 f_1(p))}. \]

The former is shown to be more stable and reliable because the weak identifiability in conditional mixture models.
Value

- lfdra vector of values in [0, 1]. Local FDR estimates of each hypothesis.
- modelan adapt_model object. The selected model if multiple models are provided.

Examples

```r
# Load estrogen data
data(estrogen)
pvals <- as.numeric(estrogen$pvals)
x <- data.frame(x = as.numeric(estrogen$ord_high))
dist <- beta_family()

# Subsample the data for convenience
inds <- (x$x <= 5000)
pvals <- pvals[inds]
x <- x[inds,,drop = FALSE]

# Generate models for function adapt
library("splines")
formulas <- paste0("ns(x, df = ", 6:10, ",")")
models <- lapply(formulas, function(formula){
  piargs <- muargs <- list(formula = formula)
  gen_adapt_model(name = "glm", piargs = piargs, muargs = muargs)
})

# Run ctgm_lfdr with two types of lfdr estimates
res_over <- ctgm_lfdr(x, pvals, models, type = "over")
res_raw <- ctgm_lfdr(x, pvals, models, type = "raw")

# Compare two estimates
par(mfrow = c(2, 1))
hist(res_over$lfdr)
hist(res_raw$lfdr)
```

Description

P-values and ordering of genes drawn from a microarray dataset, consisting of 22283 genes on breast cancer cells in response to estrogen, from NCBI Gene Expression Omnibus (GEO) through 'GEOquery' package, with index "GDS2324".

Usage

estrogen
Format

An object of class data.frame with 22283 rows and 3 columns.

Details

The original dataset "GDS2324" consists of gene expression measurements for \( n = 22283 \) genes, in response to estrogen treatments in breast cancer cells for five groups of patients, with different dosage levels and 5 trials in each. The task is to identify the genes responding to a low dosage. The p-value for gene \( i \) is obtained by a one-sided permutation test which evaluates evidence for a change in gene expression level between the control group (placebo) and the low-dose group. The p-values are then ordered according to permutation t-statistics comparing the control and low-dose data, pooled, against data from a higher dosage (with genes that appear to have a strong response at higher dosages placed earlier in the list).

Two orderings are considered: first, a stronger (more informative) ordering based on a comparison to the highest dosage; and second, a weaker (less informative) ordering based on a comparison to a medium dosage.

The variables are as follows:

- \( \text{pvals} \), p-values
- \( \text{ord\_high} \), stronger ordering
- \( \text{ord\_mod} \), weaker ordering

The R code to produce the data can be found in './extdata/estrogen_get_pvals.R'.
gen_adapt_model

muargs a list. Arguments for "mufun". An empty list as default
piargs_init a list. Arguments for piargs_init. An empty list as default
muargs_init a list. Arguments for muargs_init. An empty list as default
name a string. An optional argument for the user-specified name of the model. An empty string as default.

Details
pifun should be in the form of pifun(formula, data, family, weights, ...) or pifun(x, y, family, ...). The former includes glm and gam and the latter includes glmnet. The outputs should be in the form of list(fitv = , info = , ...) where fitv gives the estimate of pi(x), as a vector with the same order of x, and info should at least contain a key df if model selection is used, i.e. info = list(df = , ...)
mufun should be in the form of pifun(formula, data, family, weights, ...) or pifun(x, y, family, weights, ...). Note that mufun must take weights as an input. The outputs should be in the same form as pifun except that fitv should give the estimate of mu(x).
When pifun / mufun takes the form of (formula, family, ...), piargs / muargs should at least contain a key formula; when pifun / mufun takes the form of (x, y, family, ...), piargs / muargs can be empty.
For glm/gam/glmnet, one can use the shortcut by running gen_adapt_model with name = "glm" or "gam" or "glmnet" but without specifying pifun, mufun, pifun_init and mufun_init. See examples below.

Value
name same as the input name
algo a list recording pifun, mufun, pifun_init and mufun_init
args a list recording piargs, muargs, piargs_init and muargs_init

Examples

# Exemplary code to generate 'adapt_model' for logistic-Gamma glm with naive initialization.
# The real implementation in the package is much more complicated.

# pifun as a logistic regression
pifun <- function(formula, data, weights, ...){
  glm(formula, data, weights = weights, family = binomial(), ...)
}
# pifun_init as a constant
pifun_init <- function(x, pvals, s, ...){
  rep(0.1, length(pvals))
}
# mufun as a Gamma GLM
mufun <- function(formula, data, weights, ...){
  glm(formula, data, weights = weights, family = Gamma(), ...)
}
# mufun_init as a constant
mufun_init <- function(x, pvals, s, ...){
  rep(1.5, length(pvals))
}

library("splines") # for using ns() in the formula
piargs <- list(formula = "ns(x, df = 8)"
muargs <- list(formula = "ns(x, df = 8)"
name <- "glm"

mod <- gen_adapt_model(pifun, mufun, pifun_init, mufun_init,
piargs, muargs, name = name)
mod

# Using shortcut for GLM. See the last paragraph of Details.
mod2 <- gen_adapt_model(name = "glm", piargs = piargs, muargs = muargs)
mod2

---

### gen_exp_family

**Generate exp_family Objects for Exponential Families**

#### Description

exp_family objects contain all required information in an exponential family to perform the E-step. The exponential function is encoded by

\[
h(p; \mu) = \exp\{(\eta(\mu) - \eta(\mu^*))g(p) - (A(\mu) - A(\mu^*))\}
\]

where \(g(p)\) is an arbitrary transformation, \(\mu\) is the mean parameter, \(\eta\) is the natural parameter, and \(A(\mu)\) is the partition function. The extra redundant parameter \(\mu^*\) is to guarantee that \(U([0, 1])\) belongs to the class.

#### Usage

```r
gen_exp_family(g, ginv, eta, mustar, A, name = NULL, family = NULL)
```

beta_family()

inv_gaussian_family()

#### Arguments

- `g`: a function. An transformation of p-values
- `ginv`: a function. The inverse function of \(g\)
- `eta`: a function. The natural parameter as a function of the mean parameter \(\mu\)
- `mustar`: a scalar. The mean parameter that gives \(U([0, 1])\)
- `A`: a function. The partition function
name a string. A name for the family. NULL by default
family an object of class "family" from stats package. The family used for model fitting in \texttt{glm}, \texttt{gam}, \texttt{glmnet}, etc.

Details

Beta family (\texttt{beta_family()}): modeling p-values as Beta-distributed random variables, i.e. \( g(p) = -\log(p) \), \( \eta(\mu) = -1/\mu \), \( \mu^* = 1 \), \( A(\mu) = \log(\mu) \), name = "beta" and family = Gamma(). Beta-family is highly recommended for general problems and used as default.

Inverse-gaussian family (\texttt{inv_gaussian_family()}): modeling p-values as transformed z-scores, i.e. \( g(p) = \Phi^{-1}(p)(\Phi \text{is the c.d.f. of a standard normal random variable}) \), \( \eta(\mu) = \mu \), \( \mu^* = 0 \), \( A(\mu) = \mu^2/2 \), name = "inv_gaussian" and family = gaussian().

Value

an object of class "exp_family". This includes all inputs and \( h \), the density function.

---

**plot\_1d**

Plotting Functions for AdaPT with 1D Covariates

**Description**

Plotting the outputs of \texttt{adapt} when \( x \) is 1-dimensional, including threshold curves and level curves of local FDR.

**Usage**

\begin{verbatim}
plot_1d_thresh(obj, x, pvals, alpha, title, xlab = "x", xlim = NULL, disp_ymax = 0.2, num_yticks = 3, rand_seed_perturb = NA, ...)
plot_1d_lfdr(obj, x, pvals, alpha, title, xlab = "x", xlim = NULL, disp_ymax = 0.2, num_yticks = 3, legend_pos = "topright", ...)
\end{verbatim}

**Arguments**

- \texttt{obj} an 'adapt' object
- \texttt{x} covariates (i.e. side-information). Should be compatible to models and 1-dimensional.
- \texttt{pvals} a vector of values in \([0, 1]\). P-values
- \texttt{alpha} a positive scalar in \((0, 1)\). Target FDR level
- \texttt{title} a string. Title of the figure
- \texttt{xlab} a string. Label of the x-axis
- \texttt{xlim} a vector of length 2. Limits of x-axis
- \texttt{disp_ymax} a positive scalar in \([0, 1]\). Maximum value displayed in the y-axis
num_yticks  a positive integer. Number of ticks in the y-axis
rand_seed_perturb  random seed if jitter is added. NA if no jittering is needed
...  other arguments passed to par
legend_pos  a string. Position of the legend

Examples

# Load estrogen data
data(estrogen)
pvals <- as.numeric(estrogen$pvals)
x <- data.frame(x = as.numeric(estrogen$ord_high))
dist <- beta_family()

# Subsample the data for convenience
inds <- (x$x <= 5000)
pvals <- pvals[inds]
x <- x[inds,, drop = FALSE]

# Run adapt_glm
library("splines")
formulas <- paste0("ns(x, df = ", 6:10, ")")
res <- adapt_glm(x = x, pvals = pvals, pi_formulas = formulas,
                 mu_formulas = formulas, dist = dist, nfits = 10)

# Plots
par(mfrow = c(2, 1))
plot_1d_thresh(res, x, pvals, 0.1, "P-value Thresholds (alpha = 0.1)",
               disp_ymax = 0.5)
plot_1d_lfdr(res, x, pvals, 0.1, "Level Curves of lfdr (alpha = 0.1)",
             disp_ymax = 0.5)

plot_2d  

Plotting Functions for AdaPT with 2D Covariates

Description

Plotting the outputs of adapt when x is 2-dimensional, including threshold curves and level curves of local FDR.

Usage

plot_2d_thresh(obj, x, pvals, alpha, title, xlab = NULL, ylab = NULL,
               keyaxes = list(), ...)

plot_2d_lfdr(obj, x, pvals, alpha, title, targetp, xlab = NULL, ylab = NULL,
             keyaxes = list(), ...)
Arguments

obj an 'adapt' object
x covariates (i.e. side-information). Should be compatible to models and 2-dimensional.
pvals a vector of values in [0, 1]. P-values
alpha a positive scalar in (0, 1). Target FDR level
title a string. Title of the figure
xlab, ylab a string. Label of x/y-axis
keyaxes a list of arguments passed into axis. The graphical setting for the legend bar. An empty list by default
... other arguments passed to `par`
targetp a real in (0, 1). See Details

Details

The breaks in the legend of plot_2d_thresh correspond to the maximum, the 95
plot_2d_lfdr gives the contour plot of local FDR estimates when all p-values are equal to targetp.
It is recommended to run plot_2d_lfdr for multiple targetp's ranging from 0.001, 0.005, 0.01, 0.05.

Examples

```r
# Generate a 2-dim x
n <- 400
x1 <- x2 <- seq(-100, 100, length.out = 20)
x <- expand.grid(x1, x2)
colnames(x) <- c("x1", "x2")

# Generate p-values (one-sided z test)
# Set all hypotheses in the central circle with radius 30 to be 
# non-nulls. For non-nulls, z~N(2,1) and for nulls, z~N(0,1).
H0 <- apply(x, 1, function(coord){sum(coord^2) < 900})
mu <- ifelse(H0, 2, 0)
set.seed(0)
zvals <- rnorm(n) + mu
pvals <- 1 - pnorm(zvals)

# Run adapt_gam with a 2d spline basis
library("mgcv")
formula <- "s(x1, x2)"
dist <- beta_family()
res <- adapt_gam(x = x, pvals = pvals, pi_formulas = formula,
                 mu_formulas = formula, dist = dist, nfits = 5)

# Plots
plot_2d_thresh(res, x, pvals, 0.3, "P-value Thresholds (alpha = 0.3)"
plot_2d_lfdr(res, x, pvals, 0.3, "Local FDR Estimates (alpha = 0.3, p = 0.01)", 0.01)
```
Index

* datasets
  estrogen, 12
  adapt, 2, 5–9, 13
  adapt_gam, 5, 7, 8
  adapt_glm, 6, 6, 8
  adapt_glmnet, 6, 7, 8
  beta_family, 2, 5, 7, 8, 11
  beta_family (gen_exp_family), 15
  corr_lfdr, 9
  ctgm_lfdr, 10
  estrogen, 12
  family, 16
  gam, 5, 6, 14, 16
  gen_adapt_model, 3, 11, 13, 14
  gen_exp_family, 2, 5, 7, 8, 11, 15
  glm, 6, 7, 14, 16
  glmnet, 8, 14, 16
  inv_gaussian_family (gen_exp_family), 15
  ns, 5–7
  par, 17, 18
  plot_1d, 16
  plot_1d_lfdr (plot_1d), 16
  plot_1d_thresh (plot_1d), 16
  plot_2d, 17
  plot_2d_lfdr (plot_2d), 17
  plot_2d_thresh (plot_2d), 17
  s, 5, 6