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tebalance — Check balance after teffects or stteffects estimation

Description Syntax Remarks and examples Methods and formulas References Also see

Description

The tebalance postestimation commands produce diagnostic statistics, test statistics, and diagnostic plots to assess whether a teffects or an stteffects command balanced the covariates over treatment levels

Syntax

tebalance subcommand ... [, options]

subcommand	Description
summarize overid	compare means and variances in raw and balanced data overidentification test
density box	kernel density plots for raw and balanced data box plots for each treatment level for balanced data
DOX	box plots for each treatment level for balanced data

Remarks and examples

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This entry provides an overview of the commands in tebalance. We recommend that you read this entry before proceeding to [CAUSAL] **tebalance summarize**, [CAUSAL] **tebalance overid**, [CAUSAL] **tebalance density**, or [CAUSAL] **tebalance box** for command-specific syntax and details.

A covariate is said to be balanced when its distribution does not vary over treatment levels.

Covariates are balanced in experimental data because treatment assignment is independent of the covariates because of the study design. In contrast, covariates must be balanced by weighting or matching in observational data because treatment assignment is related to the covariates that also affect the outcome of interest.

The estimators implemented in teffects and stteffects use a model or matching method to make the outcome conditionally independent of the treatment by conditioning on covariates. If this model or matching method is well specified, it should balance the covariates. Balance diagnostic techniques and tests check the specification of the conditioning method used by a teffects or an stteffects estimator; see [CAUSAL] teffects intro advanced for an introduction to teffects, and [CAUSAL] stteffects intro for an introduction to stteffects.

tebalance implements four methods to check for balance after teffects and stteffects. Which tebalance methods are available depends on the teffects estimation method, as summarized in the table below.

tebalance			1	Works aft	er teffect	s		rks after effects
method	Description	ipw	\mathtt{aipw}	ipwra	nnmatch	psmatch	ipw	ipwra
summarize	standardized differences and variance ratios	Х	Х	х	Х	Х	Х	Х
overid	χ^2 test for balance	X	X	X			X	X
density	diagnostic kernel density plots	X	X	X	X	X	X	X
box	diagnostic box plots				X	X		

tebalance overid implements a formal test, while the other three methods are exploratory diagnostic techniques. There is no balance check after teffects ra, stteffects ra, or stteffects wra, because they use neither a treatment model nor a matching method.

Austin (2009, 2011) and Guo and Fraser (2015, sec. 5.52) provide introductions to covariate balance. Imai and Ratkovic (2014) derived a test for balance implemented in tebalance overid.

The remainder of this entry provides a quick introduction to using tebalance to check for balance after teffects. See [CAUSAL] stteffects intro for examples after stteffects.

Example 1: Balance after estimators that use weighting

Inverse-probability-weighted (IPW) estimators use a model for the treatment to make the outcome conditionally independent of the treatment. If this model is well specified, it will also balance the covariates.

Using an extract from Cattaneo (2010), we use teffects ipw 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/r18/cattaneo2
 (Excerpt from Cattaneo (2010) Journal of Econometrics 155: 138-154)
- . teffects ipw (bweight) (mbsmoke mmarried mage prenatal1 fbaby)

Iteration 0: EE criterion = 1.873e-22
Iteration 1: EE criterion = 3.315e-26

Treatment-effects estimation Number of obs = 4,642

Estimator : inverse-probability weights

Outcome model : weighted mean Treatment model: logit

bweight	Coefficient	Robust std. err.	z	P> z	[95% conf.	interval]
ATE mbsmoke (Smoker vs Nonsmoker)	-236.1038	23.86187	-9.89	0.000	-282.8722	-189.3354
POmean mbsmoke Nonsmoker	3402.552	9.539555	356.68	0.000	3383.855	3421.249

Rubin (2008) recommends finding a model that balances the covariates before looking at results for the estimated treatment effect. Thus we do not interpret the above results, and we note that we could pay closer heed to Rubin's recommendation by preceding the teffects command with quietly to suppress the output.

Imai and Ratkovic (2014) derived a test for balance by viewing the restrictions imposed by balance as overidentifying conditions. This test is implemented in tebalance overid, and we use it to test whether the above treatment model balanced the covariates.

```
tebalance overid
Iteration 0: Criterion = .02146858
Iteration 1: Criterion =
                          .02159149
                                     (backed up)
Iteration 2: Criterion =
                          .02177783
Iteration 3: Criterion = .02260102
Iteration 4: Criterion = .02267956
Iteration 5: Criterion = .02292367
Iteration 6: Criterion = .02431655
Iteration 7: Criterion = .02457028
Iteration 8: Criterion =
                         .02488569
                          .02529483
Iteration 9: Criterion =
Iteration 10: Criterion =
                          .0254588
Iteration 11: Criterion = .02550245
Iteration 12: Criterion = .02552864
Iteration 13: Criterion = .02554462
Iteration 14: Criterion = .02554512
Iteration 15: Criterion = .02554514
Overidentification test for covariate balance
HO: Covariates are balanced
        chi2(5) = 38.1464
        Prob > chi2 = 0.0000
```

We reject the null hypothesis that the treatment model balanced the covariates.

Let's use tebalance summarize to see where the problem lies. To get an idea of the extent to which the covariates are unbalanced, we begin by summarizing the covariates by group in the raw data by specifying the baseline option.

. tebalance summarize, baseline

Covariate balance summary

		Raw	Weighted
Number of obs Treated obs	=	4,642 864	4,642.0 2,315.3
Control obs	=	3,778	2,326.7

	Me	ans	Varia	ances
	Control	Treated	Control	Treated
mmarried	.7514558	.4733796	.1868194	.2495802
mage	26.81048	25.16667	31.87141	28.10429
prenatal1	.8268925	.6898148	.1431792	.2142183
fbaby	.4531498	.3715278	.2478707	.2337654

The output indicates that the covariates may not be balanced in the raw data. For example, the distribution of the mother's age may differ over the treatment groups. We can investigate the differences further with standardized differences and variance ratios. A perfectly balanced covariate has a standardized difference of zero and variance ratio of one. There are no standard errors on these statistics, so inference is informal. Austin (2009) provides a recent introduction to these diagnostics, although they have been used at least since Rosenbaum and Rubin (1985).

By omitting the baseline option, we obtain these diagnostic statistics for the raw data and the weighted data.

. tebalance summarize

Covariate balance summary

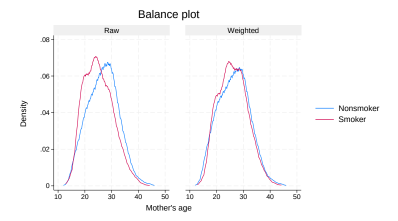
	Raw	Weighted
Number of obs = Treated obs = Control obs =	4,642 864 3,778	4,642.0 2,315.3 2,326.7

	Standardized	differences	Vari	ance ratio
	Raw	Weighted	Raw	Weighted
mmarried	5953009	0105562	1.335944	1.009079
mage	300179	0672115	.8818025	.8536401
prenatal1	3242695	0156339	1.496155	1.023424
fbaby	1663271	.0257705	.9430944	1.005698

Reviewing the output, we see that for mmarried, prenatal1, and fbaby, our model improved the level of balance. The weighted standardized differences are all close to zero and the variance ratios are all close to one. However, output indicates that mage may not be balanced by our model. The weighted standardized difference is close to zero, but the weighted variance ratio still appears to be considerably less than one.

Now, let's use tebalance density to look at how the densities of mage for treated and control groups differ.

. tebalance density mage



The plots also indicate a lack of balance in mage between the treatment groups.

To try to achieve better balance, we specify a richer model with interactions between mage and the other covariates and look at the resulting standardized differences.

- . quietly teffects ipw (bweight) (mbsmoke mmarried mage prenatal1 fbaby
- > c.mage#(c.mage i.mmarried prenatal1))
- . tebalance summarize

Covariate balance summary

	Raw	Weighted
Number of obs =	4,642	4,642.0
Treated obs =	864	2,329.1
Control obs =	3,778	2,312.9

	Standardized	differences	Varia	ance ratio
	Raw	Weighted	Raw	Weighted
mmarried	5953009	.0053497	1.335944	.9953184
mage	300179	.0410889	.8818025	1.076571
prenatal1	3242695	.0009807	1.496155	.9985165
fbaby	1663271	0130638	.9430944	.9965406
mage#	3028275	.0477465	.8274389	1.109134
mmarried# mage Married	6329701	.0197209	1.157026	1.034108
prenatal1# mage Yes	4053969	.0182109	1.226363	1.032561

A

The standardized difference and variance ratio results for mage look closer to the expected values of zero and one, so we proceed to the formal test.

```
. tebalance overid
Iteration 0: Criterion =
                           .0602349
Iteration 1:
             Criterion =
                          .06172749
                                      (backed up)
Iteration 2:
             Criterion =
                           .06428588
                                      (backed up)
Iteration 3: Criterion =
                           .06489623
                                      (backed up)
Iteration 4: Criterion =
                           .06527284
                                      (backed up)
                           .06643426
Iteration 5: Criterion =
Iteration 6: Criterion =
                           .07134338
Iteration 7: Criterion =
                          .07638414
Iteration 8: Criterion =
                           .07673211
Iteration 9: Criterion =
                           .07681959
Iteration 10: Criterion =
                            .077044
Iteration 11: Criterion =
                           .07759547
                           .07771973
Iteration 12: Criterion =
Iteration 13: Criterion =
                           .0777271
Iteration 14: Criterion =
                           .07773395
Iteration 15: Criterion =
                          .07774839
Iteration 16: Criterion =
                          .07775314
Iteration 17: Criterion = .07775324
Overidentification test for covariate balance
HO: Covariates are balanced
         chi2(8)
                     = 11.8612
         Prob > chi2 = 0.1575
```

We do not reject the null hypothesis that the specified treatment model balances the covariates.

Example 2: Balance after estimators that use matching

Instead of weighting, we might want to use a matching estimator. We can select teffects nnmatch or teffects psmatch for balance and estimation; in this example, we use teffects nnmatch.

```
. teffects nnmatch (bweight mmarried mage prenatal1 fbaby)
```

> (mbsmoke), bias(mage) ematch(mmarried prenatal1 fbaby)

Treatment-effect	s estimation	Number of obs	=	4,642
Estimator :	nearest-neighbor matching	Matches: requested	=	1
Outcome model :	matching	min	=	1
Distance metric:	Mahalanobis	max	=	139

bweight	Coefficient	AI robust std. err.	z	P> z	[95% conf.	interval]
ATE mbsmoke (Smoker vs Nonsmoker)	-240.4589	28.43008	-8.46	0.000	-296.1808	-184.7369

4

Again we ignore the estimated effect and first check for balance. We begin by reviewing whether the summary statistics indicate good balance.

. tebalance summarize (refitting the model using the generate() option) Covariate balance summary

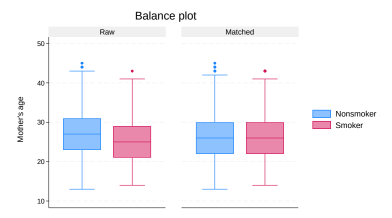
	Raw	Matched
=	4,642	9,284
=	864	4,642
=	3,778	4,642
		= 864

	Standardized differences		Varia	Variance ratio	
	Raw	Matched	Raw	Matched	
mmarried	5953009	-2.42e-16	1.335944	1	
mage	300179	0040826	.8818025	.9815517	
prenatal1	3242695	-2.78e-16	1.496155	1	
fbaby	1663271	2.24e-16	.9430944	1	

We do not have standard errors on these statistics, so we cannot make any formal conclusions, but the summary statistics appear to indicate much better balance than the IPW results. tebalance summarize has to refit the model to recover the matched sample because the generate() option was not specified on the teffects nnmatch command. The reestimation does not affect the results, although the computation takes longer; see example 3 for details.

Because it is a matching estimator, and not an IPW estimator, we cannot use tebalance overid after teffects nnmatch. The matching estimators, however, provide diagnostic box plots using tebalance box that are not available after the IPW estimators.

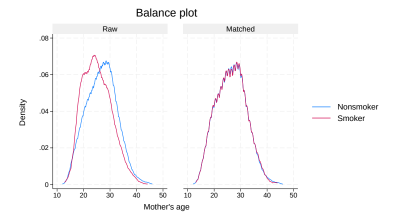
. tebalance box mage (refitting the model using the generate() option)



The box plots of the matched data also indicate covariate balance.

Let's look at the kernel density plots using the matched data.

. tebalance density mage
(refitting the model using the generate() option)



The plots using the matched data appear to be balanced.

□ Technical note

teffects implements matching estimators, IPW estimators, regression-adjustment (RA) estimators, and estimators that combine IPW and RA. Matching estimators define a matched sample, and IPW estimators define a weighted sample, each of which can be used to compute covariate balance statistics. RA estimators do not define an adjusted sample that can be used to compute covariate balance statistics, and tebalance does not work after teffects ra. Only the IPW component of the estimators that combine RA and IPW defines a weighted sample that can be used to compute balance statistics. So, tebalance produces the same results after teffects aipw or teffects ipwra as it does after teffects ipw.

4

Example 3: Faster results after a matching estimator

The tebalance commands run in example 2 executed more slowly than necessary. tebalance issued the note

refitting the model using the generate() option

after the commands

- . tebalance summarize
- . tebalance box mage

and

. tebalance density mage

After teffects nnmatch or teffects psmatch, tebalance computes the balance statistics on the matched sample defined by the matching estimator. teffects nnmatch and teffects psmatch leave behind only variables that identify the matched sample when the generate() option is specified. Unless the generate() option is specified with teffects nnmatch or teffects psmatch, each tebalance command must rerun the estimation command to recover the matched sample.

Typing

- . teffects nnmatch (bweight mmarried mage fbaby prenatal1)
- > (mbsmoke), bias(mage) ematch(mmarried fbaby prenatal1)
- > generate(matchv)

would generate variables that identify the matched sample that the tebalance commands could use. See *Remarks and examples* in [CAUSAL] **tebalance box**, [CAUSAL] **tebalance density**, and [CAUSAL] **tebalance summarize** for examples using the option generate() on teffects psmatch to speed up the postestimation computations.

4

Methods and formulas

Methods and formulas are presented under the following headings:

Introduction
Matched samples
IPW samples
Testing the propensity-score model specification

Introduction

For covariate z, we observe values $\{z_1, z_2, \ldots, z_N\}$. Define a treatment indicator variable for J treatment levels as $t_i \in \{1, 2, \ldots, J\}$, for $i = 1, \ldots, N$, and frequency weights as $\{w_1, w_2, \ldots, w_N\}$. The sample mean and variance of z for level t are

$$\widehat{\mu}_z(t) = \frac{\sum\limits_{i}^{N} I(t_i = t) w_i z_i}{N_t} \quad \text{and}$$

$$\widehat{\sigma}_z^2(t) = \frac{\sum\limits_{i}^{N} I(t_i = t) w_i \left\{ z_i - \widehat{\mu}_z(t) \right\}^2}{N_t - 1}$$

where $N_t = \sum_{i}^{N} w_i I(t_i = t)$, and

$$I(t_i = t) = \begin{cases} 1 & \text{if } t_i = t \\ 0 & \text{otherwise} \end{cases}$$

As shown in Austin (2011), the standardized differences for covariate z between level t and the control t_0 are computed as

$$\delta_z(t) = \frac{\widehat{\mu}_z(t) - \widehat{\mu}_z(t_0)}{\sqrt{\frac{\widehat{\sigma}_z^2(t) + \widehat{\sigma}_z^2(t_0)}{2}}}$$
(1)

The variance ratio is $\rho_z(t) = {\{\widehat{\sigma}_z^2(t)\}}/{\{\widehat{\sigma}_z^2(t_0)\}}$.

Matched samples

We now turn our attention to the matched samples for the potential-outcome mean (POM), average treatment effect (ATE), and average treatment effect on the treated (ATET) estimators. We estimate the covariate for the counter-factual treatment by taking the mean of the matched observations

$$\dot{z}_i = \frac{\sum_{j \in \Omega(i)} w_j z_j}{\sum_{j \in \Omega(i)} w_j}$$

where $\Omega(i)=(k_1,k_2,\ldots,k_{m_j})$ is the set of observation indices that are matched to observation i of the opposite treatment condition. The observed covariate and matched covariate pairs, (z_i,\dot{z}_i) , $i=1,\ldots,N$, are used in the box plot (see [G-2] **graph box**) and the kernel density plot (see [R] **kdensity**). The ATET sample is limited to those observations from the conditional treatment, \widetilde{t} , and their matched covariate means.

In Methods and formulas of [CAUSAL] **teffects nnmatch**, we define $K_m(i)$ as the number of times observation i is used in a match with observation j of the opposite treatment condition, $i \in \Omega(j)$, weighted by the total number of matches for observation j. Specifically,

$$K_{m}(i) = \sum_{j=1}^{N} I\left\{i \in \Omega\left(j\right)\right\} \frac{w_{j}}{\sum\limits_{k \in \Omega\left(j\right)} w_{k}}$$

These weights are used in the estimation of the mean and variance for the matched dataset. For the POM and ATE models, the estimated mean and variance are computed as

$$\begin{split} \widehat{\mu}_{\dot{z}}(t) &= \frac{\sum_{i}^{N} I(t_{i} = t) w_{i} z_{i} \left\{1 + K_{m}(i)\right\}}{M_{t}} \quad \text{ and } \\ \widehat{\sigma}_{\dot{z}}^{2}(t) &= \frac{\sum_{i}^{N} I(t_{i} = t) w_{i} \{1 + K_{m}(i)\} \{z_{i} - \widehat{\mu}_{\dot{z}}(t)\}^{2}}{M_{t} - 1} \end{split}$$

where
$$M_t = \sum_{i=1}^{N} I(t_i = t) w_i \{1 + K_m(i)\}.$$

The standardized differences between the control level and all other levels for the matched covariate distribution are computed as in (1), but $\widehat{\mu}_{\dot{z}}(t)$ is substituted for $\widehat{\mu}_{z}(t)$ and $\widehat{\sigma}_{z}^{2}(t)$ for $\widehat{\sigma}_{z}^{2}(t)$.

For the ATET model, there is no matched sample for the treatment levels other than the conditional treatment \widetilde{t} . The covariate mean and variance for the conditional treatment are the same as that of the raw data, $\mu_z(\widetilde{t})$ and $\sigma_z(\widetilde{t})$. However, the covariate mean and variance for the sample matched to the conditional treatment, $t \neq \widetilde{t}$, are computed as

$$\begin{split} \widetilde{\mu}_{\dot{z}}(t) &= \frac{\sum_{i}^{N} I(t_i = t) w_i z_i K_m(i)}{M_t} \quad \text{ and } \\ \widetilde{\sigma}_{\dot{z}}^2(t) &= \frac{\sum_{i}^{N} I(t_i = t) w_i K_m(i) \left\{ z_i - \widetilde{\mu}_{\dot{z}}(t) \right\}^2}{M_t - 1} \end{split}$$

where
$$M_t = \sum_{i=1}^{N} I(t_i = t) w_i K_m(i)$$
.

IPW samples

Computation of the inverse-probability weights is discussed in Methods and formulas of [CAUSAL] teffects aipw and in Methods and formulas of [CAUSAL] stteffects ipwra. For the POM and ATE estimators, we defined the normalized IPW weights as $\overline{d}_i(t) = N_t d_i(t) / \sum_{i=1}^{N} d_i(t)$, where $d_i(t) = I(t_i = t)/p(\mathbf{z}_i, t, \widehat{\boldsymbol{\gamma}})$ for treatment level t and individual i.

For the ATET estimator, we use the normalized weights $\overline{f}_i = N f_i / \sum_i^N f_i$, where $f_i =$ $p(\mathbf{z}_i, \widetilde{t}, \widehat{\gamma})/p(\mathbf{z}_i, t_i, \widehat{\gamma})$ are the treatment-adjusted inverse-probability weights, and \widetilde{t} is the conditional treatment.

We will simplify notation by defining a single weight

$$\overline{w}_i(t) = \begin{cases} \overline{d}_i(t) & \text{if model is ATE or POM} \\ \overline{f}_i(t) & \text{if model is ATET} \end{cases}$$

The covariate mean and variance for treatment level t are

$$\widetilde{\mu}_{\dot{z}}(t) = \frac{\sum_{i}^{N} I(t_{i} = t) w_{i} \overline{w}_{i} x_{i}}{M_{t}} \quad \text{and}$$

$$\widetilde{\sigma}_{\dot{z}}^{2}(t) = \frac{I(t_{i} = t) w_{i} \overline{w}_{i} \left\{ z_{i} - \widetilde{\mu}_{\dot{z}}(t) \right\}^{2}}{M_{t} - 1}$$

where $M_t = \sum_{i=1}^{N} I(t_i = t) w_i \overline{w}_i$.

The kernel density is computed by kdensity for each covariate conditioned on each treatment level using the raw covariate with iweights equal to $w_i \overline{w}_i$.

Testing the propensity-score model specification

We estimate the probability of treatment conditioned on a set of covariates with a propensityscore model. Imai and Ratkovic (2014) derive a test for whether the estimated propensity score balances the covariates. The score equations for parameters of the propensity-score model define an exactly identified generalized method of moments (GMM) estimator. Imai and Ratkovic (2014) use the conditions imposed by mean balance as overidentifying conditions. A standard GMM test for the validity of the overidentifying conditions is then a test for covariate balance. See [R] gmm for a discussion of this overidentifying test, which is known as Hansen's J test in the econometrics literature.

Here are the details about the score equations and the overidentifying balance conditions. Recall from Methods and formulas of [CAUSAL] teffects aipw and Methods and formulas of [CAUSAL] stteffects ipwra, we have the first-order condition of the treatment model

$$\frac{1}{N} \sum_{i=1}^{N} \mathbf{s}_{\text{tm},i}(\mathbf{z}_i, \widehat{\boldsymbol{\gamma}}) = 0$$

For a two-level treatment-effects model with conditional treatment \widetilde{t} and control t_0 , the score is

$$\mathbf{s}_{\text{tm},i}(\mathbf{z}_{i},\boldsymbol{\gamma}) = \frac{I\left(t_{i} = \widetilde{t}\right)}{p(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma})} \frac{\partial p(\mathbf{z}_{i},t,\boldsymbol{\gamma})}{\partial \boldsymbol{\gamma}'} - \left\{ \frac{I(t_{i} = t_{0})}{1 - p\left(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma}\right)} \right\} \left. \frac{\partial p\left(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma}\right)}{\partial \boldsymbol{\gamma}'} \right|_{\boldsymbol{\gamma} = \widehat{\boldsymbol{\gamma}}}$$

The score reduces to

$$\mathbf{s}_{\text{tm},i}\left(\mathbf{z}_{i},\widehat{\boldsymbol{\gamma}}\right) = \left[\frac{I\left(t_{i} = \widetilde{t}\right) - p\left(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma}\right)}{p\left(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma}\right)\left\{1 - p\left(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma}\right)\right\}}\right] \frac{\partial p\left(\mathbf{z}_{i},\widetilde{t},\boldsymbol{\gamma}\right)}{\partial \boldsymbol{\gamma}'}\bigg|_{\boldsymbol{\gamma} = \widehat{\boldsymbol{\gamma}}}$$

The corresponding covariate balancing moment conditions are

$$\mathbf{w}_{\mathrm{tm},i}(\mathbf{z}_{i},\boldsymbol{\gamma}) = \left[\frac{I\left(t_{i} = \widetilde{t}\right) - p\left(\mathbf{z}_{i}, \widetilde{t}, \boldsymbol{\gamma}\right)}{p\left(\mathbf{z}_{i}, \widetilde{t}, \boldsymbol{\gamma}\right) \left\{1 - p\left(\mathbf{z}_{i}, \widetilde{t}, \boldsymbol{\gamma}\right)\right\}} \right] \mathbf{z}_{i}$$

for the POM and ATE models. For the ATET model with conditional treatment \widetilde{t} , we multiply by $p(\mathbf{z}_i, \widetilde{t}, \gamma)$ and scale by $N/N_{\widetilde{t}}$:

$$\mathbf{w}_{\mathrm{tm},i}(\mathbf{z}_{i},\boldsymbol{\gamma}) = \frac{N}{N_{\widetilde{t}}} \left\{ \frac{I\left(t_{i} = \widetilde{t}\right) - p\left(\mathbf{z}_{i}, \widetilde{t}, \boldsymbol{\gamma}\right)}{1 - p(\mathbf{z}_{i}, \widetilde{t}, \boldsymbol{\gamma})} \right\} \mathbf{z}_{i}$$

We stack the moment conditions

$$\mathbf{g}_{\text{tm}}(\mathbf{Z}, \boldsymbol{\gamma}) = \frac{1}{N} \sum_{i=1}^{N} \left\{ \begin{aligned} \mathbf{s}_{\text{tm},i}(\mathbf{z}_{i}, \boldsymbol{\gamma}) \\ \mathbf{w}_{\text{tm},i}(\mathbf{z}_{i}, \boldsymbol{\gamma}) \end{aligned} \right\}$$
$$= \frac{1}{N} \sum_{i=1}^{N} \mathbf{g}_{\text{tm},i}(\mathbf{z}_{i}, \boldsymbol{\gamma})$$

The overidentified GMM estimator is then

$$\widetilde{\gamma} = \operatorname{argmin}_{\gamma} N \ \mathbf{g}_{tm}(\mathbf{Z}, \gamma)' \ \mathbf{W}_{tm}(\mathbf{Z}, \gamma)^{-1} \ \mathbf{g}_{tm}(\mathbf{Z}, \gamma)$$
 (2)

where

$$\mathbf{W}_{\mathrm{tm}}(\mathbf{Z}, \boldsymbol{\gamma}) = \frac{1}{N} \sum_{i=1}^{N} E_{T} \left\{ \mathbf{g}_{\mathrm{tm}, i}(\mathbf{z}, \boldsymbol{\gamma}) \ \mathbf{g}_{\mathrm{tm}, i}(\mathbf{z}, \boldsymbol{\gamma})' \right\}$$

and the expectation is taken with respect to treatment distribution. The weight matrix $\mathbf{W}_{\mathrm{tm}}(\mathbf{Z}, \gamma)$ is computed explicitly (Imai and Ratkovic 2014), and (2), written as a maximization problem, is solved using m1.

Finally, Hansen's J statistic is evaluated at its minimum,

$$J = N\mathbf{g}_{\mathrm{tm}}(\mathbf{Z}, \widetilde{\boldsymbol{\gamma}})' \ \mathbf{W}_{\mathrm{tm}}(\mathbf{Z}, \widetilde{\boldsymbol{\gamma}})^{-1} \ \mathbf{g}_{\mathrm{tm}}(\mathbf{Z}, \widetilde{\boldsymbol{\gamma}})$$

and is asymptotically distributed χ^2 with degrees of freedom d,

$$d = \operatorname{rank}\left\{\mathbf{W}_{\operatorname{tm}}\left(\mathbf{Z},\widetilde{\boldsymbol{\gamma}}\right)\right\} - \operatorname{rank}\left[\frac{1}{N}\sum_{i=1}^{N}E_{T}\left\{\mathbf{s}_{\operatorname{tm},i}(\mathbf{z}_{i},\widetilde{\boldsymbol{\gamma}})\ \mathbf{s}_{\operatorname{tm},i}\left(\mathbf{z}_{i},\widetilde{\boldsymbol{\gamma}}\right)'\right\}\right]$$

References

- Austin, P. C. 2009. Balance diagnostics for comparing the distribution of baseline covariates between treatment groups in propensity-score matched samples. *Statistics in Medicine* 28: 3083–3107. https://doi.org/10.1002/sim.3697.
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Also see

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

[CAUSAL] tebalance box — Covariate balance box

[CAUSAL] tebalance density — Covariate balance density

[CAUSAL] tebalance overid — Test for covariate balance

[CAUSAL] **tebalance summarize** — Covariate-balance summary statistics

[CAUSAL] teffects — Treatment-effects estimation for observational data

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