Package ‘personalized’

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Type Package

Title Estimation and Validation Methods for Subgroup Identification and Personalized Medicine

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Description Provides functions for fitting and validation of models for subgroup identification and personalized medicine / precision medicine under the general subgroup identification framework of Chen et al. (2017) <doi:10.1111/biom.12676>. This package is intended for use for both randomized controlled trials and observational studies.


BugReports https://github.com/jaredhuling/personalized/issues

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Author Jared Huling [aut, cre] (&lt;https://orcid.org/0000-0003-0670-4845&gt;), Aaron Potvien [ctb], Alexandros Karatzoglou [cph], Alex Smola [cph]

Maintainer Jared Huling &lt;jaredhuling@gmail.com&gt;

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check.overlap  

Check propensity score overlap

Description

Results in a plot to check whether the propensity score has adequate overlap between treatment groups

Usage

check.overlap(x, trt, propensity.func, type = c("histogram", "density", "both"), bins = 50L, alpha = ifelse(type == "both", 0.35, 0.5))

Arguments

- **x**  
  The design matrix (not including intercept term)

- **trt**  
  Treatment vector with each element equal to a 0 or a 1, with 1 indicating treatment status is active.

- **propensity.func**  
  Function that inputs the design matrix x and the treatment vector trt and outputs the propensity score, i.e., \( \text{Pr}(\text{trt} = 1 \mid X = x) \). Function should take two arguments 1) x and 2) trt. See example below. For a randomized controlled trial this can simply be a function that returns a constant equal to the proportion of patients assigned to the treatment group, i.e., \( \text{propensity.func} = \text{function}(x, \text{trt}) 0.5 \).

- **type**  
  Type of plot to create. Options are either a histogram (type = "histogram") for each treatment group, a density (type = "density") for each treatment group, or to plot both a density and histogram (type = "code")

- **bins**  
  Integer number of bins for histograms when type = "histogram"
check.overlap

alpha value between 0 and 1 indicating transparency level (1 for solid, 0 for fully transparent)

Examples

library(personalized)

set.seed(123)
n.obs <- 250
n.vars <- 15
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.25 + 0.5 * x[,11] - 0.5 * x[,12]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
  # fit propensity score model
  propens.model <- cv.glmnet(y = trt,
                            x = x, family = "binomial")
  pi.x <- predict(propens.model, s = "lambda.min",
                  newx = x, type = "response")[,1]
  pi.x
}

check.overlap(x = x,
               trt = trt01,
               propensity.func = prop.func)

# now add density plot with histogram
check.overlap(x = x,
               trt = trt01,
               type = "both",
               propensity.func = prop.func)

# simulated non-randomized treatment with multiple levels
xbetat_1 <- 0.15 + 0.5 * x[,9] - 0.25 * x[,12]
xbetat_2 <- 0.15 - 0.5 * x[,11] + 0.25 * x[,15]
trt.1.prob <- exp(xbetat_1) / (1 + exp(xbetat_1) + exp(xbetat_2))
trt.2.prob <- exp(xbetat_2) / (1 + exp(xbetat_1) + exp(xbetat_2))
trt.3.prob <- 1 - (trt.1.prob + trt.2.prob)
prob.mat <- cbind(trt.1.prob, trt.2.prob, trt.3.prob)
trt <- apply(prob.mat, 1, function(rr) rmultinom(1, 1, prob = rr))
trt <- apply(trt, 2, function(rr) which(rr == 1))

# use multinomial logistic regression model with lasso penalty for propensity
propensity.multinom.lasso <- function(x, trt)
{

fit.subgroup

Fitting subgroup identification models

Description


Usage


if (!is.factor(trt)) trt <- as.factor(trt)
gfit <- cv.glmnet(y = trt, x = x, family = "multinomial")

# predict returns a matrix of probabilities:
# one column for each treatment level
propens <- drop(predict(gfit, newx = x, type = "response", s = "lambda.min", nfolds = 5, alpha = 0))

# return the probability corresponding to the
# treatment that was observed
probs <- propens[,match(levels(trt), colnames(propens))]

probs

check.overlap(x = x,
trt = trt,
type = "histogram",
propensity.func = propensity.multinom.lasso)
Arguments

- **x**: The design matrix (not including intercept term)
- **y**: The response vector
- **trt**: Treatment vector with each element equal to a 0 or a 1, with 1 indicating treatment status is active.
- **propensity.func**: Function that inputs the design matrix x and the treatment vector trt and outputs the propensity score, i.e., \( \text{Pr}(\text{trt} = 1 | X = x) \). Function should take two arguments 1) x and 2) trt. See example below. For a randomized controlled trial this can simply be a function that returns a constant equal to the proportion of patients assigned to the treatment group, i.e.: propensity.func = function(x, trt) 0.5.
- **loss**: Choice of both the M function from Chen, et al (2017) and potentially the penalty used for variable selection. All loss options starting with `sq_loss` use \( M(y, v) = (v - y)^2 \), all options starting with `logistic_loss` use the logistic loss: \( M(y, v) = y \log(1 + \exp{-v}) \), and all options starting with `cox_loss` use the negative partial likelihood loss for the Cox PH model. All options ending with `lasso` have a lasso penalty added to the loss for variable selection. `sq_loss_lasso_gam` and `logistic_loss_lasso_gam` first use the lasso to select variables and then fit a generalized additive model with nonparametric additive terms for each selected variable. `sq_loss_gam` involves a squared error loss with a generalized additive model and no variable selection. `sq_loss_gbm` involves a squared error loss with a gradient-boosted decision trees model for the benefit score; this allows for flexible estimation using machine learning and can be useful when the underlying treatment-covariate interaction is complex.

- **Continuous Outcomes**
  - "sq_loss_lasso" - \( M(y, v) = (v - y)^2 \) with linear model and lasso penalty
  - "owl_logistic_loss_lasso" - \( M(y, v) = y \log(1 + \exp{-v}) \) (method of Regularized Outcome Weighted Subgroup Identification)
  - "owl_logistic_flip_loss_lasso" - \( M(y, v) = y \log(1 + \exp{-|y|v}) \)
  - "owl_hinge_loss" - \( M(y, v) = \max(0, 1 - v) \) (method of Estimating individualized treatment rules using outcome weighted learning)
  - "owl_hinge_flip_loss" - \( M(y, v) = \max(0, 1 - |y|v) \)
  - "sq_loss_lasso_gam" - \( M(y, v) = (v - y)^2 \) with variables selected by lasso penalty and generalized additive model fit on the selected variables
  - "sq_loss_gam" - \( M(y, v) = (v - y)^2 \) with generalized additive model fit on all variables
  - "owl_logistic_loss_gam" - \( M(y, v) = y \log(1 + \exp{-v}) \) with generalized additive model fit on all variables
  - "owl_logistic_flip_loss_gam" - \( M(y, v) = y \log(1 + \exp{-|y|v}) \) with generalized additive model fit on all variables
  - "owl_logistic_loss_lasso_gam" - \( M(y, v) = y \log(1 + \exp{-v}) \) with variables selected by lasso penalty and generalized additive model fit on the selected variables
- "owl_logistic_flip_loss_lasso_gam" - $M(y, v) = \frac{|y|\log(1 + \exp{-\text{sign}(y)v})}{v}$ with variables selected by lasso penalty and generalized additive model fit on the selected variables
- "sq_loss_gbm" - $M(y, v) = (v - y)^2$ with gradient-boosted decision trees model

**Binary Outcomes**
- All losses for continuous outcomes can be used plus the following:
  - "logistic_loss_lasso" - $M(y, v) = -[yv - \log(1 + \exp{-v})]$ with linear model and lasso penalty
  - "logistic_loss_lasso_gam" - $M(y, v) = -[yv - \log(1 + \exp{-v})]$ with variables selected by lasso penalty and generalized additive model fit on the selected variables
  - "logistic_loss_gam" - $M(y, v) = -[yv - \log(1 + \exp{-v})]$ with generalized additive model fit on all variables
  - "logistic_loss_gbm" - $M(y, v) = -[yv - \log(1 + \exp{-v})]$ with gradient-boosted decision trees model

**Count Outcomes**
- All losses for continuous outcomes can be used plus the following:
  - "poisson_loss_lasso" - $M(y, v) = -[yv - \exp(v)]$ with linear model and lasso penalty
  - "poisson_loss_lasso_gam" - $M(y, v) = -[yv - \exp(v)]$ with variables selected by lasso penalty and generalized additive model fit on the selected variables
  - "poisson_loss_gam" - $M(y, v) = -[yv - \exp(v)]$ with generalized additive model fit on all variables
  - "poisson_loss_gbm" - $M(y, v) = -[yv - \exp(v)]$ with gradient-boosted decision trees model

**Time-to-Event Outcomes**
- "cox_loss_lasso" - $M$ corresponds to the negative partial likelihood of the cox model with linear model and additionally a lasso penalty
- "cox_loss_gbm" - $M$ corresponds to the negative partial likelihood of the cox model with gradient-boosted decision trees model

`method` subgroup ID model type. Either the weighting or A-learning method of Chen et al., (2017)

`match.id` a (character, factor, or integer) vector with length equal to the number of observations in `x` indicating using integers or levels of a factor vector which patients are in which matched groups. Defaults to NULL and assumes the samples are not from a matched cohort. Matched case-control groups can be created using any method (propensity score matching, optimal matching, etc). If each case is matched with a control or multiple controls, this would indicate which case-control pairs or groups go together. If `match.id` is supplied, then it is unnecessary to specify a function via the `propensity.func` argument. A quick usage example: if the first patient is a case and the second and third are controls matched to it, and the fourth patient is a case and the fifth through seventh patients are matched with it, then the user should specify `match.id = c(1,1,1,2,2,2,2)` or `match.id = c(rep("Grp1", 3),rep("Grp2", 4))`
augment.func function which inputs the response y, the covariates x, and trt and outputs predicted values (on the link scale) for the response using a model constructed with x. augment.func() can also be simply a function of x and y. This function is used for efficiency augmentation. When the form of the augmentation function is correct, it can provide efficient estimation of the subgroups. Some examples of possible augmentation functions are:

Example 1: augment.func <- function(x, y) {lmod <- lm(y ~ x); return(fitted(lmod))}

Example 2:

```r
augment.func <- function(x, y, trt) {
  data <- data.frame(x, y, trt)
  lmod <- lm(y ~ x * trt)
  ## get predictions when trt = 1
  data$trt <- 1
  preds_1 <- predict(lmod, data)
  ## get predictions when trt = -1
  data$trt <- -1
  preds_n1 <- predict(lmod, data)
  ## return predictions averaged over trt
  return(0.5 * (preds_1 + preds_n1))
}
```

For binary and time-to-event outcomes, make sure that predictions are returned on the scale of the predictors

Example 3:

```r
augment.func <- function(x, y) {
  bmod <- glm(y ~ x, family = binomial())
  return(predict(bmod, type = "link"))
}
```

fit.custom.loss A function which minimizes a user-specified custom loss function M(y,v) to be used in model fitting. If provided, fit.custom.loss should take the modified design matrix (which includes an intercept term) as an argument and the responses and optimize a custom weighted loss function.

The loss function M(y, v) to be minimized MUST meet the following two criteria:

1. \( D_M(y, v) = \partial M(y, v) / \partial v \) must be increasing in v for each fixed y. \( D_M(y, v) \) is the partial derivative of the loss function M(y, v) with respect to v
2. \( D_M(y, 0) \) is monotone in y

An example of a valid loss function is \( M(y, v) = (y - v)^2 \). In this case \( D_M(y, v) = -2(y - v) \). See Chen et al. (2017) for more details on the restrictions on the loss function M(y, v).

The provided function MUST return a list with the following elements:
predict a function that inputs a design matrix and a 'type' argument for
the type of predictions and outputs a vector of predictions on the scale
of the linear predictor. Note that the matrix provided to 'fit.custom.loss'
has a column appended to the first column of x corresponding to the treat-
ment main effect. Thus, the prediction function should deal with this, e.g.
\[ \text{predict(model, cbind(1, x))} \]
• model a fitted model object returned by the underlying fitting function
• coefficients if the underlying fitting function yields a vector of coeffi-
cient estimates, they should be provided here

The provided function **MUST** be a function with the following arguments:

1. x design matrix
2. y vector of responses
3. weights vector for observations weights. The underlying loss function
**MUST** have samples weighted according to this vector. See below example
4. ... additional arguments passed via '...'. This can be used so that users
can specify more arguments to the underlying fitting function if so desired.

The provided function can also optionally take the following arguments:

• match.id vector of case/control cluster IDs. This is useful if cross valida-
tion is used in the underlying fitting function in which case it is advisable
to sample whole clusters randomly instead of individual observations.
• offset if efficiency augmentation is used, the predictions from the out-
come model from augment.func will be provided via the offset argu-
ment, which can be used as an offset in the underlying fitting function as a
means of incorporating the efficiency augmentation model’s predictions
• trt vector of treatment statuses
• family family of outcome
• n.trts numer of treatment levels. Can be useful if there are more than 2
treatment levels

Example 1: Here we minimize \( M(y, v) = (y - v)^2 \)

```r
fit.custom.loss <- function(x, y, weights, ...) {
  df <- data.frame(y = y, x)

  # minimize squared error loss with NO lasso penalty
  lmf <- lm(y ~ x - 1, weights = weights,
           data = df, ...)

  # save coefficients
  cfs = coef(lmf)

  # create prediction function. Notice
  # how a column of 1's is appended
  # to ensure treatment main effects are included
  # in predictions
  prd = function(x, type = "response") {
    
    # additional code here ...
  }
```
```r
dfte <- cbind(1, x)
colnames(dfte) <- names(cfs)
predict(lmf, data.frame(dfte))

# return list of required components
list(predict = prd, model = lmf, coefficients = cfs)
```

**Example 2:** \( M(y, v) = y \exp(-v) \)

```r
fit.expo.loss <- function(x, y, weights, ...)
{
    ## define loss function to be minimized
    expo.loss <- function(beta, x, y, weights)
    { sum(weights * y * exp(-drop(tcrossprod(x, t(beta)))))
    }

    # use optim() to minimize loss function
    opt <- optim(rep(0, NCOL(x)), fn = expo.loss, x = x, y = y, weights = weights)

    coefs <- opt$par

    pred <- function(x, type = "response")
    { tcrossprod(cbind(1, x), t(coefs))
    }

    # return list of required components
    list(predict = pred, model = opt, coefficients = coefs)
}
```

**cutpoint** numeric value for patients with benefit scores above which (or below which if larger.outcome.better = FALSE) will be recommended to be in the treatment group. Can also set cutpoint = "median", which will use the median value of the benefit scores as the cutpoint or can set specific quantile values via "quantx" where "x" is a number between 0 and 100 representing the quantile value; e.g. cutpoint = "quant75" will use the 75th percent upper quantile of the benefit scores as the quantile.

**larger.outcome.better** boolean value of whether a larger outcome is better/preferable. Set to TRUE if a larger outcome is better/preferable and set to FALSE if a smaller outcome is better/preferable. Defaults to TRUE.

**reference.trt** which treatment should be treated as the reference treatment. Defaults to the first level of trt if trt is a factor or the first alphabetical or numerically first treatment level. Not used for multiple treatment fitting with OWL-type losses.

**retcall** boolean value. if TRUE then the passed arguments will be saved. Do not set to FALSE if the validate.subgroup() function will later be used for your fitted subgroup model. Only set to FALSE if memory is limited as setting to TRUE saves the design matrix to the fitted object
... options to be passed to underlying fitting function. For all loss options with 'lasso', this will be passed to `cv.glmnet`. For all loss options with 'gam', this will be passed to `gam` from the `mgcv` package. Note that for all loss options that use `gam()` from the `mgcv` package, the user cannot supply the `gam` argument method because it is also an argument of `fit.subgroup`, so instead, to change the `gam` method argument, supply `method.gam`, i.e. `method.gam = "REML"`. For all loss options with 'hinge', this will be passed to both `weighted.ksvm` and `ipop` from the `kernlab` package.

Value

An object of class "subgroup_fitted".

- **predict**: A function that returns predictions of the covariate-conditional treatment effects
- **model**: An object returned by the underlying fitting function used. For example, if the lasso use used to fit the underlying subgroup identification model, this will be an object returned by `cv.glmnet`.
- **coefficients**: If the underlying subgroup identification model is parametric, `coefficients` will contain the estimated coefficients of the model.
- **call**: The call that produced the returned object. If `retcall = TRUE`, this will contain all objects supplied to `fit.subgroup()`.
- **family**: The family corresponding to the outcome provided.
- **loss**: The loss function used.
- **method**: The method used (either weighting or A-learning).
- **propensity.func**: The propensity score function used.
- **larger.outcome.better**: If larger outcomes are preferred for this model.
- **cutpoint**: Benefit score cutoff value used for determining subgroups.
- **var.names**: The names of all variables used.
- **n.trts**: The number of treatment levels.
- **comparison.trts**: All treatment levels other than the reference level.
- **reference.trt**: The reference level for the treatment. This should usually be the control group/level.
- **trts**: All treatment levels.
- **trt.received**: The vector of treatment assignments.
- **pi.x**: A vector of propensity scores.
- **y**: A vector of outcomes.
- **benefit.scores**: A vector of conditional treatment effects, i.e. benefit scores.
- **recommended.trts**: A vector of treatment recommendations (i.e. for each patient, which treatment results in the best expected potential outcomes).
subgroup.trt.effects
(Biased) estimates of the conditional treatment effects and conditional outcomes.
These are essentially just empirical averages within different combinations of
treatment assignments and treatment recommendations

individual.trt.effects
estimates of the individual treatment effects as returned by treat.effects

References


See Also

validate.subgroup for function which creates validation results for subgroup identification models, predict.subgroup_fitted for a prediction function for fitted models from fit.subgroup, plot.subgroup_fitted for a function which plots results from fitted models, and print.subgroup_fitted for arguments for printing options for fit.subgroup(). from fit.subgroup.

Examples

library(personalized)

set.seed(123)
n.obs <- 500
n.vars <- 15
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,7] - 0.5 * x[,9]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)

trt <- 2 * trt01 - 1

# simulate response
# delta below drives treatment effect heterogeneity
delta <- 2 * (0.5 + x[,2] - x[,3] - x[,11] + x[,1] * x[,12])
xbeta <- x[,1] + x[,11] - 2 * x[,12]^2 + x[,13] + 0.5 * x[,15] ^ 2
xbeta <- xbeta + delta * trt

# continuous outcomes
y <- drop(xbeta) + rnorm(n.obs, sd = 2)

# binary outcomes
y.binary <- 1 * (xbeta + rnorm(n.obs, sd = 2) > 0)

# count outcomes
y.count <- round(abs(xbeta + rnorm(n.obs, sd = 2)))

# time-to-event outcomes
surv.time <- exp(-20 - xbeta + rnorm(n.obs, sd = 1))
cens.time <- exp(rnorm(n.obs, sd = 3))
y.time.to.event <- pmin(surv.time, cens.time)
status <- 1 * (surv.time <= cens.time)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
  # fit propensity score model
  propens.model <- cv.glmnet(y = trt,
                             x = x, family = "binomial")
  pi.x <- predict(propens.model, s = "lambda.min",
                  newx = x, type = "response")[,1]
  pi.x
}

############################################################
# Continuous outcomes
############################################################

subgrp.model <- fit.subgroup(x = x, y = y,
                             trt = trt01,
                             propensity.func = prop.func,
                             loss = "sq_loss_lasso",
                             nfolds = 10)  # option for cv.glmnet

summary(subgrp.model)

# estimates of the individual-specific treatment effect estimates:
subgrp.model$individual.trt.effects

# fit lasso + gam model with REML option for gam

subgrp.modelg <- fit.subgroup(x = x, y = y,
                               trt = trt01,
                               propensity.func = prop.func,
                               loss = "sq_loss_lasso_gam",
                               method.gam = "REML",  # option for gam
                               nfolds = 5)  # option for cv.glmnet

subgrp.modelg
### Using an augmentation function

Augmentation functions involve modeling the conditional mean $E[Y|T, X]$ and returning predictions that are averaged over the treatment values.

```r
augment.fnc <- function(x, y, trt) {
  data <- data.frame(x, y, trt)
  xm <- model.matrix(y~trt*x-1, data = data)

  lmod <- cv.glmnet(y = y, x = xm)
  ## get predictions when trt = 1
  data$trt <- 1
  xm <- model.matrix(y-trt*x-1, data = data)
  preds_1 <- predict(lmod, xm, s = "lambda.min")

  ## get predictions when trt = -1
  data$trt <- -1
  xm <- model.matrix(y-trt*x-1, data = data)
  preds_n1 <- predict(lmod, xm, s = "lambda.min")

  ## return predictions averaged over trt
  return(0.5 * (preds_1 + preds_n1))
}
```

```r
subgrp.model.aug <- fit.subgroup(x = x, y = y,
  trt = trt01,
  propensity.fnc = prop.fnc,
  augment.fnc = augment.fnc,
  loss = "sq_loss_lasso",
  nfolds = 10) # option for cv.glmnet
```

### Binary outcomes

Use logistic loss for binary outcomes.

```r
subgrp.model.bin <- fit.subgroup(x = x, y = y.binary,
  trt = trt01,
  propensity.fnc = prop.fnc,
  loss = "logistic_loss_lasso",
  type.measure = "auc", # option for cv.glmnet
  nfolds = 5) # option for cv.glmnet
```

### Count outcomes

Use poisson loss for count/poisson outcomes.
subgrp.model.poisson <- fit.subgroup(x = x, y = y.count,
  trt = trt01,
  propensity.func = prop.func,
  loss = "poisson_loss_lasso",
  type.measure = "mse", # option for cv.glmnet
  nfocus = 5) # option for cv.glmnet

subgrp.model.poisson

########################### Time-to-event outcomes ###########################

library(survival)

subgrp.model.cox <- fit.subgroup(x = x, y = Surv(y.time.to.event, status),
  trt = trt01,
  propensity.func = prop.func,
  loss = "cox_loss_lasso",
  nfocus = 5) # option for cv.glmnet

subgrp.model.cox

########################### Using custom loss functions ###########################

## Use custom loss function for binary outcomes

fit.custom.loss.bin <- function(x, y, weights, offset, ...) {
  df <- data.frame(y = y, x)

  # minimize logistic loss with NO lasso penalty
  # with allowance for efficiency augmentation
  glmf <- glm(y ~ x - 1, weights = weights,
              offset = offset, # offset term allows for efficiency augmentation
              family = binomial(), ...)

  # save coefficients
  cfs = coef(glmf)

  # create prediction function.
  prd = function(x, type = "response") {
    dfte <- cbind(1, x)
    colnames(dfte) <- names(cfs)
    ## predictions must be returned on the scale
    ## of the linear predictor
    predict(glmf, data.frame(dfte), type = "link")
  }

  # return list of required components
  list(predict = prd, model = glmf, coefficients = cfs)
}
LaLonde

LaLonde <- function(x, y, weights, ...) {
  expo.loss <- function(beta, x, y, weights) {
    sum(weights * y * exp(-drop(x * beta)))
  }

  # use optim() to minimize loss function
  opt <- optim(rep(0, NCOL(x)), fn = expo.loss, x = x, y = y, weights = weights)

  coefs <- opt$par

  pred <- function(x, type = "response") {
    tcrossprod(cbind(1, x), t(coefs))
  }

  # return list of required components
  list(predict = pred, model = opt, coefficients = coefs)
}

# use exponential loss for positive outcomes
subgrp.model.expo <- fit.subgroup(x = x, y = y.count,
                                  trt = trt01,
                                  propensity.func = prop.func,
                                  fit.custom.loss = fit.expo.loss)

subgrp.model.expo

LaLonde

National Supported Work Study Data

Description

The LaLonde dataset comes from the National Supported Work Study, which sought to evaluate the effectiveness of an employment training program on wage increases.
Usage

LaLonde

Format

A data frame with 722 observations and 12 variables:

outcome whether earnings in 1978 are larger than in 1975; 1 for yes, 0 for no

Treat whether the individual received the treatment; "Yes" or "No"

Age age in years

Educ education in years

Black black or not; factor with levels "Yes" or "No"

Hisp hispanic or not; factor with levels "Yes" or "No"

White white or not; factor with levels "Yes" or "No"

Marr married or not; factor with levels "Yes" or "No"

NoDegree No high school degree; factor with levels "Yes" (for no HS degree) or "No"

LogRe75 log of earnings in 1975

U75 unemployed in 1975; factor with levels "Yes" or "No"

WtsExtrap extrapolation weights to the 1978 Panel Study for Income Dynamics dataset

Source

The National Supported Work Study.

References


Examples

data(LaLonde)
y <- LaLonde$outcome

trt <- LaLonde$treat

ox.varnames <- c("age", "educ", "black", "hisp", "white", "marr", "nodegr", "log.re75", "u75")

# covariates
data.x <- LaLonde[, x.varnames]

# construct design matrix (with no intercept)
x <- model.matrix(~ -1 + ., data = data.x)
const.propens <- function(x, trt)
{
  mean.trt <- mean(trt == "Trt")
  rep(mean.trt, length(trt))
}

subgrp_fit_w <- fit.subgroup(x = x, y = y, trt = trt,
  loss = "logistic_loss_lasso",
  propensity.func = const.propens,
  cutpoint = 0,
  type.measure = "auc",
  nfolds = 10)

summary(subgrp_fit_w)

plot.subgroup_fitted  Plotting results for fitted subgroup identification models

Description

Plots results for estimated subgroup treatment effects
Plots validation results for estimated subgroup treatment effects

Usage

## S3 method for class 'subgroup_fitted'
plot(x, type = c("boxplot", "density",
  "interaction", "conditional"), avg.line = TRUE, ...)

## S3 method for class 'subgroup_validated'
plot(x, type = c("boxplot", "density",
  "interaction", "conditional", "stability"), avg.line = TRUE, ...)

Arguments

- `x`  
  fitted object returned by `validate.subgroup()` or `fit.subgroup()` function
- `type`  
  type of plot. "density" results in a density plot for the results across all observations (if `x` is from `fit.subgroup()`) or if `x` is from `validate.subgroup()` across iterations of either the bootstrap or training/test re-fitting. For the latter case the test results will be plotted. "boxplot" results in boxplots across all observations/iterations of either the bootstrap or training/test re-fitting. For the latter case the test results will be plotted. "interaction" creates an interaction plot for the different subgroups (crossing lines here means a meaningful subgroup). "conditional" For subgroup_fitted objects, plots smoothed (via a GAM smoother) means of the outcomes as a function of the estimated benefit score separately for the treated and untreated groups. For subgroup_validated objects, boxplots of summary statistics within subgroups will be plotted as subgroups are defined by different cutoffs of the benefit scores. These cutoffs can be specified via the `benefit.score.quantiles` argument of `validate.subgroup`. 
avg.line: boolean value of whether or not to plot a line for the average value in addition to the density (only valid for type = "density")

- not used

See Also

- `fit.subgroup` for function which fits subgroup identification models.
- `validate.subgroup` for function which creates validation results and `fit.subgroup` for function which fits subgroup identification models.

Examples

```r
library(personalized)

set.seed(123)
n.obs <- 250
n.vars <- 15
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,1] - 0.5 * x[,13]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)

trt <- 2 * trt01 - 1

delta <- 2 * (0.5 + x[,2] - x[,3] - x[,1] + x[,1] * x[,12])
xbeta <- x[,1] + x[,11] - 2 * x[,12]^2 + x[,13]
xbeta <- xbeta + delta * trt

# continuous outcomes
y <- drop(xbeta) + rnorm(n.obs, sd = 2)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
  # fit propensity score model
  propens.model <- cv.glmnet(y = trt,
                             x = x, family = "binomial")
  pi.x <- predict(propens.model, s = "lambda.min",
                  newx = x, type = "response")[,1]

  pi.x
}

subgrp.model <- fit.subgroup(x = x, y = y,
                             trt = trt01,
                             propensity.func = prop.func,
                             loss = "sq_loss_lasso",
                             nfolds = 5)  # option for cv.glmnet
```
subgrp.model$subgroup.trt.effects

plot(subgrp.model)

plot(subgrp.model, type = "boxplot")

plot(subgrp.model, type = "interaction")

plot(subgrp.model, type = "conditional")

valmod <- validate.subgroup(subgrp.model, B = 3, 
  method = "training_test", 
  benefit.score.quantiles = c(0.25, 0.5, 0.75), 
  train.fraction = 0.75)

valmod$avg.results

plot(valmod)

plot(valmod, type = "interaction")

# see how summary statistics of subgroups change
# when the subgroups are defined based on different cutoffs
# (25th quantile of bene score, 50th, and 75th)
plot(valmod, type = "conditional")

# visualize the frequency of particular variables
# of being selected across the resampling iterations with
# 'type = "stability"
# not run:
# plot(valmod, type = "stability")

---

**plotCompare**

Plot a comparison results for fitted or validated subgroup identification models

---

**Description**

Plots comparison of results for estimated subgroup treatment effects

**Usage**

```r
plotCompare(..., type = c("boxplot", "density", "interaction", 
  "conditional"), avg.line = TRUE)
```

**Arguments**

- `...`: the fitted (model or validation) objects to be plotted. Must be either objects returned from `fit.subgroup()` or `validate.subgroup()`
plotCompare

**type**

type of plot. "density" results in a density plot for the results across all observations (if \( x \) is from \texttt{fit.subgroup()} or if \( x \) is from \texttt{validate.subgroup()} across iterations of either the bootstrap or training/test re-fitting. For the latter case the test results will be plotted. "boxplot" results in boxplots across all observations/iterations of either the bootstrap or training/test re-fitting. For the latter case the test results will be plotted. "interaction" creates an interaction plot for the different subgroups (crossing lines here means a meaningful subgroup). "conditional" plots smoothed (via a GAM smoother) means of the outcomes as a function of the estimated benefit score separately for the treated and untreated groups.

**avg.line**

boolean value of whether or not to plot a line for the average value in addition to the density (only valid for type = "density")

**See Also**

\texttt{fit.subgroup} for function which fits subgroup identification models and \texttt{validate.subgroup} for function which creates validation results.

**Examples**

```r
library(personalized)
set.seed(123)
n.obs <- 100
n.vars <- 15
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,1] - 0.5 * x[,4]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)

trt <- 2 * trt01 - 1

# simulate response
delta <- 2 * (0.5 + x[,2] - x[,3] - x[,11] + x[,1] * x[,12])
xbeta <- x[,1] + x[,11] - 2 * x[,12]^2 + x[,13]
xbeta <- xbeta + delta * trt

# continuous outcomes
y <- drop(xbeta) + rnorm(n.obs, sd = 2)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
  # fit propensity score model
  propens.model <- cv.glmnet(y = trt,
                            x = x, family = "binomial")
  pi.x <- predict(propens.model, s = "lambda.min",
                  newx = x, type = "response")[,1]
```
predict.subgroup_fitted

Function to predict either benefit scores or treatment recommendations

Description
Predicts benefit scores or treatment recommendations based on a fitted subgroup identification model.
Function to obtain predictions for weighted ksvm objects.

Usage

## S3 method for class 'subgroup_fitted'
predict(object, newx, type = c("benefit.score", "trt.group"), cutpoint = 0, ...)

## S3 method for class 'wksvm'
predict(object, newx, type = c("class", "linear.predictor"), ...)

Arguments

- **object**: fitted object returned by validate.subgrp() function.
  For predict.wksvm(), this should be a fitted wksvm object from the weighted.ksvm() function.
- **newx**: new design matrix for which predictions will be made.
type of prediction. type = "benefit.score" results in predicted benefit scores and type = "trt.group" results in prediction of recommended treatment group.

For predict.wksvm(), type = 'class' yields predicted class and type = 'linear.predictor' yields estimated function (the sign of which is the estimated class)

cutpoint numeric value for patients with benefit scores above which (or below which if larger.outcome.better = FALSE) will be recommended to be in the treatment group. Can also set cutpoint = "median", which will use the median value of the benefit scores as the cutpoint or can set specific quantile values via "quantx" where "x" is a number between 0 and 100 representing the quantile value; e.g. cutpoint = "quant75" will use the 75th percent upper quantile of the benefit scores as the quantile.

See Also

fit.subgroup for function which fits subgroup identification models.

weighted.ksvm for fitting weighted.ksvm objects

Examples

library(personalized)

set.seed(123)
n.obs <- 1000
n.vars <- 50
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,21] - 0.5 * x[,41]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)

trt <- 2 * trt01 - 1

# simulate response
delta <- 2 * (0.5 + x[,2] - x[,3] - x[,11] + x[,1] * x[,12])
xbeta <- x[,1] + x[,11] - 2 * x[,12] 2 + x[,13]
xbeta <- xbeta + delta * trt

# continuous outcomes
ty <- drop(xbeta) + rnorm(n.obs, sd = 2)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
    # fit propensity score model
    propens.model <- cv.glmnet(y = trt,
                               x = x, family = "binomial")
print.individual_treatment_effects

pi.x <- predict(propens.model, s = "lambda.min", 
   newx = x, type = "response")[,1]
pi.x
}

subgrp.model <- fit.subgroup(x = x, y = y, 
   trt = trt01, 
   propensity.fnc = prop.fnc, 
   loss = "sq.loss.lasso", 
   nfolds = 5)  # option for cv.glmnet

subgrp.model$subgroup.trt.effects
benefit.scores <- predict(subgrp.model, newx = x, type = "benefit.score")
rec.trt.grp <- predict(subgrp.model, newx = x, type = "trt.group")

print.individual_treatment_effects

Printing individualized treatment effects

Description

Prints results for estimated subgroup treatment effects

Usage

## S3 method for class 'individual_treatment_effects'
print(x, 
   digits = max(getOption("digits") - 3, 3), ...)

Arguments

x a fitted object from either treat.effects or treatment.effects
digits minimal number of significant digits to print.
... further arguments passed to or from print.default.

print.subgroup_fitted

Printing results for fitted subgroup identification models

Description

Prints results for estimated subgroup treatment effects
Prints summary results for estimated subgroup treatment effects
Usage

```r
## S3 method for class 'subgroup_fitted'
print(x, digits = max(getOption("digits") - 3, 3), ...)
```

```r
## S3 method for class 'subgroup_validated'
print(x, digits = max(getOption("digits") - 3, 3), sample.pct = FALSE, which.quant = NULL, ...)
```

```r
## S3 method for class 'subgroup_summary'
print(x, p.value = 0.001, digits = max(getOption("digits") - 3, 3), ...)
```

Arguments

- `x`: a fitted object from either `fit.subgroup`, `validate.subgroup`, or `summarize.subgroups()`
- `digits`: minimal number of significant digits to print.
- `...`: further arguments passed to or from `print.default`.
- `sample.pct`: boolean variable of whether to print the percent of the test sample within each subgroup. If false the sample size itself, not the percent is printed. This may not be informative if the test sample size is much different from the total sample size.
- `which.quant`: when `validate.subgroup()` is called with a vector of quantile values specified for `benefit.score.quantiles`, i.e. `benefit.score.quantiles = c(0.25, 0.5, 0.75)`, the argument `which.quant` can be a vector of indexes specifying which quantile cutoff value validation results to display, i.e. `which.quant = c(1,3)` in the above example results in the display of validation results for subgroups defined by cutoff values of the benefit score defined by the 25th and 75th quantiles of the benefit score.
- `p.value`: a p-value threshold for mean differences below which covariates will be displayed. P-values are adjusted for multiple comparisons by the Hommel approach. For example, setting `p.value = 0.05` will display all covariates that have a significant difference between subgroups with p-value less than 0.05. Defaults to 0.001.

See Also

- `validate.subgroup` for function which creates validation results and `fit.subgroup` for function which fits subgroup identification models.
- `summarize.subgroups` for function which summarizes subgroup covariate values.
subgroup.effects  Computes treatment effects within various subgroups

Description

Computes treatment effects within various subgroups to estimate subgroup treatment effects

Usage

subgroup.effects(benefit.scores, y, trt, pi.x, cutpoint = 0,
larger.outcome.better = TRUE, reference.trt = NULL)

Arguments

- `benefit.scores`: vector of estimated benefit scores
- `y`: The response vector
- `trt`: treatment vector with each element equal to a 0 or a 1, with 1 indicating treatment status is active.
- `pi.x`: The propensity score for each observation
- `cutpoint`: numeric value for patients with benefit scores above which (or below which if `larger.outcome.better = FALSE`) will be recommended to be in the treatment group. Can also set `cutpoint = "median"`, which will use the median value of the benefit scores as the cutpoint or can set specific quantile values via "quantx" where "x" is a number between 0 and 100 representing the quantile value; e.g. `cutpoint = "quant75"` will use the 75th percent upper quantile of the benefit scores as the quantile.
- `larger.outcome.better`: boolean value of whether a larger outcome is better. Set to TRUE if a larger outcome is better and set to FALSE if a smaller outcome is better. Defaults to TRUE.
- `reference.trt`: index of which treatment is the reference (in the case of multiple treatments). This should be known already, as for a `trt` with K-levels, there will be K-1 benefit scores (1 per column) of `benefit.scores`, where each column is a comparison of each K-1 treatments with the reference treatment. The default is the last level of `trt` if it is a factor.

See Also

`fit.subgroup` for function which fits subgroup identification models which generate benefit scores.
summarize.subgroups  Summarizing covariates within estimated subgroups

Description

Summarizes covariate values within the estimated subgroups

Usage

summarize.subgroups(x, ...)

## Default S3 method:
summarize.subgroups(x, subgroup, ...)

## S3 method for class 'subgroup_fitted'
summarize.subgroups(x, ...)

Arguments

x  a fitted object from fit.subgroup() or a matrix of covariate values
...
optional arguments to summarize.subgroups methods
subgroup  vector of indicators of same length as the number of rows in x if x is a matrix. A value of 1 in the ith position of subgroup indicates patient i is in the subgroup of patients recommended the treatment and a value of 0 in the ith position of subgroup indicates patient i is in the subgroup of patients recommended the control. If x is a fitted object returned by fit.subgroup(), subgroup is not needed.

Details

The p-values shown are raw p-values and are not adjusted for multiple comparisons.

See Also

fit.subgroup for function which fits subgroup identification models and print.subgroup_summary for arguments for printing options for summarize.subgroups().

Examples

library(personalized)

set.seed(123)
n.obs <- 1000
n.vars <- 50
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,21] - 0.5 * x[,41]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)

trt <- 2 * trt01 - 1

# simulate response
delta <- 2 * (0.5 + x[,2] - x[,3] - x[,11] + x[,1] * x[,12])
xbeta <- x[,1] + x[,11] - 2 * x[,12]^2 + x[,13]
xbeta <- xbeta + delta * trt

# continuous outcomes
y <- drop(xbeta) + rnorm(n.obs, sd = 2)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
  # fit propensity score model
  propens.model <- cv.glmnet(y = trt,
                             x = x, family = "binomial")
  pi.x <- predict(propens.model, s = "lambda.min",
                  newx = x, type = "response")[,1]

  pi.x
}

subgrp.model <- fit.subgroup(x = x, y = y,
                             trt = trt01,
                             propensity.func = prop.func,
                             loss = "sq_loss_lasso",
                             nfolds = 5)  # option for cv.glmnet

comp <- summarize.subgroups(subgrp.model)
print(comp, p.value = 0.01)

# or we can simply supply the matrix x and the subgroups
comp2 <- summarize.subgroups(x, subgroup = 1 * (subgrp.model$benefit.scores > 0))

print(comp2, p.value = 0.01)

summary.subgroup_fitted

Summary of results for fitted subgroup identification models

Description

Prints summary of results for estimated subgroup treatment effects
Prints summary of results for estimated weighted ksvm
Usage

```r
## S3 method for class 'subgroup_fitted'
summary(object,
digits = max(getOption("digits") - 3, 3), ...)

## S3 method for class 'wksvm'
summary(object, digits = max(getOption("digits") - 3, 3), ...)
```

Arguments

- **object**: a fitted object from either `fit.subgroup` or `validate.subgroup`
- **digits**: minimal number of significant digits to print.
- **...**: further arguments passed to or from `print.default`.

See Also

- `validate.subgroup` for function which creates validation results and `fit.subgroup` for function which fits subgroup identification models.

---

`treatment.effects`  
**Calculation of covariate-conditional treatment effects**

**Description**

Calculates covariate conditional treatment effects using estimated benefit scores

**Usage**

```r
treatment.effects(x, ...)

## Default S3 method:
treatment.effects(x, ...)
```

```r
method = c("weighting", "a_learning"), pi.x = NULL, ...)

## S3 method for class 'subgroup_fitted'
treatment.effects(x, ...)
```
Arguments

- **x**: a fitted object from `fit.subgroup()`
- **benefit.scores**: vector of estimated benefit scores
- **loss**: loss choice USED TO CALCULATE benefit.scores of both the M function from Chen, et al (2017) and potentially the penalty used for variable selection. See `fit.subgroup` for more details.
- **method**: method choice USED TO CALCULATE benefit.scores. Either the "weighting" method or "aLearning" method. See `fit.subgroup` for more details.
- **pi.x**: The propensity score for each observation

Value

A List with elements delta (if the treatment effects are a difference/contrast, i.e. $E[Y|T = 1, X] - E[Y|T = 1, X]$) and gamma (if the treatment effects are a ratio, i.e. $E[Y|T = 1, X]/E[Y|T = 1, X]$)

See Also

- `fit.subgroup` for function which fits subgroup identification models.
- `print.individual_treatment_effects` for printing of objects returned by `treat.effects` or `treatment.effects`

Examples

```r
library(personalized)

set.seed(123)
n.obs <- 1000
n.vars <- 50
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,2] - 0.5 * x[,41]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)
trt <- 2 * trt01 - 1

# simulate response
delta <- 2 * (0.5 + x[,2] - x[,3] - x[,11] + x[,1] * x[,12])
xbeta <- x[,1] + x[,11] - 2 * x[,12]+2 + x[,13]
xbeta <- xbeta + delta * trt

# continuous outcomes
y <- drop(xbeta) + rnorm(n.obs, sd = 2)

# time-to-event outcomes
```

validate.subgroup

Validating fitted subgroup identification models

Description

Usage

validate.subgroup(model, B = 50L, method = c("training_test_replication", "boot_bias_correction"), train.fraction = 0.5, benefit.score.quantiles = c(0.1666667, 0.3333333, 0.5, 0.6666667, 0.8333333), parallel = FALSE)

Arguments

model fitted model object returned by fit.subgroup() function

B integer. number of bootstrap replications or refitting replications.

method validation method. "boot_bias_correction" for the bootstrap bias correction method of Harrell, et al (1996) or "training_test_replication" for repeated training and test splitting of the data (train.fraction should be specified for this option)

train.fraction fraction (between 0 and 1) of samples to be used for training in training/test replication. Only used for method = "training_test_replication"

benefit.score.quantiles a vector of quantiles (between 0 and 1) of the benefit score values for which to return bootstrapped information about the subgroups. I.e if one of the quantile values is 0.5, the median value of the benefit scores will be used as a cutoff to determine subgroups and summary statistics will be returned about these subgroups

parallel Should the loop over replications be parallelized? If FALSE, then no, if TRUE, then yes. If user sets parallel = TRUE and the fitted fit.subgroup() object uses the parallel version of an internal model, say for cv.glmnet(), then the internal parallelization will be overridden so as not to create a conflict of parallelism.

Details

Estimates of various quantities conditional on subgroups and treatment statuses are provided and displayed via the print.subgroup_validated function:

1. "Conditional expected outcomes" The first results shown when printing a subgroup_validated object are estimates of the expected outcomes conditional on the estimated subgroups (i.e. which subgroup is 'recommended' by the model) and conditional on treatment/intervention status. If there are two total treatment options, this results in a 2x2 table of expected conditional outcomes.

2. "Treatment effects conditional on subgroups" The second results shown when printing a subgroup_validated object are estimates of the expected outcomes conditional on the estimated subgroups. If the treatment takes levels $j \in \{1, \ldots, K\}$, a total of $K$ conditional treatment effects will be shown. For example, of the outcome is continuous, the $j$th conditional treatment effect is defined as $E(Y|Trt = j, Subgroup = j) - E(Y|Trt = j, Subgroup = / = j)$, where $Subgroup = j$ if treatment $j$ is recommended, i.e. treatment $j$ results in the largest/best expected potential outcomes given the fitted model.
3. "Overall treatment effect conditional on subgroups" The third quantity displayed shows the overall improvement in outcomes resulting from all treatment recommendations. This is essentially an average over all of the conditional treatment effects weighted by the proportion of the population recommended each respective treatment level.

**Value**

An object of class "subgroup_validated"

- avg.results: Estimates of average conditional treatment effects when subgroups are determined based on the provided cutoff value for the benefit score. For example, if cutoff = 0 and there is a treatment and control only, then the treatment is recommended if the benefit score is greater than 0.
- se.results: Standard errors of the estimates from avg.results.
- boot.results: Contains the individual results for each replication. avg.results is comprised of averages of the values from boot.results.
- avg.quantile.results: Estimates of average conditional treatment effects when subgroups are determined based on different quantile cutoff values for the benefit score. For example, if benefit.score.quantiles = 0.75 and there is a treatment and control only, then the treatment is recommended if the benefit score is greater than the 75th upper quantile of all benefit scores. If multiple quantile values are provided, e.g. benefit.score.quantiles = c(0.15, 0.5, 0.85), then results will be provided for all quantile levels.
- se.quantile.results: Standard errors corresponding to avg.quantile.results.
- boot.results.quantiles: Contains the individual results for each replication. avg.quantile.results is comprised of averages of the values from boot.results.quantiles.
- family: Family of the outcome. For example, "gaussian" for continuous outcomes.
- method: Method used for subgroup identification model. Weighting or A-learning.
- n.trts: The number of treatment levels.
- comparison.trts: All treatment levels other than the reference level.
- reference.trt: The reference level for the treatment. This should usually be the control group/level.
- larger.outcome.better: If larger outcomes are preferred for this model.
- cutpoint: Benefit score cutoff value used for determining subgroups.
- val.method: Method used for validation.
- iterations: Number of replications used in the validation process.

**References**

validate.subgroup


See Also

fit.subgroup for function which fits subgroup identification models, plot.subgroup_validated for plotting of validation results, and print.subgroup_validated for arguments for printing options for validate.subgroup().

Examples

library(personalized)

set.seed(123)
n.obs <- 500
n.vars <- 20
x <- matrix(rnorm(n.obs * n.vars, sd = 3), n.obs, n.vars)

# simulate non-randomized treatment
xbetat <- 0.5 + 0.5 * x[,11] - 0.5 * x[,13]
trt.prob <- exp(xbetat) / (1 + exp(xbetat))
trt01 <- rbinom(n.obs, 1, prob = trt.prob)
trt <- 2 * trt01 - 1

# simulate response
delta <- 2 * (0.5 + x[,2] - x[,3] - x[,11] + x[,1] * x[,12])
xbeta <- xbeta + delta * trt

# continuous outcomes
y <- drop(xbeta) + rnorm(n.obs, sd = 2)

# create function for fitting propensity score model
prop.func <- function(x, trt)
{
  # fit propensity score model
  propens.model <- cv.glmnet(y = trt,
      x = x, family = "binomial")
  pi.x <- predict(propens.model, s = "lambda.min",
      newx = x, type = "response")[,1]
  pi.x
}

subgrp.model <- fit.subgroup(x = x, y = y,
    trt = trt01,
    propensity.func = prop.func,
    loss = "sq_loss_lasso",
    nfolds = 5) # option for cv.glmnet
weighted.ksvm

### weighted.ksvm

*Fit weighted kernel svm model.*

---

**Description**

Fits weighted kernel SVM. To be used for OWL with hinge loss (but can be used more generally)

**Usage**

```r
weighted.ksvm(y, x, weights, C = c(0.1, 0.5, 1, 5, 10),
               kernel = "rbfdot", kpar = "automatic", nfolds = 10,
               foldid = NULL, ...)
```

**Arguments**

- `y` The response vector (either a character vector, factor vector, or numeric vector with values in -1, 1)
- `x` The design matrix (not including intercept term)
- `weights` vector of sample weights for weighted SVM
- `C` cost of constraints violation, see `ksvm`

---

```r
subgrp.model$subgroup.trt.effects
x.test <- matrix(rnorm(10 * n.obs * n.vars, sd = 3), 10 * n.obs, n.vars)

# simulate non-randomized treatment
xbetat.test <- 0.5 + 0.5 * x.test[,11] - 0.5 * x.test[,13]
trt01.test <- exp(xbetat.test) / (1 + exp(xbetat.test))
trt.test <- rbinom(10 * n.obs, 1, prob = trt01.test)

# simulate response
delta.test <- 2 * (0.5 + x.test[,2] - x.test[,3] - x.test[,11] + x.test[,1] * x.test[,12])
xbeta.test <- xbeta.test + delta.test * trt.test

y.test <- drop(xbeta.test) + rnorm(10 * n.obs, sd = 2)
valmod <- validate.subgroup(subgrp.model, B = 3,
                             method = "training_test",
                             train.fraction = 0.75)
valmod

print(valmod, which.quant = c(4, 5))
```
weighted.ksvm

kernel

kernel function used for training and prediction. See ksvm and kernels

kpar

list of hyperparameters for the kernel function. See ksvm

nfolds

number of cross validation folds for selecting value of C

foldid

optional vector of values between 1 and nfolds specifying which fold each observation is in. If specified, it will override the nfolds argument.

... extra arguments to be passed to ipop from the kernlab package

See Also

predict.wksvm for predicting from fitted weighted.ksvm objects

Examples

library(kernlab)

x <- matrix(rnorm(200 * 2), ncol = 2)

y <- 2 * (sin(x[,2]) ^ 2 * exp(-x[,2]) - 0.2 > rnorm(200, sd = 0.1)) - 1

weights <- runif(100, max = 1.5, min = 0.5)

wk <- weighted.ksvm(x = x[1:100,], y = y[1:100], C = c(0.1, 0.5, 1, 2, 10), weights = weights[1:100])

pr <- predict(wk, newx = x[101:200,])

mean(pr == y[101:200])
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