

# Package ‘apm’

July 22, 2025

**Title** Averaged Prediction Models

**Version** 0.1.1

**Description** In panel data settings, specifies set of candidate models, fits them to data from pre-treatment validation periods, and selects model as average over candidate models, weighting each by posterior probability of being most robust given its differential average prediction errors in pre-treatment validation periods. Subsequent estimation and inference of causal effect's bounds accounts for both model and sampling uncertainty, and calculates the robustness changepoint value at which bounds go from excluding to including 0. The package also includes a range of diagnostic plots, such as those illustrating models' differential average prediction errors and the posterior distribution of which model is most robust.

**License** GPL (>= 2)

**Depends** R (>= 3.5.0)

**Imports** stats, ggplot2 (>= 3.5.1), ggh4x (>= 0.2.8), ggrepel (>= 0.9.6), MASS, sandwich, pbapply (>= 1.7-2), fwb (>= 0.3.0), chk (>= 0.10.0)

**Suggests** parallel, knitr, rmarkdown

**Encoding** UTF-8

**RoxygenNote** 7.3.2

**LazyData** true

**VignetteBuilder** knitr

**URL** <https://github.com/tl2624/apm/>, <https://tl2624.github.io/apm/>

**BugReports** <https://github.com/tl2624/apm/issues>

**NeedsCompilation** no

**Author** Thomas Leavitt [aut, cre] (ORCID: <https://orcid.org/0000-0002-3668-6409>),  
Laura Hatfield [aut] (ORCID: <https://orcid.org/0000-0003-0366-3929>),  
Noah Greifer [aut] (ORCID: <https://orcid.org/0000-0003-3067-7154>)

**Maintainer** Thomas Leavitt <thomas.leavitt@baruch.cuny.edu>

**Repository** CRAN

**Date/Publication** 2025-05-22 20:00:02 UTC

## Contents

apm_est . . . . .	2
apm_mod . . . . .	5
apm_pre . . . . .	7
plot.apm_pre_fits . . . . .	10
ptpdata . . . . .	12
robustness_bound . . . . .	13

<b>Index</b>	<b>15</b>
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apm_est	<i>Estimate ATTs from models fits</i>
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### Description

apm\_est() computes the ATTs from the models previously fit by [apm\\_pre\(\)](#), choosing the optimal one by minimizing the largest absolute average prediction error across validation times. Optionally, this process can be simulated to arrive at a distribution of ATTs that accounts for the uncertainty in selecting the optimal model. plot() plots the resulting ATT(s).

### Usage

```
apm_est(
  fits,
  post_time,
  M = 0,
  R = 1000L,
  all_models = FALSE,
  cl = NULL,
  verbose = TRUE,
  ...
)

## S3 method for class 'apm_est'
summary(object, level = 0.95, M = NULL, ...)

## S3 method for class 'apm_est'
plot(x, label = TRUE, size.weights = TRUE, ...)
```

### Arguments

fits	an apm_pre_fits object; the output of a call to <a href="#">apm_pre()</a> .
post_time	the value of the time variable considered post-treatment, for which the ATT is to be estimated.

M	the sensitivity parameter for set identification. For <code>apm_est()</code> , the default is 0, i.e., under point identification. For <code>summary()</code> , this can be set to one or more positive values to produce uncertainty bounds for each value. Only allowed when not set to 0 in the call to <code>apm_est()</code> . See Details.
R	the number of bootstrap iterations used to compute the sampling variance of the ATT. Default is 1000. More is better but takes longer.
all_models	logical; whether to compute ATTs for all models (TRUE) or just those with BMA weights greater than 0 (FALSE, default). This will not effect the final estimates but leaving as FALSE can speed up computation when some models have BMA weights of 0.
cl	a cluster object created by <code>parallel::makeCluster()</code> , an integer to indicate number of child-processes (ignored on Windows) for parallel evaluations, or "future" to use a future backend. NULL (default) refers to sequential evaluation. See <code>fwb::fwb()</code> for details and issues related to replicability.
verbose	logical; whether to print information about the progress of the estimation, including a progress bar. Default is TRUE.
...	other arguments passed to <code>fwb::fwb()</code> .
level	the desired confidence level. Set to 0 to ignore sampling variation in computing the interval bounds. Default is .95.
x, object	an <code>apm_est</code> object; the output of a call to <code>apm_est()</code> .
label	logical; whether to label the ATT estimates. Default is TRUE.
size.weights	logical; whether to size the points based on their BMA weights. Default is TRUE.

## Details

`apm_est()` estimates the ATT from each model and combines them to form the BMA-weighted estimate of the ATT. Uncertainty for the BMA-weighted ATT is computed by combining two variance components, one that account for sampling and another that accounts for model selection. The component due to sampling is computed by bootstrapping the process of fitting the outcome model for the post-treatment outcome identified by `post_time` and computing the difference between the observed outcome mean difference and the model-predicted outcome mean difference. The fractional weighted bootstrap as implemented in `fwb::fwb()` is used to ensure no units are dropped from the analysis. In each bootstrap sample, the BMA-weighted ATT estimate is computed as the weighted average of the ATTs computed from the models using the fixed BMA weights computed by `apm_pre()`, and the variance is computed as the empirical variance over the bootstrapped estimates. The variance component due to model selection is computed as the BMA-weighted variance of the original ATTs.

When `M` is greater than 0, bounds for set identification and their uncertainty are additionally computed. This involves bootstrapping the fitting of the pre-period models along with post-treatment models on order to compute the maximum absolute difference in average prediction errors for each model across validation periods. Each bootstrap sample produces a margin of error for each model computed as  $M \times \delta_m$  where  $\delta_m$  is the maximum absolute difference in average prediction errors for model  $m$ . Upper and lower bounds for the set-identified BMA-weighted ATT are computed as  $ATT_m \pm M \times \delta_m$ . The same procedure as above is then used to compute the variance of these bounds.

`summary()` displays the BMA-weighted ATT estimate, its standard error, and Wald confidence intervals. When `M` is greater than 0, bounds for the set-identified ATT are displayed in the confidence interval bound columns. The lower bound is computed as  $LB - \sigma_{LB}Z_l$  and the upper bound as  $UB + \sigma_{UB}Z_l$ , where `LB` and `UB` are the lower and upper bounds,  $\sigma_{LB}$  and  $\sigma_{UB}$  are their variances accounting for sampling and model selection, and  $Z_l$  is the critical Z-statistic for confidence level  $l$ . To display the set-identification bounds themselves, one should set `level = 0`.

## Value

`apm_est()` returns an `apm_est` object, which contains the ATT estimates and their variance estimates. The following components are included:

**BMA\_att** the BMA-weighted ATT

**atts** a 1-column matrix containing the ATT estimates from each model (when `all_models = FALSE`, only models with positive BMA weights are included)

**BMA\_var** the total variance estimate for the BMA-weighted ATT incorporating the variance due to sampling and due to model selection

**BMA\_var\_b** the bootstrap-based component of the variance estimate for the BMA-weighted ATT due to sampling

**BMA\_var\_m** the component of the variance estimate for the BMA-weighted ATT due to model selection

**M** the value of the sensitivity parameter `M`

**post\_time** the value supplied to `post_time`

**observed\_means** a matrix of the observed outcome means at each pre-treatment validation period

**pred\_errors** an array containing the average prediction errors for each model and each pre-treatment validation period

**pred\_error\_diffs** a matrix containing the difference in average prediction errors between groups for each model and each pre-treatment validation period

**BMA\_weights** the BMA weights computed by `apm_pre()` (when `all_models = FALSE`, only positive BMA weights are included)

**boot\_out** an `fwb` object containing the bootstrap results

`plot()` returns a `ggplot` object displaying the ATT for each model plotted against the maximum absolute difference in average prediction errors for that model. The model with the lowest maximum absolute difference in average prediction errors is displayed in red.

`summary()` produces a table with the BMA-weighted ATT, its estimated standard error, and confidence interval limits. When `M` is greater than 0, additional rows for each value of `M` are included with the lower and upper bound. When `level` is greater than 0, these bounds include the uncertainty due to sampling and model selection; otherwise, they correspond to the set identification bounds for the ATT.

## See Also

[apm\\_pre\(\)](#) for computing the BMA weights; [fwb::fwb\(\)](#) for the fractional weighted bootstrap.

## Examples

```
data("ptpdata")

# Combination of 4 models: 2 time trends, 2 lags
models <- apm_mod(list(crude_rate ~ 1),
                  lag = 0:1,
                  time_trend = 0:1)

models

# Fit the models to data; unit_var must be supplied for
# fixed effects
fits <- apm_pre(models,
                data = ptpdata,
                group_var = "group",
                time_var = "year",
                val_times = 2004:2007,
                unit_var = "state",
                nsim = 100,
                verbose = FALSE)

est <- apm_est(fits,
               post_time = 2008,
               M = 1,
               R = 20,
               verbose = FALSE)

est

# ATT estimate and bounds for M = 1
summary(est)

# Bounds for other values of M
summary(est, M = c(.5, 1, 1.5, 2))

# Set-ID bounds without uncertainty
summary(est, level = 0)

plot(est)
```

---

apm\_mod

*Generate models used to fit outcomes*

---

## Description

`apm_mod()` generates a list of models characterized by a basic model formulas and other options (e.g., lags, families, etc.) that are supplied to `apm_pre()`. These values are completely crossed to create a grid of model specifications, and multiple sets of model specifications can be combined using `c()` (see Examples).

**Usage**

```

apm_mod(
  formula_list,
  family = "gaussian",
  lag = 0L,
  diff_k = 0L,
  log = FALSE,
  time_trend = 0L,
  fixef = FALSE,
  identiy_only_log = TRUE
)

```

**Arguments**

<code>formula_list</code>	a list of model formulas with the outcome on the left side and predictions (or just an intercept) on the right side.
<code>family</code>	a list of family specifications; see <code>family()</code> for allowable options. These will eventually be passed to <code>glm()</code> when fitting the models in <code>apm_pre()</code> . "negbin" can also be supplied to request a negative binomial model with a log link fit using <code>MASS::glm.nb()</code> . Default is "gaussian" to specify a linear model.
<code>lag</code>	a vector of integers indicating the desired outcome lags to be used as predictors. For example, a lag value of 3 means the outcome lagged once, twice, and three times will be included as predictors. Default is 0 for no lags.
<code>diff_k</code>	a vector of integers indicating the desired outcome lag to be used as an offset. For example, a <code>diff_k</code> value of 1 means the prior time point's outcome will be included as an offset, equivalent to using the outcome minus its corresponding lag as the outcome of the corresponding model. Default is 0 for no lags. Any models with a <code>diff_k</code> value less than a lag value will be removed automatically. When used with a family with a log link, the lags are automatically log-transformed; an error will be thrown by <code>apm_pre()</code> if nonpositive values are present in the outcome.
<code>log</code>	a logical vector indicating whether the outcome should be log-transformed. Default is FALSE to use the original outcome. When lag or <code>diff_k</code> are greater than 0, the outcome lags will also be log-transformed if TRUE. When the family has a log link and <code>diff_k</code> is greater than zero, the lag in the offset will be log transformed.
<code>time_trend</code>	a vector of integers indicating the desired powers to be included in a time trend. For example, a <code>time_trend</code> value of 2 means the time variable and its square will be included as predictors in the model. A value of 0 (the default) means time is not included as a predictor.
<code>fixef</code>	a logical vector indicating whether unit fixed effects should be included as predictors. Default is FALSE to omit unit fixed effects.
<code>identiy_only_log</code>	logical; whether to omit any models in which log is TRUE but the link in the family specification corresponds to something other than "identity". Default is TRUE, and this should probably not be changed.

**Value**

An `apm_models` object, which is a list containing the full cross (less any omitted combinations) of the model features specified in the arguments, with each combination a list. These have a `print()` method and can be combined using `c()`. Each model is named automatically, but these can be set manually using `names()` as well. Models can be removed by setting their value to `NULL`; see Examples.

**See Also**

`formula()`, `family()`

**Examples**

```
data("ptpdata")

# Combination of 8 models: 1 baseline formulas,
# 2 families, 2 lags, 2 time trends
models1 <- apm_mod(crude_rate ~ 1,
                  family = list("gaussian", "quasipoisson"),
                  time_trend = 0:1,
                  lag = 0:1, fixef = TRUE)

models1

# Add a single other model with a square time trend
models2 <- apm_mod(crude_rate ~ 1,
                  family = "gaussian",
                  time_trend = 2,
                  fixef = FALSE)

models2

(models <- c(models1, models2))

# Remove a model
models[[4]] <- NULL
models
```

---

apm\_pre

*Fit validation models to pre-treatment data*

---

**Description**

`apm_pre()` fits models to the pre-treatment data to compute the observed prediction errors for each model in each period and compute the Bayesian model averaging (BMA) weights eventually used in `apm_est()` to estimate the treatment effect.

**Usage**

```

apm_pre(
  models,
  data,
  weights = NULL,
  group_var,
  time_var,
  val_times,
  unit_var,
  nsim = 1000L,
  cl = NULL,
  verbose = TRUE
)

## S3 method for class 'apm_pre_fits'
summary(object, order = NULL, ...)

```

**Arguments**

<code>models</code>	an <code>apm_models</code> object; the output of a call to <code>apm_mod()</code> .
<code>data</code>	a dataset containing all the variables named in the supplied models (i.e., the outcome and any predictors) as well as any variable named below.
<code>weights</code>	an optional vector of weights (e.g., sampling weights) used to fit weighted regression models.
<code>group_var</code>	string; the name of the treatment variable in <code>data</code> defining the "to be treated" and "not to be treated" groups. The corresponding variable should take on values of 0 and 1 only.
<code>time_var</code>	string; the name of the variable in <code>data</code> containing the time variable.
<code>val_times</code>	a numeric vector corresponding to the pre-treatment times that will be used as validation times when select the model with the optimal average expected prediction error.
<code>unit_var</code>	string; the name of the unit ID variable in <code>data</code> .
<code>nsim</code>	the number of simulated draws from the joint posterior of the fitted models to use to compute the BMA weights. Default is 1000. More is better but takes longer.
<code>cl</code>	a cluster object created by <code>parallel::makeCluster()</code> , or an integer to indicate number of child-processes (integer values are ignored on Windows) for parallel evaluations. It can also be "future" to use a future backend. NULL (default) refers to sequential evaluation. See the <code>cl</code> argument of <code>pbapply::pblapply()</code> for details.
<code>verbose</code>	logical; whether to print information about the progress of the estimation, including a progress bar. Default is TRUE.
<code>object</code>	an <code>apm_pre_fit</code> object; the output of a call to <code>apm_pre()</code> .
<code>order</code>	how to order the summary; NULL (the default) is the same ordering as the models supplied to <code>apm_pre()</code> , "weights" orders the models by their computed BMA



weights with the highest weights on top, and "errors" orders the models by their maximum absolute difference in prediction errors with the smallest errors on top.

... ignored.

## Details

`apm_pre()` creates a grid of all models and all time points and fits all corresponding models. For each validation time supplied to `val_times`, each model is fit using all previous times. For example, for a validation time of 5, a model is fit with data only from periods 1-4.

`lm()`, `glm()`, or `MASS::glm.nb()` are used to fit the given models. The joint covariance matrix of all the coefficients is computed using the SUEST method described in Mize et al. (2019, p164), which is also used by the STATA command `suest`. This is equivalent to the covariance matrix computed by stacking the score equations for the models and fitting them using M-estimation and yields the equivalent of the HC0 covariance matrix for all within-model covariances. The covariance is clustered by `unit_id`.

To compute the BMA weights, random variate drawn from a multivariate normal distribution `nsim` times with mean vector equal to the concatenated model coefficients and covariance equal to the joint covariance matrix described above. For each iteration, the absolute average prediction errors are calculated for each model and validation period. A model is considered the "winner" if its largest absolute average prediction error across validation periods is the smallest among all models. The BMA weight for each model is equal to the proportion of iterations in which that model was the "winner".

## Value

`apm_pre()` returns an `apm_pre_fits` object, which is a list containing the models supplied to `models`, a grid of all fitted models, a list of all model fit objects, a list of all estimated coefficients, the joint covariance of the coefficients, the dataset supplied to `data`, and other components supplied to `apm_pre()`.

`summary()` produces a data frame containing the BMA weights and maximum absolute difference in mean prediction errors for each model, ordered according order. An asterisk appears next to the model with the smallest error.

## See Also

`lm()`, `glm()`, and `MASS::glm.nb()` for the functions used to fit the models; `apm_est()` to compute the ATT and its uncertainty; `plot.apm_pre_fits()` for plotting an `apm_pre_fits` object.

## Examples

```
data("ptpdata")

# Combination of 8 models: 2 baseline formulas,
# 2 families, 2 lags
models <- apm_mod(crude_rate ~ 1,
                  family = list("gaussian", "quasipoisson"),
                  time_trend = 0:1,
                  lag = 0:1)
```

```

models

# Fit the models to data
fits <- apm_pre(models, data = ptpdata,
               group_var = "group",
               time_var = "year",
               val_times = 1999:2007,
               unit_var = "state",
               nsim = 200,
               verbose = FALSE)

fits

summary(fits)

plot(fits, type = "weights")

plot(fits, type = "error")

```

---

plot.apm\_pre\_fits      *Plot outputs of apm\_pre()*

---

### Description

plot() displays the Bayesian model averaging (BMA) weights for each model (computed by apm\_fit() as the posterior probability of selection) and the distribution of the difference in average prediction errors.

### Usage

```

## S3 method for class 'apm_pre_fits'
plot(
  x,
  type = "weights",
  abs = TRUE,
  ncol = 4L,
  clip_at = 15,
  model = ".optimal",
  ...
)

```

### Arguments

x                    an apm\_pre\_fits object; the output of a call to [apm\\_pre\(\)](#).

type                which values to plot: allowable options include "weights" to plot the BMA weights/posterior probabilities (default), "errors" to plot the difference in average predictions errors for all models across validation periods, "predict" to plot the time series and model predictions for each model, and "corrected" to

	plot the corrected predictions for the treated group for each model. Abbreviations allowed.
abs	logical; when type = "errors", whether to plot the differences in average prediction errors in absolute value (TRUE, default) or not (FALSE).
ncol	when type is "errors", "predict", or "corrected", the number of columns to use to display the plots. Default is 4.
clip_at	when type = "errors", the value (in robust z-score units) at which to clip the y-axis of the plot to prevent outliers from distorting it. Default is 15. Set to Inf to prevent clipping.
model	string; when type = "predict" or type = "corrected", the model(s) to plot. Allowable values include ".optimal" to plot the model with the smallest maximum absolute difference in average prediction errors, ".all" to plot all models (excluding the BMA-weighted predictions), or the names of one or more specific models. Abbreviations allowed.
...	ignored.

### Details

When type = "weights", plot() displays a bar plot with a bar for each model with height equal to the BMA weight/posterior probability of selection for that model. (Note that the plot margins can sometimes cut off the model names; use theme(plot.margins =) after loading ggplot2 to extend the left margin of the plot to ensure all text is visible. Alternatively, the axis text can be rotated using theme(axis.text.x =).)

When type = "errors", plot() displays a lattice of bar plots with a plot for each model displaying the difference in average prediction errors for each validation period. The period with the largest difference in average prediction errors will be shaded black. The model with the smallest maximum absolute difference in average prediction errors will have a gray label.

When type = "predict", plot() displays a lattice of line plots with a plot for each model displaying the observed and predicted outcomes for each validation period under each model. The observed outcomes are displayed as points, while the predicted outcomes are displayed as lines.

When type = "corrected", plot() displays a lattice of line plots with a plot for each model displaying the observed and corrected predictions for the treated group for each validation period under each model. The observed outcomes are displayed as points, while the corrected predictions are displayed as lines. Corrected predictions are computed as the observed outcome in the treated group minus the prediction error in the treated group plus the prediction error in the control group.

### Value

A ggplot object, which can be manipulated using ggplot2 syntax (after loading ggplot2).

### See Also

[apm\\_pre\(\)](#) to compute the difference in average prediction errors and BMA weights; [ggplot2::geom\\_col\(\)](#), which is used to create the plots.

**Examples**

```

data("ptpdata")

# Combination of 8 models: 2 baseline formulas,
# 2 families, 2 lags
models <- apm_mod(crude_rate ~ 1,
                  family = "gaussian",
                  time_trend = 0:1,
                  lag = 0:1,
                  diff_k = 0:1)

models

# Fit the models to data
fits <- apm_pre(models, data = ptpdata,
                group_var = "group",
                time_var = "year",
                val_times = 1999:2007,
                unit_var = "state",
                nsim = 50,
                verbose = FALSE)

fits

plot(fits, type = "weights")

plot(fits, type = "error", ncol = 2)

plot(fits, type = "predict", model = ".optimal")

plot(fits, type = "corrected", model = ".optimal")

```

---

ptpdata

*Dataset on Annual Homicide Rates*


---

**Description**

A dataset of homicide rates across 9 states from 1994 to 2008. Missouri repealed its permit-to-purchase (PTP) law in 2007.

**Usage**

```
ptpdata
```

**Format**

A dataframe of 7 variables with 135 observations (9 states, 15 years).

**Details**

details for dataset2

**state** the name of the state; states present include Arkansas, Illinois, Iowa, Kansas, Kentucky, Missouri, Nebraska, Oklahoma, and Tennessee.

**year** the year of each observation. Years range from 1994 to 2008.

**deaths** the number of homicide deaths in the corresponding state in the corresponding year.

**crude\_rate** the homicide rate in the corresponding state in the corresponding year.

**age\_adj\_rate** the age-adjusted homicide rate in the corresponding state in the corresponding year.

**group** whether each observation belongs to the "to-be-treated" group; 1 for Missouri and 0 for all other states.

**treat** whether each observation is treated; 1 for Missouri in 2008 and 0 otherwise.

---

robustness_bound	<i>Compute the robustness changepoint</i>
------------------	---

---

**Description**

robustness\_bound() computes the value of the sensitivity parameter M at which the robustness bounds change from excluding to including an ATT of 0.

**Usage**

```
robustness_bound(object, level = 0.95)
```

**Arguments**

object	an apm_est object; the output of a call to <a href="#">apm_est()</a> . M must have been set to a nonzero value to use robustness_bound().
level	the desired confidence level. Set to 0 to ignore sampling variation in computing the interval bounds. Default is .95.

**Value**

A single number corresponding to the changepoint value of M. If there is no positive value of M for which the interval bounds cross 0, NA will be returned.

**See Also**

[summary.apm\\_est\(\)](#) for examining the ATT and bounds for a given value of M; [uniroot\(\)](#) for the function that finds the changepoint value of M.

**Examples**

```
data("ptpdata")

# Combination of 4 models: 2 time trends, 2 lags
models <- apm_mod(list(crude_rate ~ 1),
                  lag = 0:1,
                  time_trend = 0:1)

models

# Fit the models to data; unit_var must be supplied for
# fixed effects
fits <- apm_pre(models,
                data = ptpdata,
                group_var = "group",
                time_var = "year",
                val_times = 2004:2007,
                unit_var = "state",
                nsim = 100,
                verbose = FALSE)

est <- apm_est(fits,
               post_time = 2008,
               M = 1,
               R = 20,
               verbose = FALSE)

est

# ATT estimate and bounds for M = 1
summary(est)

#Changepoint value of M ignoring estimation uncertainty
(M <- robustness_bound(est, level = 0))

summary(est, level = 0, M = M)

#Changepoint value of M accounting for estimation uncertainty
(M <- robustness_bound(est, level = .95))

summary(est, level = .95, M = M)
```

# Index

## \* datasets

ptpdata, [12](#)

apm\_est, [2](#)  
apm\_est(), [7](#), [9](#), [13](#)  
apm\_mod, [5](#)  
apm\_mod(), [8](#)  
apm\_pre, [7](#)  
apm\_pre(), [2-6](#), [10](#), [11](#)

family(), [6](#), [7](#)  
formula(), [7](#)  
fwb::fwb(), [3](#), [4](#)

ggplot2::geom\_col(), [11](#)  
glm(), [6](#), [9](#)

lm(), [9](#)

MASS::glm.nb(), [6](#), [9](#)

names(), [7](#)

parallel::makeCluster(), [3](#), [8](#)  
pbapply::pblapply(), [8](#)  
plot.apm\_est (apm\_est), [2](#)  
plot.apm\_pre\_fits, [10](#)  
plot.apm\_pre\_fits(), [9](#)  
ptpdata, [12](#)

robustness\_bound, [13](#)

summary.apm\_est (apm\_est), [2](#)  
summary.apm\_est(), [13](#)  
summary.apm\_pre\_fits (apm\_pre), [7](#)

uniroot(), [13](#)