

Package ‘pensynth’

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Type Package

Title Penalized Synthetic Control Estimation

Version 0.8.2

Description Estimate penalized synthetic control models and perform hold-out validation to determine their penalty parameter. This method is based on the work by Abadie & L'Hour (2021) <[doi:10.1080/01621459.2021.1971535](https://doi.org/10.1080/01621459.2021.1971535)>. Penalized synthetic controls smoothly interpolate between one-to-one matching and the synthetic control method.

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URL <https://github.com/vankesteren/pensynth>

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cv_pensynth	<i>Hold-out validated penalized synthetic control estimator</i>
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Description

Compute a penalized synthetic control estimator with hold-out validation for the lambda penalty parameter. Lambda will be determined by minimizing the mean squared error on a hold-out set of pre-intervention outcome time-series.

Usage

```
cv_pensynth(
  X1,
  X0,
  Z1,
  Z0,
  v = 1,
  nlambda = 100,
  opt_pars = clarabel::clarabel_control(),
  standardize = TRUE,
  return_solver_info = FALSE,
  verbose = interactive(),
  adaptive_lambda = TRUE
)
```

Arguments

X1	N_covars by N_treated matrix of treated unit covariates
X0	N_covars by N_donors matrix of donor unit covariates
Z1	N_targets by N_treated matrix of treated unit hold-out outcome(s)
Z0	N_targets by N_donors matrix of donor unit hold-out outcomes
v	N_covars vector of variable weights, default 1
nlambda	integer length of lambda sequence (see details)
opt_pars	clarabel settings using <code>clarabel::clarabel_control()</code>
standardize	boolean whether to standardize the input matrices (default TRUE)

```

return_solver_info
    boolean whether to return diagnostic information concerning solver (default
    FALSE)
verbose
    boolean whether to print progress messages. Default on if in an interactive
    session.
adaptive_lambda
    boolean whether to allow the selected lambda to differ across treated units (de-
    fault TRUE)

```

Details

The lambda sequence is an exponentially increasing sequence where The minimum lambda is always $1e-11$, the max lambda is determined by the data.

For multiple treated units, is `adaptive_lambda` is set to `FALSE`, the (shared) minimum lambda will be selected by local regression of $\sqrt{\text{mse}}$ on $\log(\text{lambda})$.

Value

A list of optimal weights, optimal lambda(s), the lambda sequence(s), the associated weights, and the mses. If there are multiple treated units, this list contains sublists for each unit. If `return_solver_info` is `TRUE`, the list will also contain diagnostic information about the solvers.

See Also

[pensynth\(\)](#), [plot.cvpensynth\(\)](#), [placebo_test\(\)](#), [simulate_data_synth\(\)](#)

Examples

```

set.seed(45)
dat <- simulate_data_synth()
res <- with(dat, cv_pensynth(X1, X0, Z1, Z0))
plot(res)

```

in_convex_hull

Check whether treated unit is in the convex hull of donors

Description

This function finds out if the treated unit is in the convex hull of the donor units.

Usage

```
in_convex_hull(X1, X0, verbose = interactive(), ...)
```

Arguments

X1	N_covars by N_treated matrix of treated unit covariates
X0	N_covars by N_donors matrix of donor unit covariates
verbose	boolean whether to print progress messages. Default on if in an interactive session.
...	additional arguments passed to <code>clarabel::clarabel()</code>

Details

This function does not actually construct the convex hull (which is infeasible in higher dimensions), but rather it checks whether the following linear program has a feasible solution:

$$\min q'w \text{ s.t. } Aw = b$$

with w constrained to be above 0 and sum to 1, the feasibility of this linear program directly corresponds to whether the treated is in the convex hull

When the treated unit very close to the boundary of the convex hull this method usually cannot determine this exactly and this function may return NA with the warning "Solver terminated due to lack of progress"

Value

bool whether the treated unit is in the convex hull of the donor units. NA if this cannot be determined. Vector if X1 has multiple columns.

Examples

```
# create some data
set.seed(45)
X0 <- matrix(runif(20), nrow = 2)
X1 <- matrix(c(.5, .5))

# test if X1 is in the convex hull:
in_convex_hull(X1, X0)

# also works with multiple units in X1:
X1 <- cbind(X1, c(1.3, -3))
in_convex_hull(X1, X0)
```

Description

For a given set of variable weights (v) this function estimates the unit weights for a synthetic control with penalization according to Abadie & L'Hour (2021). This function deals with only a single treated unit.

Usage

```
pensynth(  
  X1,  
  X0,  
  v = 1,  
  lambda = 0,  
  opt_pars = clarabel::clarabel_control(),  
  standardize = TRUE,  
  verbose = interactive()  
)
```

Arguments

X1	N_covars by N_treated matrix of treated unit covariates
X0	N_covars by N_donors matrix of donor unit covariates
v	N_covars vector of variable weights (default 1)
lambda	numeric penalization parameter
opt_pars	clarabel settings using clarabel::clarabel_control()
standardize	boolean whether to standardize the input matrices (default TRUE)
verbose	boolean whether to print progress messages. Default on if in an interactive session.

Details

This routine uses the same notation of the original [Synth::synth\(\)](#) implementation but uses a different, faster quadratic program solver (namely, [clarabel::clarabel\(\)](#)). Additionally, it implements the penalization procedure described in Abadie & L'Hour (2021), such that the loss function is as in equation 5 of that paper.

Variable weights are not optimized by this function, meaning they need to be pre-specified. This is by design.

The original synthetic control method can be recovered by setting $\lambda = 0$. For determining λ based on hold-out data, see [cv_pensynth\(\)](#).

Value

A list with two values: w, the estimated weights; and solution, the result of the optimization.

References

Abadie, A., & L'Hour, J. (2021). A penalized synthetic control estimator for disaggregated data. *Journal of the American Statistical Association*, 116(536), 1817-1834.

See Also

[cv_pensynth\(\)](#), [placebo_test\(\)](#), [simulate_data_synth\(\)](#), [Synth::synth\(\)](#)

Examples

```

# generate some data
X0 <- matrix(
  c(1, 1.3,
    0.5, 1.8,
    1.1, 2.4,
    1.8, 1.8,
    1.3, 1.8), 2)
X1 <- matrix(c(0.8, 1.65), 2)

# run classic synthetic control (no penalization)
res <- pensynth(X1, X0)

# plot donor units in covariate space
plot(t(X0), asp = 1, xlab = "X1", ylab = "X2",
     main = "Covariate space plot")
# add the treated unit
points(t(X1), pch = 2)
# add the synthetic control
points(t(X0*%res$w), pch = 3)

# run synthetic control with penalty
res <- pensynth(X1, X0, lambda = 0.5)
# the resulting synthetic control is
# biased towards its closest neighbours
points(t(X0 %%% res$w), pch = 4)

```

placebo_test

Placebo permutation test for pensynth

Description

Perform a permutation test on a pensynth object, in the sense of Abadie, Diamond, and Hainmueller (2010). The pensynth method is performed multiple times, treating each donor as the treated unit and the treated unit with the remaining donors as the donor units.

Usage

```

placebo_test(object, Y1, Y0, verbose = TRUE)

## S3 method for class 'pensynth'
placebo_test(object, Y1, Y0, verbose = TRUE)

## S3 method for class 'cvpensynth'
placebo_test(object, Y1, Y0, verbose = TRUE)

```

Arguments

object	a fitted pensynth or cvpensynth object
Y1	the post-intervention outcome of the treated unit
Y0	the post-intervention outcome of the donor units
verbose	boolean whether to print progress messages. Default on if in an interactive session. (with N_donors columns)

Details

Note that this function updates the original call in order to re-estimate the synthetic control on the permuted data. Ensure that the data is available to the placebo test function (i.e., avoid complex environment functions such as `with()`), and ensure that the data does not change between estimating the original object and calling this function.

Value

A list with two elements

- E1, the treated unit effect(s), computed as $Y1 - Y0 \%*\% w$
- E0, the donor unit effects, computed in the same way but using the permutation test's weights.
- ATE1, the estimated ATE of the treated unit(s)
- ATE0, the estimated ATE of the donor units

References

Abadie, A., Diamond, A., & Hainmueller, J. (2010). Synthetic control methods for comparative case studies: Estimating the effect of California's tobacco control program. *Journal of the American statistical Association*, 105(490), 493-505.

See Also

[pensynth\(\)](#), [cv_pensynth\(\)](#), [plot.pensynthtest\(\)](#), [stats::update\(\)](#)

Examples

```
set.seed(45)

# simulate data with an effect of 0.8 SD
dat <- simulate_data_synth(treatment_effect = .8)

# fit a model
fit <- pensynth(dat$X1, dat$X0, lambda = 1e-5)

# Perform placebo test
test <- placebo_test(fit, dat$Y1, dat$Y0)
plot(test)
abline(h = .8, lty = 2)
legend("bottomright", lty = 2, legend = "true effect")
```

```
# compute a pseudo p-value based on ATE in
# the post-intervention time period
ref_dist <- stats::ecdf(test$ATE0)
1 - ref_dist(test$ATE1)
```

plot.cvpensynth *Plotting for hold-out validated penalized synthetic control objects*

Description

Displays a mean squared error curve and weights curve as a function of lambda, the penalization parameter.

Usage

```
## S3 method for class 'cvpensynth'
plot(x, treated_unit = 1, ...)
```

Arguments

x	a cvpensynth output object
treated_unit	integer index of the treated unit to display
...	additional arguments passed to plot()

Value

No return value, called for side effects

See Also

[cv_pensynth\(\)](#) [pensynth\(\)](#)

plot.pensynthtest *Plotting a pensynth permutation object*

Description

Plotting the reference distribution and the estimated treatment effect for the treated unit for the pensynth permutation test.

Usage

```
## S3 method for class 'pensynthtest'
plot(x, treated_unit = 1, ...)
```

Arguments

x a pensynthtest object
treated_unit integer index of the treated unit to display
... additional parameters passed to plot

Value

No return value, called for side effects

See Also

[base::plot\(\)](#)

predict.cvpensynth *Create prediction from cvpensynth model*

Description

Matrix multiplies the values in newdata by the unit weights extracted from the cvpensynth object to produce predicted values.

Usage

```
## S3 method for class 'cvpensynth'  
predict(object, newdata, lambda, ...)
```

Arguments

object a fitted cvpensynth model
newdata $N_values * N_donors$ matrix of values for the donor units.
lambda desired lambda value (defaults to optimal lambda)
... ignored

Details

For a chosen lambda that is not in the list of tested lambdas in the cvpensynth object, the closest lambda (on the log scale) will be chosen.

Value

a matrix (column vector) of predicted values

predict.pensynth *Create prediction from pensynth model*

Description

Matrix multiplies the values in newdata by the unit weights extracted from the pensynth object to produce predicted values.

Usage

```
## S3 method for class 'pensynth'
predict(object, newdata, ...)
```

Arguments

object	a fitted cvpensynth model
newdata	N_values * N_donors matrix of values for the donor units.
...	ignored

Value

a matrix (column vector) of predicted values

print.cvpensynth *Print cvpensynth model*

Description

Print cvpensynth model

Usage

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

Arguments

x	a cvpensynth object
...	ignored

Value

the cvpensynth object, invisibly

print.pensynth	<i>Print pensynth model</i>
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Description

Print pensynth model

Usage

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

Arguments

x	a pensynth object
...	ignored

Value

the pensynth object, invisibly

simulate_data_factor	<i>Simulate data according to factor model</i>
----------------------	--

Description

This function simulates data according to a latent factor model:

1. Simulate time-varying latent factors, which are the same for all units
2. Simulate time-invariant factor loadings, separately for each donor unit
3. Create sparse unit weights for each treated unit
4. Compute the loadings for the treated units as donor-unit loadings * weights
5. Simulate observed outcome time-series as factors * loadings + error
6. Do the same for each covariate, holding loadings equal. Average across pre-intervention time-points.

Usage

```
simulate_data_factor(
  N_donor = 50,
  N_treated = 1,
  N_nonzero = 4,
  N_covar = 5,
  N_pre = 12,
  N_post = 6,
  N_factors = 3,
  treatment_effect = 1,
  sd_factors = sqrt(2)/2,
  ar1_factors = 0.8,
  sd_loadings = sqrt(2)/2,
  sd_errors = 0.5,
  covar_means = TRUE
)
```

Arguments

N_donor	number of donors
N_treated	number of treated units
N_nonzero	number of true nonzero weights
N_covar	number of covariates
N_pre	number of pre-intervention timepoints
N_post	number of post-intervention timepoints
N_factors	number of latent factors to simulate
treatment_effect	the size of the true treatment effect
sd_factors	the standard deviation of the (unit-invariant, time-varying) factors
ar1_factors	autoregressive effect of the factors
sd_loadings	the standard deviation of the (time-invariant) factor loadings
sd_errors	the standard deviation of the independent errors
covar_means	whether to average the covariates across the pre-intervention times (experimental)

Details

Note that treatment effect can be a single number, but it may also be a vector of length N_{post} , indicating the effect size at each post-intervention measurement. occasion. It may also be a matrix of size N_{post} by N_{treated} .

Standard values of `sd_factors`, `sd_loadings`, and `sd_errors` have been chosen such that the observed variables have expected variance of 1.

Value

A list with the following elements

- W the true unit weights
- X0 the donor unit covariates
- X1 the treated unit covariates
- Z0 the donor unit pre-intervention outcomes
- Z1 the treated unit pre-intervention outcomes
- Y0 the donor unit post-intervention outcomes
- Y1 the treated unit post-intervention outcomes

See Also

[pensynth\(\)](#), [cv_pensynth\(\)](#), [placebo_test\(\)](#), [simulate_data_synth\(\)](#)

Examples

```
# simulate data with an effect of 0.8 SD
dat <- simulate_data_factor(N_treated = 3)

plot(
  NA,
  ylim = c(-5, 5),
  xlim = c(1, 18),
  main = "Simulated data",
  ylab = "Outcome value",
  xlab = "Timepoint"
)
for (n in 1:ncol(dat$Z0))
  lines(1:18, c(dat$Z0[, n], dat$Y0[, n]), col = "grey")
for (n in 1:ncol(dat$Z1)) {
  lines(1:18, c(dat$Z1[, n], dat$Y1[, n]), lwd = 2, col = n)
  lines(1:18, (rbind(dat$Z0, dat$Y0) %*% dat$W)[,n], lty = 2, lwd = 2, col = n)
}

abline(v = nrow(dat$Z1) + 0.5, lty = 3)
legend(
  x = "bottomleft",
  legend = c(
    "Donor units",
    "Treated unit",
    "Synth. control"
  ),
  lty = c(1, 1, 2),
  lwd = c(1, 2, 2),
  col = c("grey", "black", "black")
)
text(nrow(dat$Z1) + 0.5, -5, "Intervention\ntimepoint", pos = 4, font = 3)
```

simulate_data_synth *Simulate data according to synthetic control model*

Description

This function simulates a basic form of synthetic control data, mainly for testing purposes.

Usage

```
simulate_data_synth(  
  N_donor = 50,  
  N_treated = 1,  
  N_covar = 5,  
  N_pre = 12,  
  N_post = 6,  
  N_nonzero = 4,  
  treatment_effect = 1,  
  sd_resid_X = 0.1,  
  sd_resid_ZY = 0.1,  
  ar1_outcome = 0.8  
)
```

```
simulate_data(  
  N_donor = 50,  
  N_treated = 1,  
  N_covar = 5,  
  N_pre = 12,  
  N_post = 6,  
  N_nonzero = 4,  
  treatment_effect = 1,  
  sd_resid_X = 0.1,  
  sd_resid_ZY = 0.1,  
  ar1_outcome = 0.8  
)
```

Arguments

N_donor	number of donors
N_treated	number of treated units
N_covar	number of covariates
N_pre	number of pre-intervention timepoints
N_post	number of post-intervention timepoints
N_nonzero	number of true nonzero weights
treatment_effect	the size of the true treatment effect

sd_resid_X	the residual standard deviation of X1
sd_resid_ZY	the residual standard deviation of Z1 and Y1
ar1_outcome	autoregressive effect of the outcome

Details

Note that treatment effect can be a single number, but it may also be a vector of length N_{post} , indicating the effect size at each post-intervention measurement. occasion. It may also be a matrix of size N_{post} by N_{treated} .

Value

A list with the following elements

- W the true unit weights
- X0 the donor unit covariates
- X1 the treated unit covariates
- Z0 the donor unit pre-intervention outcomes
- Z1 the treated unit pre-intervention outcomes
- Y0 the donor unit post-intervention outcomes
- Y1 the treated unit post-intervention outcomes

See Also

[pensynth\(\)](#), [cv_pensynth\(\)](#), [placebo_test\(\)](#), [simulate_data_factor\(\)](#)

Examples

```
# simulate data with an effect of 0.8 SD
dat <- simulate_data_synth(treatment_effect = 0.8)

plot(
  NA,
  ylim = c(-3, 3),
  xlim = c(1, 18),
  main = "Simulated data",
  ylab = "Outcome value",
  xlab = "Timepoint"
)
for (n in 1:ncol(dat$Z0))
  lines(1:18, c(dat$Z0[, n], dat$Y0[, n]), col = "grey")
lines(1:18, c(dat$Z1, dat$Y1), lwd = 2)
lines(1:18, rbind(dat$Z0, dat$Y0) %*% dat$W, lty = 2, lwd = 2)
abline(v = length(dat$Z1) + 0.5, lty = 3)
legend(
  x = "bottomleft",
  legend = c(
    "Donor units",
    "Treated unit",
  )
)
```

```
      "Synth. control"  
    ),  
    lty = c(1, 1, 2),  
    lwd = c(1, 2, 2),  
    col = c("grey", "black", "black")  
  )  
  text(length(dat$Z1) + 0.5, -3, "Intervention\ntimepoint", pos = 4, font = 3)
```

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