bootstrap — Bootstrap sampling and estimation

Description

bootstrap performs nonparametric bootstrap estimation of specified statistics (or expressions) for a Stata command or a user-written program. Statistics are bootstrapped by resampling the data in memory with replacement. bootstrap is designed for use with nonestimation commands, functions of coefficients, or user-written programs. To bootstrap coefficients, we recommend using the vce(bootstrap) option when allowed by the estimation command.

Quick start

Bootstrap the mean of v1 returned by summarize in r(mean)

```
bootstrap mean=r(mean): summarize v1
```

Bootstrap the statistic r(mystat) returned by program myprog1

```
bootstrap stat=r(mystat): myprog1 v1
```

As above, but use 100 replications

```
bootstrap stat=r(mystat), reps(100): myprog1 v1
```

As above, and save the results from each replication in mydata.dta

```
bootstrap stat=r(mystat), reps(100) saving(mydata): myprog1 v1
```

Bootstrap a difference in coefficients estimated by regress

```
bootstrap diff=(_b[x2]-_b[x1]): regress y x1 x2 x3
```

Bootstrap the coefficients stored in e(b) by myprog2

```
bootstrap _b: myprog2 y x1 x2 x3
```

As above, but with bootstrap samples taken independently within strata identified by svar

```
bootstrap _b, strata(svar): myprog2 y x1 x2 x3
```

Resample clusters defined by cvar and create newcvar identifying resampled clusters

```
bootstrap _b, cluster(cvar) idcluster(newcvar): myprog2 y x1 x2 x3
```

Menu

Statistics > Resampling > Bootstrap estimation
## Syntax

```
bootstrap exp_list [ , options eform_option ] : command
```

<table>
<thead>
<tr>
<th>options</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>reps(#)</code></td>
<td>perform # bootstrap replications; default is <code>reps(50)</code></td>
</tr>
<tr>
<td><code>strata(varlist)</code></td>
<td>variables identifying strata</td>
</tr>
<tr>
<td><code>size(#)</code></td>
<td>draw samples of size #; default is <code>_N</code></td>
</tr>
<tr>
<td><code>cluster(varlist)</code></td>
<td>variables identifying resampling clusters</td>
</tr>
<tr>
<td><code>idcluster(newvar)</code></td>
<td>create new cluster ID variable</td>
</tr>
<tr>
<td><code>saving(filename, ...)</code></td>
<td>save statistics to <code>filename</code>; save statistics in double precision; save results to <code>filename</code> every # replications</td>
</tr>
<tr>
<td><code>bca</code></td>
<td>compute acceleration for BCa confidence intervals</td>
</tr>
<tr>
<td><code>ties</code></td>
<td>adjust BC/BCa confidence intervals for ties</td>
</tr>
<tr>
<td><code>mse</code></td>
<td>use MSE formula for variance estimation</td>
</tr>
<tr>
<td><code>level(#)</code></td>
<td>set confidence level; default is <code>level(95)</code></td>
</tr>
<tr>
<td><code>notable</code></td>
<td>suppress table of results</td>
</tr>
<tr>
<td><code>noheader</code></td>
<td>suppress table header</td>
</tr>
<tr>
<td><code>nolegend</code></td>
<td>suppress table legend</td>
</tr>
<tr>
<td><code>verbose</code></td>
<td>display the full table legend</td>
</tr>
<tr>
<td><code>nodots</code></td>
<td>suppress replication dots</td>
</tr>
<tr>
<td><code>dots(#)</code></td>
<td>display dots every # replications</td>
</tr>
<tr>
<td><code>noisily</code></td>
<td>display any output from <code>command</code></td>
</tr>
<tr>
<td><code>trace</code></td>
<td>trace <code>command</code></td>
</tr>
<tr>
<td><code>title(text)</code></td>
<td>use <code>text</code> as title for bootstrap results</td>
</tr>
<tr>
<td><code>display_options</code></td>
<td>control columns and column formats, row spacing, line width, display of omitted variables and base and empty cells, and factor-variable labeling</td>
</tr>
<tr>
<td><code>eform_option</code></td>
<td>display coefficient table in exponentiated form</td>
</tr>
<tr>
<td><code>nodrop</code></td>
<td>do not drop observations</td>
</tr>
<tr>
<td><code>nowarn</code></td>
<td>do not warn when <code>e(sample)</code> is not set</td>
</tr>
<tr>
<td><code>force</code></td>
<td>do not check for <code>weights</code> or <code>svy</code> commands; seldom used</td>
</tr>
<tr>
<td><code>reject(exp)</code></td>
<td>identify invalid results</td>
</tr>
<tr>
<td><code>seed(#)</code></td>
<td>set random-number seed to #</td>
</tr>
<tr>
<td><code>group(varname)</code></td>
<td>ID variable for groups within <code>cluster()</code></td>
</tr>
<tr>
<td><code>jackknifeopts(jkopts)</code></td>
<td>options for <code>jackknife</code>; see [R] <code>jackknife</code></td>
</tr>
<tr>
<td><code>coeflegend</code></td>
<td>display legend instead of statistics</td>
</tr>
</tbody>
</table>
command is any command that follows standard Stata syntax. weights are not allowed in command.

group(), jackknifeopts(), and coeflegend do not appear in the dialog box.

See [U] 20 Estimation and postestimation commands for more capabilities of estimation commands.

exp_list contains (name: elist)
    elist
eexp
eexp is specname
    [eqno]specname
specname is _b
    _b[]
    _se
    _se[]
eqno is ##
    name

exp is a standard Stata expression; see [U] 13 Functions and expressions.

Distinguish between [], which are to be typed, and [ ], which indicate optional arguments.

Options

reps(#) specifies the number of bootstrap replications to be performed. The default is 50. A total of 50–200 replications are generally adequate for estimates of standard error and thus are adequate for normal-approximation confidence intervals; see Mooney and Duval (1993, 11). Estimates of confidence intervals using the percentile or bias-corrected methods typically require 1,000 or more replications.

strata(varlist) specifies the variables that identify strata. If this option is specified, bootstrap samples are taken independently within each stratum.

size(#) specifies the size of the samples to be drawn. The default is _N, meaning to draw samples of the same size as the data. If specified, # must be less than or equal to the number of observations within strata().

If cluster() is specified, the default size is the number of clusters in the original dataset. For unbalanced clusters, resulting sample sizes will differ from replication to replication. For cluster sampling, # must be less than or equal to the number of clusters within strata().

ccluster(varlist) specifies the variables that identify resampling clusters. If this option is specified, the sample drawn during each replication is a bootstrap sample of clusters.

idcluster(newvar) creates a new variable containing a unique identifier for each resampled cluster. This option requires that ccluster() also be specified.
saving( filename[, suboptions] ) creates a Stata data file (.dta file) consisting of (for each statistic in exp_list) a variable containing the replicates.

**double** specifies that the results for each replication be saved as **doubles**, meaning 8-byte reals. By default, they are saved as **floats**, meaning 4-byte reals. This option may be used without the **saving()** option to compute the variance estimates by using double precision.

**every(#)** specifies that results be written to disk every #th replication. **every()** should be specified only in conjunction with **saving()** when **command** takes a long time for each replication. This option will allow recovery of partial results should some other software crash your computer. See [P] **postfile**.

**replace** specifies that **filename** be overwritten if it exists. This option does not appear in the dialog box.

**bca** specifies that **bootstrap** estimate the acceleration of each statistic in **exp_list**. This estimate is used to construct BCa confidence intervals. Type **estat bootstrap, bca** to display the BCa confidence interval generated by the bootstrap command.

**ties** specifies that **bootstrap** adjust for ties in the replicate values when computing the median bias used to construct BC and BCa confidence intervals.

**mse** specifies that **bootstrap** compute the variance by using deviations of the replicates from the observed value of the statistics based on the entire dataset. By default, **bootstrap** computes the variance by using deviations from the average of the replicates.

**level(#) ; see [R] Estimation options.**

**notable** suppresses the display of the table of results.

**noheader** suppresses the display of the table header. This option implies **nolegend**. This option may also be specified when replaying estimation results.

**nolegend** suppresses the display of the table legend. This option may also be specified when replaying estimation results.

**verbose** specifies that the full table legend be displayed. By default, coefficients and standard errors are not displayed. This option may also be specified when replaying estimation results.

**nodots** and **dots(#)** specify whether to display replication dots. By default, one dot character is displayed for each successful replication. A red ‘x’ is displayed if **command** returns an error or if any value in **exp_list** is missing. You can also control whether dots are displayed using **set dots**; see [R] **set**.

**nodots** suppresses display of the replication dots.

**dots(#)** displays dots every # replications. **dots(0)** is a synonym for **nodots**.

**noisily** specifies that any output from **command** be displayed. This option implies the **nodots** option.

**trace** causes a trace of the execution of **command** to be displayed. This option implies the **noisily** option.

**title(text)** specifies a title to be displayed above the table of bootstrap results. The default title is the title stored in **e(title)** by an estimation command, or if **e(title)** is not filled in, Bootstrap results is used. **title()** may also be specified when replaying estimation results.
display_options: noci, nopvalues, noomitted, vsquish, noemptycells, baselevels, allbaselevels, nofvlabel, fvwrap( ), fvwrapon(style), cformat(%,fmt), pformat(%,fmt), sformat(%,fmt), and nolstretch; see [R] Estimation options.

eform_option causes the coefficient table to be displayed in exponentiated form; see [R] eform_option.

command determines which of the following are allowed (eform(string) and eform are always allowed):

<table>
<thead>
<tr>
<th>eform_option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>eform(string)</td>
<td>use string for the column title</td>
</tr>
<tr>
<td>eform</td>
<td>exponentiated coefficient, string is ( \exp(b) )</td>
</tr>
<tr>
<td>hr</td>
<td>hazard ratio, string is Haz. Ratio</td>
</tr>
<tr>
<td>shr</td>
<td>subhazard ratio, string is SHR</td>
</tr>
<tr>
<td>irr</td>
<td>incidence-rate ratio, string is IRR</td>
</tr>
<tr>
<td>or</td>
<td>odds ratio, string is Odds Ratio</td>
</tr>
<tr>
<td>rrr</td>
<td>relative-risk ratio, string is RRR</td>
</tr>
</tbody>
</table>

Advanced

nodrop prevents observations outside e(sample) and the if and in qualifiers from being dropped before the data are resampled.

nowarn suppresses the display of a warning message when command does not set e(sample).

force suppresses the restriction that command not specify weights or be a svy command. This is a rarely used option. Use it only if you know what you are doing.

reject(exp) identifies an expression that indicates when results should be rejected. When exp is true, the resulting values are reset to missing values.

seed(#) sets the random-number seed. Specifying this option is equivalent to typing the following command prior to calling bootstrap:

```
. set seed #
```

The following options are available with bootstrap but are not shown in the dialog box:

group(varname) re-creates varname containing a unique identifier for each group across the resampled clusters. This option requires that idcluster() also be specified.

This option is useful for maintaining unique group identifiers when sampling clusters with replacement. Suppose that cluster 1 contains 3 groups. If the idcluster(newclid) option is specified and cluster 1 is sampled multiple times, newclid uniquely identifies each copy of cluster 1. If group(newgroupid) is also specified, newgroupid uniquely identifies each copy of each group.

jackknifeopts(jkopts) identifies options that are to be passed to jackknife when it computes the acceleration values for the BC\textsubscript{a} confidence intervals; see [R] jackknife. This option requires the bca option and is mostly used for passing the eclass, rclass, or n(#) option to jackknife.

coflegend; see [R] Estimation options.
Remarks and examples

Remarks are presented under the following headings:

- Introduction
- Using bootstrap
- Regression coefficients
- Expressions
- Combining bootstrap datasets
- A note about macros
- Achieved significance level
- Bootstrapping a ratio
- Warning messages and e(sample)
- Bootstrapping statistics from data with a complex structure

Introduction

With few assumptions, bootstrapping provides a way of estimating standard errors and other measures of statistical precision (Efron 1979; Efron and Stein 1981; Efron 1982; Efron and Tibshirani 1986; Efron and Tibshirani 1993; also see Davison and Hinkley [1997]; Guan [2003]; Mooney and Duval [1993]; Poi [2004]; and Stine [1990]). It provides a way to obtain such measures when no formula is otherwise available or when available formulas make inappropriate assumptions. Cameron and Trivedi (2010, chap. 13) discuss many bootstrapping topics and demonstrate how to do them in Stata.

To illustrate bootstrapping, suppose that you have a dataset containing \( N \) observations and an estimator that, when applied to the data, produces certain statistics. You draw, with replacement, \( N \) observations from the \( N \)-observation dataset. In this random drawing, some of the original observations will appear once, some more than once, and some not at all. Using the resampled dataset, you apply the estimator and collect the statistics. This process is repeated many times; each time, a new random sample is drawn and the statistics are recalculated.

This process builds a dataset of replicated statistics. From these data, you can calculate the standard error by using the standard formula for the sample standard deviation

\[
\hat{se} = \left\{ \frac{1}{k-1} \sum (\hat{\theta}_i - \overline{\theta})^2 \right\}^{1/2}
\]

where \( \hat{\theta}_i \) is the statistic calculated using the \( i \)th bootstrap sample and \( k \) is the number of replications. This formula gives an estimate of the standard error of the statistic, according to Hall and Wilson (1991). Although the average, \( \overline{\theta} \), of the bootstrapped estimates is used in calculating the standard deviation, it is not used as the estimated value of the statistic itself. Instead, the original observed value of the statistic, \( \hat{\theta} \), is used, meaning the value of the statistic computed using the original \( N \) observations.

You might think that \( \overline{\theta} \) is a better estimate of the parameter than \( \hat{\theta} \), but it is not. If the statistic is biased, bootstrapping exaggerates the bias. In fact, the bias can be estimated as \( \overline{\theta} - \hat{\theta} \) (Efron 1982, 33). Knowing this, you might be tempted to subtract this estimate of bias from \( \hat{\theta} \) to produce an unbiased statistic. The bootstrap bias estimate has an indeterminate amount of random error, so this unbiased estimator may have greater mean squared error than the biased estimator (Mooney and Duval 1993; Hinkley 1978). Thus, \( \hat{\theta} \) is the best point estimate of the statistic.

The logic behind the bootstrap is that all measures of precision come from a statistic’s sampling distribution. When the statistic is estimated on a sample of size \( N \) from some population, the sampling distribution tells you the relative frequencies of the values of the statistic. The sampling distribution, in turn, is determined by the distribution of the population and the formula used to estimate the statistic.
Sometimes the sampling distribution can be derived analytically. For instance, if the underlying population is distributed normally and you calculate means, the sampling distribution for the mean is also normal but has a smaller variance than that of the population. In other cases, deriving the sampling distribution is difficult, such as when means are calculated from nonnormal populations. Sometimes, as in the case of means, it is not too difficult to derive the sampling distribution as the sample size goes to infinity \((N \to \infty)\). However, such asymptotic distributions may not perform well when applied to finite samples.

If you knew the population distribution, you could obtain the sampling distribution by simulation: you could draw random samples of size \(N\), calculate the statistic, and make a tally. Bootstrapping does precisely this, but it uses the observed distribution of the sample in place of the true population distribution. Thus, the bootstrap procedure hinges on the assumption that the observed distribution is a good estimate of the underlying population distribution. In return, the bootstrap produces an estimate, called the bootstrap distribution, of the sampling distribution. From this, you can estimate the standard error of the statistic, produce confidence intervals, etc.

The accuracy with which the bootstrap distribution estimates the sampling distribution depends on the number of observations in the original sample and the number of replications in the bootstrap. A crudely estimated sampling distribution is adequate if you are going to extract, say, only a standard error. A better estimate is needed if you want to use the 2.5th and 97.5th percentiles of the distribution to produce a 95% confidence interval. To extract many features simultaneously about the distribution, an even better estimate is needed. Generally, replications on the order of 1,000 produce very good estimates, but only 50–200 replications are needed for estimates of standard errors. See Poi (2004) for a method to choose the number of bootstrap replications.

**Using bootstrap**

Typing

```
    . bootstrap exp_list, reps(#) : command
```

executes `command` multiple times, bootstrapping the statistics in `exp_list` by resampling observations (with replacement) from the data in memory \# times. This method is commonly referred to as the nonparametric bootstrap.

`command` defines the statistical command to be executed. Most Stata commands and user-written programs can be used with `bootstrap`, as long as they follow standard Stata syntax; see [U] 11 Language syntax. If the `bca` option is supplied, `command` must also work with `jackknife`; see [R] jackknife. The by prefix may not be part of `command`.

`exp_list` specifies the statistics to be collected from the execution of `command`. If `command` changes the contents in `e(b)`, `exp_list` is optional and defaults to `_b`.

Because bootstrapping is a random process, if you want to be able to reproduce results, set the random-number seed by specifying the `seed(#)` option or by typing

```
    . set seed #
```

where \# is a seed of your choosing, before running `bootstrap`; see [R] set seed.
Regression coefficients

Example 1

Let’s say that we wish to compute bootstrap estimates for the standard errors of the coefficients from the following regression:

```stata
use https://www.stata-press.com/data/r16/auto
(1978 Automobile Data)
regress mpg weight gear foreign
```

To run the bootstrap, we simply prefix the above regression command with the `bootstrap` command (specifying its options before the colon separator). We must set the random-number seed before calling `bootstrap`.

```stata
bootstrap, reps(100) seed(1): regress mpg weight gear foreign
```

The displayed confidence interval is based on the assumption that the sampling (and hence bootstrap) distribution is approximately normal (see *Methods and formulas* below). Because this confidence interval is based on the standard error, it is a reasonable estimate if normality is approximately true, even for a few replications. Other types of confidence intervals are available after `bootstrap`; see [R] bootstrap postestimation.
We could instead supply names to our expressions when we run `bootstrap`. For example,

```stata
   bootstrap diff = (_b[weight] - _b[gear]): regress mpg weight gear foreign
```

would bootstrap a statistic, named `diff`, equal to the difference between the coefficients on `weight` and `gear`. 

### Technical note

`regress`, like many estimation commands, allows the `vce(bootstrap)` option. For any estimation command that allows this option, we recommend using `vce(bootstrap)` over `bootstrap` because the estimation command automatically handles clustering and other model-specific details for you.

### Expressions

#### Example 2

When we use `bootstrap`, the list of statistics can contain complex expressions, as long as each expression is enclosed in parentheses. For example, to bootstrap the range of a variable `x`, we could type

```stata
   bootstrap range = (r(max) - r(min)), reps(1000): summarize x
```

Of course, we could also bootstrap the minimum and maximum and later compute the range.

```stata
   bootstrap max = r(max), min = r(min), reps(1000) saving(mybs): summarize x
   use mybs, clear
   (bootstrap: summarize)
   generate range = max - min
   bstat range, stat(19.5637501)
```

The difference between the maximum and minimum of `x` in the sample is 19.5637501.

The `stat()` option to `bstat` specifies the observed value of the statistic (`range`) to be summarized. This option is useful when, as shown above, the statistic of ultimate interest is not specified directly to `bootstrap` but instead is calculated by other means.

Here the observed values of `r(max)` and `r(min)` are stored as characteristics of the dataset created by `bootstrap` and are thus available for retrieval by `bstat`; see `[R] bstat`. The observed range, however, is unknown to `bstat`, so it must be specified.

### Combining bootstrap datasets

You can combine two datasets from separate runs of `bootstrap` by using `append` (see `[D] append`) and then get the bootstrap statistics for the combined datasets by running `bstat`. The runs must have been performed independently (having different starting random-number seeds), and the original dataset, command, and bootstrap statistics must have been all the same.
A note about macros

In example 2, we executed the command

```
. bootstrap max=r(max) min=r(min), reps(1000) saving(mybs): summarize x
```

We did not enclose `r(max)` and `r(min)` in single quotes, as we would in most other contexts, because it would not produce what was intended:

```
. bootstrap `r(max)' `r(min)' , reps(1000) saving(mybs): summarize x
```

To understand why, note that `r(max)' , like any reference to a local macro, will evaluate to a literal string containing the contents of `r(max)` before bootstrap is even executed. Typing the command above would appear to Stata as if we had typed

```
. bootstrap 14.5441234 33.4393293, reps(1000) saving(mybs): summarize x
```

Even worse, the current contents of `r(min)` and `r(max)` could be empty, producing an even more confusing result. To avoid this outcome, refer to statistics by name (for example, `r(max)`) and not by value (for example, ``r(max)'`).

Achieved significance level

Example 3

Suppose that we wish to estimate the achieved significance level (ASL) of a test statistic by using the bootstrap. ASL is another name for p-value. An example is

\[
\text{ASL} = \Pr (\hat{\theta}^* \geq \hat{\theta} | H_0)
\]

for an upper-tailed, alternative hypothesis, where \( H_0 \) denotes the null hypothesis, \( \hat{\theta} \) is the observed value of the test statistic, and \( \hat{\theta}^* \) is the random variable corresponding to the test statistic, assuming that \( H_0 \) is true.

Here we will compare the mean miles per gallon (mpg) between foreign and domestic cars by using the two-sample \( t \) test with unequal variances. The following results indicate the \( p \)-value to be 0.0034 for the two-sided test using Satterthwaite’s approximation. Thus, assuming that mean mpg is the same for foreign and domestic cars, we would expect to observe a \( t \) statistic more extreme (in absolute value) than 3.1797 in about 0.3% of all possible samples of the type that we observed. Thus, we have evidence to reject the null hypothesis that the means are equal.
.ttest mpg, by(foreign) unequal
Two-sample t test with unequal variances

<table>
<thead>
<tr>
<th>Group</th>
<th>Obs</th>
<th>Mean</th>
<th>Std. Err.</th>
<th>Std. Dev.</th>
<th>[95% Conf. Interval]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Domestic</td>
<td>52</td>
<td>19.82692</td>
<td>0.657777</td>
<td>4.743297</td>
<td>18.50638  21.14747</td>
</tr>
<tr>
<td>Foreign</td>
<td>22</td>
<td>24.77273</td>
<td>1.40951</td>
<td>6.611187</td>
<td>21.84149  27.70396</td>
</tr>
<tr>
<td>combined</td>
<td>74</td>
<td>21.2973</td>
<td>0.672551</td>
<td>5.785503</td>
<td>19.9569  22.63769</td>
</tr>
</tbody>
</table>

| diff    | -4.945804 | 1.555438 | -8.120053 | -1.771556 |

Ho: diff = 0  
Ha: diff < 0  
Ha: diff != 0  
Ha: diff > 0

Satterthwaite’s degrees of freedom = 30.5463
Pr(T < t) = 0.0017  Pr(|T| > |t|) = 0.0034  Pr(T > t) = 0.9983

We also place the value of the test statistic in a scalar for later use.
.scalar tobs = r(t)

Efron and Tibshirani (1993, 224) describe an alternative to Satterthwaite’s approximation that estimates the ASL by bootstrapping the statistic from the test of equal means. Their idea is to recenter the two samples to the combined sample mean so that the data now conform to the null hypothesis but that the variances within the samples remain unchanged.

.summarize mpg, meanonly
.scalar omean = r(mean)
.summarize mpg if foreign==0, meanonly
.replace mpg = mpg - r(mean) + scalar(omean) if foreign==0

variable mpg was int now float  
(52 real changes made)
.summarize mpg if foreign==1, meanonly
.replace mpg = mpg - r(mean) + scalar(omean) if foreign==1
(22 real changes made)
.sort foreign
.by foreign: summarize mpg

-> foreign = Domestic

<table>
<thead>
<tr>
<th>Variable</th>
<th>Obs</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>mpg</td>
<td>52</td>
<td>21.2973</td>
<td>4.743297</td>
<td>13.47037</td>
<td>35.47038</td>
</tr>
</tbody>
</table>

-> foreign = Foreign

<table>
<thead>
<tr>
<th>Variable</th>
<th>Obs</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>mpg</td>
<td>22</td>
<td>21.2973</td>
<td>6.611187</td>
<td>10.52457</td>
<td>37.52457</td>
</tr>
</tbody>
</table>

Each sample (foreign and domestic) is a stratum, so the bootstrapped samples must have the same number of foreign and domestic cars as the original dataset. This requirement is facilitated by the strata() option to bootstrap. By typing the following, we bootstrap the test statistic using the modified dataset and save the values in bsauto2.dta:
. keep mpg foreign
. set seed 1
. bootstrap t=r(t), rep(1000) strata(foreign) saving(bsauto2) nodots: ttest mpg, by(foreign) unequal

Warning: Because ttest is not an estimation command or does not set e(sample), bootstrap has no way to determine which observations are used in calculating the statistics and so assumes that all observations are used. This means that no observations will be excluded from the resampling because of missing values or other reasons.

If the assumption is not true, press Break, save the data, and drop the observations that are to be excluded. Be sure that the dataset in memory contains only the relevant data.

Bootstrap results
Number of strata = 2 Number of obs = 74
Replications = 1,000
command: ttest mpg, by(foreign) unequal
t: r(t)

<table>
<thead>
<tr>
<th>Observed</th>
<th>Bootstrap</th>
<th>Normal-based</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coef.</td>
<td>Std. Err.</td>
<td>z</td>
</tr>
<tr>
<td>t</td>
<td>1.75e-07</td>
<td>1.051867</td>
</tr>
</tbody>
</table>

We can use the data in bsauto2.dta to estimate ASL via the fraction of bootstrap test statistics that are more extreme than 3.1797.

. use bsauto2, clear
(bootstrap: ttest)
. generate indicator = abs(t)>=abs(scalar(tobs))
. summarize indicator, meanonly
. display "ASLboot = " r(mean)
ASLboot = .004

The result is ASL_{boot} = 0.004. Assuming that the mean mpg is the same between foreign and domestic cars, we would expect to observe a t statistic more extreme (in absolute value) than 3.1797 in about 0.4% of all possible samples of the type we observed. This finding is still strong evidence to reject the hypothesis that the means are equal.

Bootstrapping a ratio

Example 4

Suppose that we wish to produce a bootstrap estimate of the ratio of two means. Because summarize stores results for only one variable, we must call summarize twice to compute the means. Actually, we could use collapse to compute the means in one call, but calling summarize twice is much faster. Thus, we will have to write a small program that will return the results we want.
We write the program below and save it to a file called `ratio.ado` (see [U] 17 Ado-files). Our program takes two variable names as input and saves them in the `local` macros `y` (first variable) and `x` (second variable). It then computes one statistic: the mean of ‘y’ divided by the mean of ‘x’. This value is returned as a scalar in `r(ratio)`. `ratio` also returns the ratio of the number of observations used to the mean for each variable.

```stata
program myratio, rclass
    version 16.1
    args y x
    confirm var `y'
    confirm var `x'
    tempname ymean yn
    summarize `y', meanonly
    scalar `ymean' = r(mean)
    return scalar n_`y' = r(N)
    summarize `x', meanonly
    return scalar n_`x' = r(N)
    return scalar ratio = `ymean'/r(mean)
end
```

Remember to test any newly written commands before using them with `bootstrap`.

```stata
. use https://www.stata-press.com/data/r16/auto, clear
   (1978 Automobile Data)
. summarize price

<table>
<thead>
<tr>
<th>Variable</th>
<th>Obs</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>price</td>
<td>74</td>
<td>6165.26</td>
<td>2949.496</td>
<td>3291</td>
<td>15906</td>
</tr>
</tbody>
</table>

. scalar mean1=r(mean)

. summarize weight

<table>
<thead>
<tr>
<th>Variable</th>
<th>Obs</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>weight</td>
<td>74</td>
<td>3019.46</td>
<td>777.1936</td>
<td>1760</td>
<td>4840</td>
</tr>
</tbody>
</table>

. scalar mean2=r(mean)

. di scalar(mean1)/scalar(mean2)
2.0418412

. myratio price weight
. return list
scalars:
   r(ratio) = 2.041841210168278
   r(n_weight) = 74
   r(n_price) = 74
```
The results of running `bootstrap` on our program are

```
  . use https://www.stata-press.com/data/r16/auto
     (1978 Automobile Data)
  . set seed 1
  . bootstrap ratio=r(ratio), reps(1000) nowarn nodots: myratio price weight
Bootstrap results                               Number of obs = 74
                          Replications = 1,000
command:  myratio price weight
         ratio:  r(ratio)
```

| Observed | Bootstrap Coef. | Bootstrap Std. Err. | Bootstrap z | P>|z| | Normal-based [95% Conf. Interval] |
|---------|----------------|--------------------|-------------|------|----------------------------------|
| ratio   | 2.041841       | 0.0953559          | 21.41       | 0.000 | 1.854947 2.228735               |

As mentioned previously, we should specify the `saving()` option if we wish to save the bootstrap dataset.

**Warning messages and e(sample)**

`bootstrap` is not meant to be used with weighted calculations. `bootstrap` determines the presence of weights by parsing the prefixed command with standard syntax. However, commands like `stcox` and `streg` require that weights be specified in `stset`, and some user commands may allow weights to be specified by using an option instead of the standard syntax. Both cases pose a problem for `bootstrap` because it cannot determine the presence of weights under these circumstances. In these cases, we can only assume that you know what you are doing.

`bootstrap` does not know which variables of the dataset in memory matter to the calculation at hand. You can speed their execution by dropping unnecessary variables because, otherwise, they are included in each bootstrap sample.

You should thus drop observations with missing values. Leaving in missing values causes no problem in one sense because all Stata commands deal with missing values gracefully. It does, however, cause a statistical problem. Bootstrap sampling is defined as drawing, with replacement, samples of size \( N \) from a set of \( N \) observations. `bootstrap` determines \( N \) by counting the number of observations in memory, not counting the number of nonmissing values on the relevant variables. The result is that too many observations are resampled; the resulting bootstrap samples, because they are drawn from a population with missing values, are of unequal sizes.

If the number of missing values relative to the sample size is small, this will make little difference. If you have many missing values, however, you should first drop the observations that contain them.

**Example 5**

To illustrate, we use the previous example but replace some of the values of `price` with missing values. The number of values of `price` used to compute the mean for each bootstrap is not constant. This is the purpose of the `Warning` message.
. use https://www.stata-press.com/data/r16/auto
(1978 Automobile Data)
. replace price = . if inlist(_n,1,3,5,7)
(4 real changes made, 4 to missing)
. set seed 1
. bootstrap ratio=r(ratio) np=r(n_price) nw=r(n_weight), reps(100) nodots:
> myratio price weight

Warning: Because myratio is not an estimation command or does not set
  e(sample), bootstrap has no way to determine which observations are
  used in calculating the statistics and so assumes that all
  observations are used. This means that no observations will be
  excluded from the resampling because of missing values or other
  reasons.
If the assumption is not true, press Break, save the data, and drop
the observations that are to be excluded. Be sure that the dataset
in memory contains only the relevant data.

Bootstrap results

| Command: myratio price weight | Number of obs = 74 |
| Replications = 100 |
| myratio price weight |
| ratio: r(ratio) |
| np: r(n_price) |
| nw: r(n_weight) |

| Observed | Bootstrap | Normal-based |
| Coef. | Std. Err. | z | P>|z| | [95% Conf. Interval] |
| --- | --- | --- | --- | --- | --- |
| ratio | 2.063051 | .0981706 | 21.01 | 0.000 | 1.870641 | 2.255462 |
| np | 70 | 2.071939 | 33.78 | 0.000 | 65.93908 | 74.06092 |
| nw | 74 | . | . | . | . | . |

Bootstrapping statistics from data with a complex structure

Here we describe how to bootstrap statistics from data with a complex structure, for example, longitudinal or panel data, or matched data. bootstrap, however, is not designed to work with complex survey data. It is important to include all necessary information about the structure of the data in the bootstrap syntax to obtain correct bootstrap estimates for standard errors and confidence intervals.

bootstrap offers several options identifying the specifics of the data. These options are strata(), cluster(), idcluster(), and group(). The usage of strata() was described in example 3 above. Below, we demonstrate several examples that require specifying the other three options.

Example 6

Suppose that auto.dta in example 1 above are clustered by rep78. We want to obtain bootstrap estimates for the standard errors of the difference between the coefficients on weight and gear_ratio, taking into account clustering.

We supply the cluster(rep78) option to bootstrap to request resampling from clusters rather than from observations in the dataset.
. use https://www.stata-press.com/data/r16/auto, clear
(1978 Automobile Data)
. keep if rep78<.
(5 observations deleted)
. bootstrap diff=(_b[weight]-_b[gear]), seed(1) cluster(rep78): regress mpg
> weight gear foreign
(running regress on estimation sample)
Bootstrap replications (50)

.................................................. 50
Linear regression Number of obs = 69
Replications = 50
command:  regress mpg weight gear foreign
diff:  _b[weight]-_b[gear]
(Replications based on 5 clusters in rep78)

<table>
<thead>
<tr>
<th></th>
<th>Observed</th>
<th>Bootstrap</th>
<th>Normal-based</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coef.</td>
<td>Std. Err.</td>
<td>z</td>
</tr>
<tr>
<td>diff</td>
<td>-1.910396</td>
<td>2.538558</td>
<td>-0.75</td>
</tr>
</tbody>
</table>

We drop missing values in rep78 before issuing the command because bootstrap does not allow missing values in cluster(). See the section above about using bootstrap when variables contain missing values.

We can also obtain these same results by using the following syntax:

. bootstrap diff=(_b[weight]-_b[gear]), seed(1): regress mpg weight gear foreign,
> vce(cluster rep78)

When only clustered information is provided to the command, bootstrap can pick up the vce(cluster clustvar) option from the main command and use it to resample from clusters.

Example 7

Suppose now that we have matched data and want to use bootstrap to obtain estimates of the standard errors of the exponentiated difference between two coefficients (or, equivalently, the ratio of two odds ratios) estimated by clogit. Consider the example of matched case–control data on birthweight of infants described in example 2 of [R] clogit.

The infants are paired by being matched on mother’s age. All groups, defined by the pairid variable, have 1:1 matching. clogit requires that the matching information, pairid, be supplied to the group() (or, equivalently, strata()) option to be used in computing the parameter estimates. Because the data are matched, we need to resample from groups rather than from the whole dataset. However, simply supplying the grouping variable pairid in cluster() is not enough with bootstrap, as it is with clustered data.
. use https://www.stata-press.com/data/r16/lowbirth2, clear
(Applied Logistic Regression, Hosmer & Lemeshow)
. bootstrap ratio=exp(_b[smoke]-_b[ptd]), seed(1) cluster(pairid): clogit low
> lwt smoke ptd ht ui i.race, group(pairid)
(running *clogit* on estimation sample)

Bootstrap replications (50)

................................. 50

Bootstrap results

<table>
<thead>
<tr>
<th>command:</th>
<th>Observed</th>
<th>Bootstrap</th>
<th>Normal-based</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coef.</td>
<td>Std. Err.</td>
<td>z  P&gt;</td>
</tr>
<tr>
<td>ratio</td>
<td>.6654095</td>
<td>2.043274</td>
<td>0.33 0.745 -3.339334 4.670153</td>
</tr>
</tbody>
</table>

For the syntax above, imagine that the first pair was sampled twice during a replication. Then, the bootstrap sample has four subjects with *pairid* equal to one, which clearly violates the original 1:1 matching design. As a result, the estimates of the coefficients obtained from this bootstrap sample will be incorrect.

Therefore, in addition to resampling from groups, we need to ensure that resampled groups are uniquely identified in each of the bootstrap samples. The *idcluster*(newcluster) option is designed for this. It requests that at each replication *bootstrap* create the new variable, *newcluster*, containing unique identifiers for all resampled groups. Thus, to make sure that the correct matching is preserved during each replication, we need to specify the grouping variable in *cluster()*, supply a variable name to *idcluster()*, and use this variable as the grouping variable with *clogit*, as we demonstrate below.

. bootstrap ratio=exp(_b[smoke]-_b[ptd]), seed(1) cluster(pairid)
> idcluster(newpairid): clogit low lwt smoke ptd ht ui i.race, group(newpairid)
(running *clogit* on estimation sample)

Bootstrap replications (50)

................................. 50

Bootstrap results

<table>
<thead>
<tr>
<th>command:</th>
<th>Observed</th>
<th>Bootstrap</th>
<th>Normal-based</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coef.</td>
<td>Std. Err.</td>
<td>z  P&gt;</td>
</tr>
<tr>
<td>ratio</td>
<td>.6654095</td>
<td>1.156848</td>
<td>0.58 0.565 -1.601972 2.932791</td>
</tr>
</tbody>
</table>

Note the difference between the estimates of the bootstrap standard error for the two specifications of the *bootstrap* syntax.
Similarly, when you have panel (longitudinal) data, all resampled panels must be unique in each of the bootstrap samples to obtain correct bootstrap estimates of statistics. Therefore, both `cluster(panelvar)` and `idcluster(newpanelvar)` must be specified with `bootstrap`, and `i(newpanelvar)` must be used with the main command. Moreover, you must clear the current `xtset` settings by typing `xtset, clear` before calling `bootstrap`.

### Example 8

Continuing with our birthweight data, suppose that we have more information about doctors supervising women’s pregnancies. We believe that the data on the pairs of infants from the same doctor may be correlated and want to adjust standard errors for possible correlation among the pairs. `clogit` offers the `vce(cluster clustvar)` option to do this.

Let’s add a cluster variable to our dataset. One thing to keep in mind is that to use `vce(cluster clustvar)`, groups in `group()` must be nested within clusters.

```
use https://www.stata-press.com/data/r16/lowbirth2, clear
(Applied Logistic Regression, Hosmer & Lemeshow)
.set seed 12345
.by pairid, sort: egen byte doctor = total(int(2*runiform()+1)*(_n == 1))
clogit low lwt smoke ptd ht ui i.race, group(pairid) vce(cluster doctor)
Iteration 0: log pseudolikelihood = -26.768693
Iteration 1: log pseudolikelihood = -25.810476
Iteration 2: log pseudolikelihood = -25.794296
Iteration 3: log pseudolikelihood = -25.794271
Iteration 4: log pseudolikelihood = -25.794271
Conditional (fixed-effects) logistic regression
Number of obs = 112
Wald chi2(1) = .
Prob > chi2 = .
Log pseudolikelihood = -25.794271 Pseudo R2 = 0.3355
(Std. Err. adjusted for 2 clusters in doctor)
```

| low  | Coef.  | Std. Err. | z     | P>|z| | [95% Conf. Interval] |
|------|--------|-----------|-------|-----|----------------------|
| lwt  | -.0183757 | .0020314 | -9.05 | 0.000 | -.0223571 | -.0143942 |
| smoke| 1.400656  | .3067525 | 4.57  | 0.000 | .7994322  | 2.00188  |
| ptd  | 1.808009  | .2092246 | 8.64  | 0.000 | 1.397936  | 2.218082 |
| ht   | 2.361152  | 1.410341 | 1.67  | 0.094 | -.4030665 | 5.12537  |
| ui   | 1.401929  | .9406248 | 1.49  | 0.136 | -.4416617 | 3.24552  |
| race | black    | .5713643 | .999265 | 0.57 | 0.567 | -1.38716 | 2.529889 |
|      | other    | -.0253148 | .8453206 | -0.03 | 0.976 | -1.682113 | 1.631483 |

To obtain correct bootstrap standard errors of the exponentiated difference between the two coefficients in this example, we need to make sure that both resampled clusters and groups within resampled clusters are unique in each of the bootstrap samples. To achieve this, `bootstrap` needs the information about clusters in `cluster()`, the variable name of the new identifier for clusters in `idcluster()`, and the information about groups in `group()`. We demonstrate the corresponding syntax of `bootstrap` below.
. bootstrap ratio=exp(_b[smoke]-_b[ptd]), seed(1) cluster(doctor)
> idcluster(uidoctor) group(pairid): clogit low lwt smoke ptd ht ui i.race,
> group(pairid)
(running clogit on estimation sample)
Bootstrap replications (50)
.................................................. 50
Bootstrap results

command: clogit low lwt smoke ptd ht ui i.race, group(pairid)
ratio: exp(_b[smoke]-_b[ptd])

(Replications based on 2 clusters in doctor)

<table>
<thead>
<tr>
<th></th>
<th>Observed</th>
<th>Bootstrap</th>
<th>Normal-based</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coef.</td>
<td>Std. Err.</td>
<td>z</td>
</tr>
<tr>
<td>ratio</td>
<td>.6654095</td>
<td>.1459234</td>
<td>4.56</td>
</tr>
</tbody>
</table>

In the above syntax, although we specify `group(pairid)` with `clogit`, it is not the group identifiers of the original `pairid` variable that are used to compute parameter estimates from bootstrap samples. The way `bootstrap` works is that, at each replication, the clusters defined by `doctor` are resampled and the new variable, `uidoctor`, uniquely identifying resampled clusters is created. After that, another new variable uniquely identifying the `(uidoctor, group)` combination is created and renamed to have the same name as the grouping variable, `pairid`. This newly defined grouping variable is then used by `clogit` to obtain the parameter estimates from this bootstrap sample of clusters. After all replications are performed, the original values of the grouping variable are restored.

Technical note

The same logic must be used when running `bootstrap` with commands designed for panel (longitudinal) data that allow specifying the `cluster(clustervar)` option. To ensure that the combination of `(clustervar, panelvar)` values are unique in each of the bootstrap samples, `cluster(clustervar)`, `id-cluster(newclustervar)`, and `group(panelvar)` must be specified with `bootstrap`, and `i(panelvar)` must be used with the main command.

Bradley Efron (1938– ) was born in 1938 in Minnesota and studied mathematics and statistics at Caltech and Stanford; he has lived in northern California since 1960. He has worked on empirical Bayes, survival analysis, exponential families, bootstrap and jackknife methods, and confidence intervals, in conjunction with applied work in biostatistics, astronomy, and physics.

Efron is a member of the U.S. National Academy of Sciences and was awarded the U.S. National Medal of Science in 2005. He is by any standards one of the world’s leading statisticians: his work ranges from deep and elegant contributions in theoretical statistics to pathbreaking involvement in a variety of practical applications.
Stored results

bootstrap stores the following in `e()`:  

Scalars
- `e(N)` sample size
- `e(N_reps)` number of complete replications
- `e(N_misreps)` number of incomplete replications
- `e(N_strata)` number of strata
- `e(N_clust)` number of clusters
- `e(k_eq)` number of equations in `e(b)`
- `e(k_exp)` number of standard expressions
- `e(k_eexp)` number of extended expressions (i.e., `_b`)
- `e(k_extra)` number of extra equations beyond the original ones from `e(b)`
- `e(level)` confidence level for bootstrap CIs
- `e(bs_version)` version for bootstrap results
- `e(rank)` rank of `e(V)`

Macros
- `e(cmdname)` command name from `command`
- `e(cmd)` same as `e(cmdname)` or `bootstrap`
- `e(command)` command as typed
- `e(cmdline)` command as typed
- `e(prefix)` `bootstrap`
- `e(title)` title in estimation output
- `e(strata)` strata variables
- `e(cluster)` cluster variables
- `e(rngstate)` random-number state used
- `e(size)` from the `size(#)` option
- `e(exp#)` expression for the `#th` statistic
- `e(ties)` ties, if specified
- `e(mse)` `mse`, if specified
- `e(vce)` `bootstrap`
- `e(vcetype)` title used to label Std. Err.
- `e(properties)` `b V`

Matrices
- `e(b)` observed statistics
- `e(b_bs)` bootstrap estimates
- `e(reps)` number of nonmissing results
- `e(bias)` estimated biases
- `e(se)` estimated standard errors
- `e(z0)` median biases
- `e(accel)` estimated accelerations
- `e(ci_normal)` normal-approximation CIs
- `e(ci_percentile)` percentile CIs
- `e(ci_bc)` bias-corrected CIs
- `e(ci_bca)` bias-corrected and accelerated CIs
- `e(V)` bootstrap variance–covariance matrix
- `e(V_modelbased)` model-based variance

When `exp_list` is `_b`, bootstrap will also carry forward most of the results already in `e()` from `command`.

Methods and formulas

Let $\hat{\theta}$ be the observed value of the statistic, that is, the value of the statistic calculated with the original dataset. Let $i = 1, 2, \ldots, k$ denote the bootstrap samples, and let $\hat{\theta}_i$ be the value of the statistic from the $i$th bootstrap sample.
When the `mse` option is specified, the standard error is estimated as

\[
\hat{se}_{\text{MSE}} = \left\{ \frac{1}{k} \sum_{i=1}^{k} (\hat{\theta}_i - \bar{\theta})^2 \right\}^{1/2}
\]

Otherwise, the standard error is estimated as

\[
\hat{se} = \left\{ \frac{1}{k-1} \sum_{i=1}^{k} (\hat{\theta}_i - \bar{\theta})^2 \right\}^{1/2}
\]

where

\[
\bar{\theta} = \frac{1}{k} \sum_{i=1}^{k} \hat{\theta}_i
\]

The variance–covariance matrix is similarly computed. The bias is estimated as

\[
\hat{\text{bias}} = \bar{\theta} - \hat{\theta}
\]

Confidence intervals with nominal coverage rates 1 - \(\alpha\) are calculated according to the following formulas. The normal-approximation method yields the confidence intervals

\[
[\hat{\theta} - z_{1-\alpha/2} \hat{se}, \hat{\theta} + z_{1-\alpha/2} \hat{se}]
\]

where \(z_{1-\alpha/2}\) is the \((1 - \alpha/2)\)th quantile of the standard normal distribution. If the `mse` option is specified, `bootstrap` will report the normal confidence interval using \(\hat{se}_{\text{MSE}}\) instead of \(\hat{se}\). `estat bootstrap` only uses \(\hat{se}\) in the normal confidence interval.

The percentile method yields the confidence intervals

\[
[\theta^*_p, \theta^*_{1-\alpha/2}]
\]

where \(\theta^*_p\) is the \(p\)th quantile (the 100\(p\)th percentile) of the bootstrap distribution \((\hat{\theta}_1, \ldots, \hat{\theta}_k)\).

Let

\[
z_0 = \Phi^{-1}\left\{ \#(\hat{\theta}_i \leq \hat{\theta}) / k \right\}
\]

where \(\#(\hat{\theta}_i \leq \hat{\theta})\) is the number of elements of the bootstrap distribution that are less than or equal to the observed statistic and \(\Phi\) is the standard cumulative normal. \(z_0\) is known as the median bias of \(\hat{\theta}\). When the `ties` option is specified, \(z_0\) is estimated as \(\#(\hat{\theta}_i < \hat{\theta}) + \#(\hat{\theta}_i = \hat{\theta}) / 2\), which is the number of elements of the bootstrap distribution that are less than the observed statistic plus half the number of elements that are equal to the observed statistic.

Let

\[
a = \frac{\sum_{i=1}^{n} (\bar{\theta}_{(\cdot)} - \hat{\theta}_{(i)})^3}{6 \left\{ \sum_{i=1}^{n} (\bar{\theta}_{(\cdot)} - \hat{\theta}_{(i)})^2 \right\}^{3/2}}
\]

where \(\hat{\theta}_{(i)}\) are the leave-one-out (jackknife) estimates of \(\hat{\theta}\) and \(\bar{\theta}_{(\cdot)}\) is their mean. This expression is known as the jackknife estimate of acceleration for \(\hat{\theta}\). Let

\[
p_1 = \Phi\left\{ z_0 + \frac{z_0 - z_{1-\alpha/2}}{1 - a(z_0 - z_{1-\alpha/2})} \right\}
\]

\[
p_2 = \Phi\left\{ z_0 + \frac{z_0 + z_{1-\alpha/2}}{1 - a(z_0 + z_{1-\alpha/2})} \right\}
\]
where $z_{1-\alpha/2}$ is the $(1-\alpha/2)$th quantile of the normal distribution. The bias-corrected and accelerated (BC$_a$) method yields confidence intervals

$$
\left[ \theta^*_p, \theta^*_p \right]
$$

where $\theta^*_p$ is the $p$th quantile of the bootstrap distribution as defined previously. The bias-corrected (but not accelerated) method is a special case of BC$_a$ with $a = 0$.

References


Also see

[R] bootstrap postestimation — Postestimation tools for bootstrap
[R] jackknife — Jackknife estimation
[R] permute — Monte Carlo permutation tests
[R] simulate — Monte Carlo simulations
[R] set rngstream — Specify the stream for the stream random-number generator
[SVY] svy bootstrap — Bootstrap for survey data
[U] 13.5 Accessing coefficients and standard errors
[U] 13.6 Accessing results from Stata commands
[U] 20 Estimation and postestimation commands