### tebalance density — Covariate balance density

Description	Quick start	Menu	Syntax
Options	Remarks and examples	Stored results	Reference
Also see			

### Description

tebalance density produces kernel density plots that are used to check for covariate balance after estimation by a teffects inverse-probability-weighted estimator, a teffects matching estimator, or an stteffects inverse-probability-weighted estimator.

# **Quick start**

Kernel density plot of the propensity score after teffects psmatch tebalance density

Kernel density plot of x1 after a teffects command or an stteffects command tebalance density x1

Same as above, but rescale the kernel bandwidth by a factor of 2 tebalance density x1, bwidth(\*2)

### Menu

 $Statistics > Causal \ inference/treatment \ effects > Balance > Graphs$ 

## Syntax

Density plots for the propensity score

tebalance density [, options]

Density plots for a covariate

tebalance density *varname* [, *options*]

options	Description
Main	
kernel( <i>kernel</i> )	specify the kernel function; default is kernel(epanechnikov)
<u>bw</u> idth(*#)	rescale default bandwidth
line#opts( <i>line_options</i> )	twoway line options for density line number #
twoway_options	any options other than by() documented in [G-3] twoway_options
byopts(byopts)	how subgraphs are combined, labeled, etc.

collect is allowed; see [U] 11.1.10 Prefix commands.

kernel	Description	
<u>tri</u> angle	triangle kernel function; the default	
epanechnikov	Epanechnikov kernel function	
epan2	alternative Epanechnikov kernel function	
<u>bi</u> weight	biweight kernel function	
<u>cos</u> ine	cosine trace kernel function	
gaussian	Gaussian kernel function	
parzen	Parzen kernel function	
<u>rec</u> tangle	rectangle kernel function	

# Options

Main

kernel(kernel) specifies the kernel function for use in calculating the kernel density estimates. The default kernel is the kernel(epanechnikov).

- bwidth(\*#) specifies the factor by which the default bandwidths are to be rescaled. A bandwidth is the half-width of the kernel, the width of the density window around each point. Each kernel density plot has its own bandwidth, and by default, each kernel density plot uses its own optimal bandwidth; see [R] kdensity. bwidth() rescales each plot's optimal bandwidth by the specified amount.
- line#opts(line\_options) specifies the line pattern, width, color, and overall style of density line number
  #. The line numbers are in the same order as the treatment levels specified in e(tlevels).
- twoway\_options are any of the options documented in [G-3] twoway\_options, excluding by(). These
  include options for titling the graph (see [G-3] title\_options) and for saving the graph to disk (see
  [G-3] saving\_option). tebalance density uses by() to differentiate between raw and weighted
  or matched samples, and some twoway\_options will be repeated for by graph and might be better
  specified as byopts().

byopts(by\_option) is as documented in [G-3] by\_option. byopts() affects how the subgraphs are combined, labeled, etc. byopts() generally affects the entire graph, and some by\_option may be better specified as twoway\_options; see [G-3] twoway\_options.

#### **Remarks and examples**

When the distribution of a covariate does not vary over the treatment levels, the covariate is said to be balanced. tebalance density produces kernel density plots of a covariate over treatment levels for the raw data and the weighted or matched sample produced by teffects or stteffects. If the weightedsample or matched-sample kernel density plots of the covariate are the same over the treatment levels, the covariate is balanced in the weighted or matched sample.

After telasso and all teffects and stteffects commands except teffects ra, stteffects ra, and stteffects wra,

```
. tebalance density varname [, options]
```

produces kernel density plots to check whether varname is balanced.

After teffects psmatch,

. tebalance density |, options |

produces kernel density plots to check whether the propensity score estimated by teffects psmatch is balanced. Our discussion of the use of tebalance density and interpretation of its results for a covariate below also apply to a propensity score.

We recommend that you read [CAUSAL] **tebalance** before proceeding; it provides an introduction to covariate balance and an overview of the implemented methods. See [CAUSAL] **stteffects intro** for a discussion of survival-time features.

#### Example 1: Checking covariate balance after psmatch

Using an extract from the data used by Cattaneo (2010), we use teffects psmatch to estimate the effect of a mother's smoking behavior (mbsmoke) on the birthweight of her child (bweight), controlling for marital status (mmarried), the mother's age (mage), whether the mother had a prenatal doctor's visit in the baby's first trimester (prenatal1), and whether this baby is the mother's first child (fbaby).

```
. use https://www.stata-press.com/data/r19/cattaneo2
(Excerpt from Cattaneo (2010) Journal of Econometrics 155: 138-154)
. teffects psmatch (bweight) (mbsmoke mmarried mage prenatal1 fbaby),
> generate(matchv)
Treatment-effects estimation
                                                 Number of obs
                                                                            4.642
                                                                    =
Estimator
               : propensity-score matching
                                                 Matches: requested =
                                                                                1
                                                                                1
Outcome model
              : matching
                                                                min =
Treatment model: logit
                                                                max =
                                                                              139
                             AI robust
     bweight
               Coefficient std. err.
                                                  P>|z|
                                                             [95% conf. interval]
                                            z
ATE
     mbsmoke
    (Smoker
         vs
                             27.74409
                -235.1714
                                         -8.48
                                                  0 000
                                                           -289 5488
                                                                         -180 794
 Nonsmoker)
```

We specified the option generate (matchv) to speed up the postestimation command that produces density plots, as discussed in example 3 under *Remarks and examples* in [CAUSAL] **tebalance** entry. We do not interpret the estimated effect produced by this preliminary model but rather check the specification. We begin by looking at the default density plots.

. tebalance density mage



The density plots for the matched sample are nearly indistinguishable, implying that matching on the estimated propensity score balanced the covariates. The density plots are too jagged for presentation, so we oversmooth them by scaling up the bandwidth used for each plot.

. tebalance density mage, bwidth(\*1.5)



Option bwidth() rescales the default optimal bandwidths by the specified scale factor. Each of the four density plots has its own sample size and optimal bandwidth. Rescaling each of the four bandwidths by 1.5 produces smoother plots.

### Stored results

After teffects or stteffects fits a binary treatment, tebalance density stores the following in r():

Scalars

	r(bwc_adj)	bandwidth for control in weighted or matched-adjusted sample
	r(Nc_adj)	observations on control in weighted or matched-adjusted sample
	r(bwt_adj)	bandwidth for treated in weighted or matched-adjusted sample
	r(Nt_adj)	observations on treated in weighted or matched-adjusted sample
	r(bwc_raw)	bandwidth for control in raw sample
	r(Nc_raw)	observations on control in raw sample
	r(bwt_raw)	bandwidth for treated in raw sample
	r(Nt_raw)	observations on treated in raw sample
Mac	ros	

r(kernel) name of kernel

After teffects or stteffects fits a multivalued treatment, tebalance density stores the following in r():

Scalars	
r(bw#_adj) r(N#_adj)	bandwidth for treatment level # in weighted or matched-adjusted sample observations on treatment level # in weighted or matched-adjusted sample
r(bw#_raw) r(N#_raw)	bandwidth for treatment level # in raw sample observations on treatment level # in raw sample
Macros	
r(kernel)	name of kernel

### Reference

Cattaneo, M. D. 2010. Efficient semiparametric estimation of multi-valued treatment effects under ignorability. *Journal of Econometrics* 155: 138–154. https://doi.org/10.1016/j.jeconom.2009.09.023.

# Also see

[CAUSAL] stteffects intro — Introduction to treatment effects for observational survival-time data

- [CAUSAL] tebalance Check balance after teffects or stteffects estimation
- [CAUSAL] teffects aipw Augmented inverse-probability weighting

[CAUSAL] teffects ipw — Inverse-probability weighting

- [CAUSAL] teffects ipwra Inverse-probability-weighted regression adjustment
- [CAUSAL] teffects nnmatch Nearest-neighbor matching
- [CAUSAL] teffects psmatch Propensity-score matching
- [CAUSAL] teoverlap Overlap plots

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