

# **MATA REFERENCE MANUAL**

## **RELEASE 12**



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## Cross-referencing the documentation

When reading this manual, you will find references to other Stata manuals. For example,

[U] **26 Overview of Stata estimation commands**

[R] **regress**

[D] **reshape**

The first example is a reference to chapter 26, *Overview of Stata estimation commands*, in the *User's Guide*; the second is a reference to the `regress` entry in the *Base Reference Manual*; and the third is a reference to the `reshape` entry in the *Data-Management Reference Manual*.

All the manuals in the Stata Documentation have a shorthand notation:

[GSM]	<i>Getting Started with Stata for Mac</i>
[GSU]	<i>Getting Started with Stata for Unix</i>
[GSW]	<i>Getting Started with Stata for Windows</i>
[U]	<i>Stata User's Guide</i>
[R]	<i>Stata Base Reference Manual</i>
[D]	<i>Stata Data-Management Reference Manual</i>
[G]	<i>Stata Graphics Reference Manual</i>
[XT]	<i>Stata Longitudinal-Data/Panel-Data Reference Manual</i>
[MI]	<i>Stata Multiple-Imputation Reference Manual</i>
[MV]	<i>Stata Multivariate Statistics Reference Manual</i>
[P]	<i>Stata Programming Reference Manual</i>
[SEM]	<i>Stata Structural Equation Modeling Reference Manual</i>
[SVY]	<i>Stata Survey Data Reference Manual</i>
[ST]	<i>Stata Survival Analysis and Epidemiological Tables Reference Manual</i>
[TS]	<i>Stata Time-Series Reference Manual</i>
[I]	<i>Stata Quick Reference and Index</i>
[M]	<i>Mata Reference Manual</i>

Detailed information about each of these manuals may be found online at

<http://www.stata-press.com/manuals/>



# **[M-0] Introduction to the Mata manual**





# Title

[M-0] intro — Introduction to the Mata manual

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<a href="#">[M-1]</a>	<b>Introduction and advice</b>
<a href="#">[M-2]</a>	<b>Language definition</b>
<a href="#">[M-3]</a>	<b>Commands for controlling Mata</b>
<a href="#">[M-4]</a>	<b>Index and guide to functions</b>
<a href="#">[M-5]</a>	<b>Functions</b>
<a href="#">[M-6]</a>	<b>Mata glossary of common terms</b>

## Description

Mata is a matrix programming language that can be used by those who want to perform matrix calculations interactively and by those who want to add new features to Stata.

This entry describes this manual and what has changed since Stata 11.

## Remarks

This manual is divided into six sections. Each section is organized alphabetically, but there is an introduction in front that will help you get around.

If you are new to Mata, here is a helpful reading list. Start by reading

<a href="#">[M-1] first</a>	Introduction and first session
<a href="#">[M-1] interactive</a>	Using Mata interactively
<a href="#">[M-1] how</a>	How Mata works

You may find other things in section [M-1] that interest you. For a table of contents, see

<a href="#">[M-1] intro</a>	Introduction and advice
-----------------------------	-------------------------

Whenever you see a term that you are unfamiliar with, see

<a href="#">[M-6] Glossary</a>	Mata glossary of common terms
--------------------------------	-------------------------------

Now that you know the basics, if you are interested, you can look deeper into Mata’s programming features:

[M-2] <a href="#">syntax</a>	Mata language grammar and syntax
------------------------------	----------------------------------

[M-2] [syntax](#) is pretty dense reading, but it summarizes nearly everything. The other entries in [M-2] repeat what is said there but with more explanation; see

[M-2] <a href="#">intro</a>	Language definition
-----------------------------	---------------------

because other entries in [M-2] will interest you. If you are interested in object-oriented programming, be sure to see [M-2] [class](#).

Along the way, you will eventually be guided to sections [M-4] and [M-5]. [M-5] documents Mata’s functions; the alphabetical order makes it easy to find a function if you know its name but makes learning what functions there are hopeless. That is the purpose of [M-4]—to present the functions in logical order. See

[M-4] <a href="#">intro</a>	Index and guide to functions
-----------------------------	------------------------------

Mathematical

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[M-4] <a href="#">solvers</a>	Matrix solvers and inverters
[M-4] <a href="#">scalar</a>	Scalar functions
[M-4] <a href="#">statistical</a>	Statistical functions
[M-4] <a href="#">mathematical</a>	Other important functions

Utility and manipulation

[M-4] <a href="#">standard</a>	Functions to create standard matrices
[M-4] <a href="#">utility</a>	Matrix utility functions
[M-4] <a href="#">manipulation</a>	Matrix manipulation functions

Stata interface

[M-4] <a href="#">stata</a>	Stata interface functions
-----------------------------	---------------------------

String, I/O, and programming

[M-4] <a href="#">string</a>	String manipulation functions
[M-4] <a href="#">io</a>	I/O functions
[M-4] <a href="#">programming</a>	Programming functions

## What's new

This section is intended for previous Stata users. If you are new to Stata, you may as well skip it.

1. **New Stata commands `getmata` and `putmata`** make it easy to transfer your data into Mata, manipulate them, and then transfer them back to Stata. `getmata` and `putmata` are especially designed for interactive use. See [\[D\] `putmata`](#).
  2. **New functions** imported from Stata,
    - a. **Tukey's Studentized range**, cumulative and inverse, `tukeyprob()` and `invtukeyprob()`.
    - b. **Dunnett's multiple range**, cumulative and inverse, `dunnettprob()` and `invdunnettprob()`.
    - c. **New date conversion functions** `dofb()` and `bofd()` convert between business dates and standard calendar dates. See [\[D\] `datetime business calendars`](#).
- See [\[D\] `functions`](#), [\[M-5\] `normal\(\)`](#), and [\[M-5\] `date\(\)`](#).
3. **Support for hidden and historical saved results.** Existing Mata functions `st_global()`, `st_numscalar()`, and `st_matrix()` now allow an optional third argument specifying the hidden or historical status. Three new functions—`st_global_hcat()`, `st_numscalar_hcat()`, `st_matrix_hcat()`—allow you to determine the saved hidden or historical status. See [\[M-5\] `st\_global\(\)`](#), [\[M-5\] `st\_numscalar\(\)`](#), and [\[M-5\] `st\_matrix\(\)`](#).
  4. **Support for new ml features.** Stata's `ml` now distinguishes the Hessian matrix produced by `technique(nr)` from the other techniques that compute a substitute for the Hessian matrix. This means that `ml` will compute the real Hessian matrix of second derivatives to determine convergence when all other convergence tolerances are satisfied and `technique(bfgs)`, `technique(bhhh)`, or `technique(dfpr)` is in effect.

Mata's commands `optimize()` and `moptimize()` have been similarly changed. See [\[M-5\] `optimize\(\)`](#) and [\[M-5\] `moptimize\(\)`](#).

## Also see

[\[M-1\] `first`](#) — Introduction and first session

[\[M-6\] `Glossary`](#)



## **[M-1] Introduction and advice**



# Title

[M-1] intro — Introduction and advice

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<a href="#">limits</a>	Limits and memory utilization
<a href="#">tolerance</a>	Use and specification of tolerances
<a href="#">permutation</a>	An aside on permutation matrices and vectors
<a href="#">LAPACK</a>	The LAPACK linear-algebra routines

## Description

This section provides an introduction to Mata along with reference material common to all sections.

## Remarks

The most important entry in this section is [\[M-1\] first](#). Also see [\[M-6\] Glossary](#).

The Stata commands `putmata` and `getmata` are useful for moving data from Stata to Mata and back again; see [\[D\] putmata](#).

Those looking for a textbook-like introduction to Mata may want to consider [Baum \(2009\)](#), particularly chapters 13 and 14.

## Reference

Baum, C. F. 2009. *An Introduction to Stata Programming*. College Station, TX: Stata Press.

## Also see

[M-0] [intro](#) — Introduction to the Mata manual

[D] [putmata](#) — Put Stata variables into Mata and vice versa



# Title

[M-1] ado — Using Mata with ado-files

## Description

This section provides advice to ado-file programmers on how to use Mata effectively and efficiently.

## Remarks

For those interested in extending Stata by adding new commands, Mata is not a replacement for ado-files. Rather, the appropriate way to use Mata is to create subroutines used by your ado-files. Below we discuss how to do that under the following headings:

[A first example](#)  
[Where to store the Mata functions](#)  
[Passing arguments to Mata functions](#)  
[Returning results to ado-code](#)  
[Advice: Use of matastrict](#)  
[Advice: Some useful Mata functions](#)

## A first example

We will pretend that Stata cannot produce sums and that we want to write a new command for Stata that will report the sum of one variable. Here is our first take on how we might do this:

```
----- begin varsum.ado -----  
  
program varsum  
    version 12  
    syntax varname [if] [in]  
    marksample touse  
    mata: calcsum("`varlist'", "`touse'")  
    display as txt "    sum = " as res r(sum)  
end  
  
version 12  
mata:  
void calcsum(string scalar varname, string scalar touse)  
{  
    real colvector  x  
    st_view(x, ., varname, touse)  
    st_numscalar("r(sum)", colsum(x))  
}  
end  
  
----- end varsum.ado -----
```

Running this program from Stata results in the following output:

```
. varsum mpg  
    sum = 1576
```

Note the following:

1. The ado-file has both ado-code and Mata code in it.

2. The ado-code handled all issues of parsing and identifying the subsample of the data to be used.
3. The ado-code called a Mata function to perform the calculation.
4. The Mata function received as arguments the names of two variables in the Stata dataset: the variable on which the calculation was to be made and the variable that identified the subsample of the data to be used.
5. The Mata function returned the result in `r()`, from where the ado-code could access it.

The outline that we showed above is a good one, although we will show you alternatives that, for some problems, are better.

## Where to store the Mata functions

Our ado-file included a Mata function. You have three choices of where to put the Mata function:

1. You can put the code for the Mata function in the ado-file, as we did. To work, your .ado file must be placed where Stata can find it.
2. You can omit code for the Mata function from the ado-file, compile the Mata function separately, and store the compiled code (the object code) in a separate file called a .mo file. You place both your .ado and .mo files where Stata can find them.
3. You can omit the code for the Mata function from the ado-file, compile the Mata function separately, and store the compiled code in a .mlib library. Here both your .ado file and your .mlib library will need to be placed where Stata can find them.

We will show you how to do each of these alternatives, but before we do, let's consider the advantages and disadvantages of each:

1. Putting your Mata source code right in the ado-file is easiest, and it sure is convenient. The disadvantage is that Mata must compile the source code into object code, and that will slow execution. The cost is small because it occurs infrequently; Mata compiles the code when the ado-file is loaded and, as long as the ado-file is not dropped from memory, Stata and Mata will use the same compiled code over and over again.
2. Saving your Mata code as .mo files saves Mata from having to compile them at all. The source code is compiled only once—at the time you create the .mo file—and thereafter, it is the already-compiled copy that Stata and Mata will use.

There is a second advantage: rather than the Mata functions (`calcsun()` here) being private to the ado-file, you will be able to use `calcsun()` in your other ado-files. `calcsun()` will become a utility always available to you. Perhaps `calcsun()` is not deserving of this honor.

3. Saving your Mata code in a .mlib library has the same advantages and disadvantages as (2); the only difference is that we save the precompiled code in a different way. The .mo/.mlib choice is of more concern to those who intend to distribute their ado-file to others. .mlib libraries allow you to place many Mata functions (subroutines for your ado-files) into one file, and so it is easier to distribute.

Let's restructure our ado-file to save `calcsun()` in a `.mo` file. First, we simply remove `calcsun()` from our ado-file, so it now reads

```

----- begin varsum.ado -----
program varsum
    version 12
    syntax varname [if] [in]
    marksample touse
    mata: calcsun("`varlist'", "`touse'")
    display as txt "    sum = " as res r(sum)
end
----- end varsum.ado -----

```

Next, we enter Mata, enter our `calcsun()` program, and save the object code:

```

: void calcsun(string scalar varname, string scalar touse)
> {
>     real colvector  x
>
>     st_view(x, ., varname, touse)
>     st_numscalar("r(sum)", colsum(x))
> }
: mata mosave calcsun(), dir(PERSONAL)

```

This step we do only once. The `mata mosave` command created file `calcsun.mo` stored in our `PERSONAL sysdir` directory; see [M-3] [mata mosave](#) and [P] [sysdir](#) for more details. We put the `calcsun.mo` file in our `PERSONAL` directory so that Stata and Mata would be able to find it, just as you probably did with the `varsum.ado` ado-file.

Typing in the `calcsun()` program interactively is too prone to error, so instead what we can do is create a do-file to perform the step and then run the do-file once:

```

----- begin varsum.do -----
version 12
mata:
void calcsun(string scalar varname, string scalar touse)
{
    real colvector  x
    st_view(x, ., varname, touse)
    st_numscalar("r(sum)", colsum(x))
}
mata mosave calcsun(), dir(PERSONAL) replace
end
----- end varsum.do -----

```

We save the do-file someplace safe in case we should need to modify our code later, either to fix bugs or to add features. For the same reason, we added a `replace` option on the command that creates the `.mo` file, so we can run the do-file over and over.

To follow the third organization—putting our `calcsun()` subroutine in a library—is just a matter of modifying `varsum.do` to do `mata mlib add` rather than `mata mosave`. See [M-3] [mata mlib](#); there are important details having to do with how you will manage all the different functions you will put in the library, but that has nothing to do with our problem here.

Where and how you store your Mata subroutines has nothing to do with what your Mata subroutines do or how you use them.

## Passing arguments to Mata functions

In designing a subroutine, you have two paramount interests: how you get data into your subroutine and how you get results back. You get data in by the values you pass as the arguments. For `calcsun()`, we called the subroutine by coding

```
mata: calcsun("`varlist'", "`touse'")
```

After macro expansion, that line turned into something like

```
mata: calcsun("mpg", "__0001dc")
```

The `__0001dc` variable is a temporary variable created by the `marksample` command earlier in our ado-file. `mpg` was the variable specified by the user. After expansion, the arguments were nothing more than strings, and those strings were passed to `calcsun()`.

Macro substitution is the most common way values are passed to Mata subroutines. If we had a Mata function `add(a, b)`, which could add numbers, we might code in our ado-file

```
mata: add(`x', `y')
```

and, if macro `'x'` contained 2 and macro `'y'` contained 3, Mata would see

```
mata: add(2, 3)
```

and values 2 and 3 would be passed to the subroutine.

When you think about writing your Mata subroutine, the arguments your ado-file will find convenient to pass and Mata will make convenient to use are

1. numbers, which Mata calls real scalars, such as 2 and 3 (`'x'` and `'y'`), and
2. names of variables, macros, scalars, matrices, etc., which Mata calls string scalars, such as `"mpg"` and `"__0001dc"` (`"`varlist'"` and `"`touse'"`).

To receive arguments of type (1), you code `real scalar` as the type of the argument in the function declaration and then use the real scalar variable in your Mata code.

To receive arguments of type (2), you code `string scalar` as the type of the argument in the function declaration, and then you use one of the Stata interface functions (in [M-4] `stata`) to go from the name to the contents. If you receive a variable name, you will especially want to read about the functions `st_data()` and `st_view()` (see [M-5] `st_data()` and [M-5] `st_view()`), although there are many other utilities for dealing with variable names. If you are dealing with local or global macros, scalars, or matrices, you will want to see [M-5] `st_local()`, [M-5] `st_global()`, [M-5] `st_numscalar()`, and [M-5] `st_matrix()`.

## Returning results to ado-code

You have many more choices on how to return results from your Mata function to the calling ado-code.

1. You can return results in `r()`—as we did in our example—or in `e()` or in `s()`.
2. You can return results in macros, scalars, matrices, etc., whose names are passed to your Mata subroutine as arguments.
3. You can highhandedly reach back into the calling ado-file and return results in macros, scalars, matrices, etc., whose names are of your devising.

In all cases, see [M-5] `st_global()`. `st_global()` is probably not the function you will use, but there is a wonderfully useful table in the *Remarks* section that will tell you exactly which function to use.

Also see all other Stata interface functions in [M-4] `stata`.

If you want to modify the Stata dataset in memory, see [M-5] `st_store()` and [M-5] `st_view()`.

## Advice: Use of matastrict

The setting `matastrict` determines whether declarations can be omitted (see [M-2] *declarations*); by default, you may. That is, `matastrict` is set off, but you can turn it on by typing `mata set matastrict on`; see [M-3] *mata set*. Some users do, and some users do not.

So now, consider what happens when you include Mata source code directly in the ado-file. When the ado-file is loaded, is `matastrict` set on, or is it set off? The answer is that it is off, because when you include the Mata source code inside an ado-file, `matastrict` is temporarily switched off when the ado-file is loaded even if the user running the ado-file has previously set it on.

For example, `varsum.ado` could read

```

----- begin varsum.ado -----
program varsum
    version 12
    syntax varname [if] [in]
    marksample touse
    mata: calcsun("varlist", "touse")
    display as txt "    sum = " as res r(sum)
end

version 12
mata:
void calcsun(varname, touse)
{
    st_view(x, ., varname, touse)           // (note absence of declarations)
    st_numscalar("r(sum)", colsum(x))
}
end
----- end varsum.ado -----

```

and it will work even when run by users who have set `matastrict` on.

Similarly, in an ado-file, you can set `matastrict` on and that will not affect the setting after the ado-file is loaded, so `varsum.ado` could read

---

 begin varsum.ado
 

---

```

program varsum
    version 12
    syntax varname [if] [in]
    marksample touse
    mata: calcsun("`varlist'", "`touse'")
    display as txt "    sum = " as res r(sum)
end
version 12
mata:
mata set matastrict on
void calcsun(string scalar varname, string scalar touse)
{
    real colvector  x
    st_view(x, ., varname, touse)
    st_numscalar("r(sum)", colsum(x))
}
end

```

---

 end varsum.ado
 

---

and not only will it work, but running `varsum` will not change the user's `matastrict` setting.

This preserving and restoring of `matastrict` is something that is done only for ado-files when they are loaded.

## Advice: Some useful Mata functions

In the `calcsun()` subroutine, we used the `colsum()` function—see [M-5] [sum\(\)](#)—to obtain the sum:

```

void calcsun(string scalar varname, string scalar touse)
{
    real colvector  x
    st_view(x, ., varname, touse)
    st_numscalar("r(sum)", colsum(x))
}

```

We could instead have coded

```

void calcsun(string scalar varname, string scalar touse)
{
    real colvector  x
    real scalar     i, sum
    st_view(x, ., varname, touse)
    sum = 0
    for (i=1; i<=rows(x); i++) sum = sum + x[i]
    st_numscalar("r(sum)", sum)
}

```

The first way is preferred. Rather than construct explicit loops, it is better to call functions that calculate the desired result when such functions exist. Unlike ado-code, however, when such functions do not exist, you can code explicit loops and still obtain good performance.

Another set of functions we recommend are documented in [M-5] [cross\(\)](#), [M-5] [crossdev\(\)](#), and [M-5] [quadcross\(\)](#).

`cross()` makes calculations of the form

$$\begin{aligned} X'X \\ X'Z \\ X'\text{diag}(w)X \\ X'\text{diag}(w)Z \end{aligned}$$

`crossdev()` makes calculations of the form

$$\begin{aligned} (X:-x)'(X:-x) \\ (X:-x)'(Z:-z) \\ (X:-x)'\text{diag}(w)(X:-x) \\ (X:-x)'\text{diag}(w)(Z:-z) \end{aligned}$$

Both these functions could easily escape your attention because the matrix expressions themselves are so easily written in Mata. The functions, however, are quicker, use less memory, and sometimes are more accurate. Also, quad-precision versions of the functions exist; [M-5] [quadcross\(\)](#).

## Also see

[M-2] [version](#) — Version control

[M-1] [intro](#) — Introduction and advice

# Title

[M-1] first — Introduction and first session

## Description

Mata is a component of Stata. It is a matrix programming language that can be used interactively or as an extension for do-files and ado-files. Thus

1. Mata can be used by users who want to think in matrix terms and perform (not necessarily simple) matrix calculations interactively, and
2. Mata can be used by advanced Stata programmers who want to add features to Stata.

Mata has something for everybody.

Primary features of Mata are that it is fast and that it is C-like.

## Remarks

This introduction is presented under the following headings:

*Invoking Mata*  
*Using Mata*  
*Making mistakes: Interpreting error messages*  
*Working with real numbers, complex numbers, and strings*  
*Working with scalars, vectors, and matrices*  
*Working with functions*  
*Distinguishing real and complex values*  
*Working with matrix and scalar functions*  
*Performing element-by-element calculations: Colon operators*  
*Writing programs*  
*More functions*  
*Mata environment commands*  
*Exiting Mata*

If you are reading the entries in the order suggested in [M-0] **intro**, see [M-1] **interactive** next.

## Invoking Mata

To enter Mata, type `mata` at Stata's dot prompt and press enter; to exit Mata, type `end` at Mata's colon prompt:

<code>. mata</code>	← type <code>mata</code> to enter Mata
<code>_____ mata (type <b>end</b> to exit) _____</code>	
<code>: 2 + 2</code>	← type Mata statements at the colon prompt
<code>  4</code>	
<code>: end</code>	← type <code>end</code> to return to Stata
<code>_____</code>	
<code>. _</code>	← you are back to Stata



## Using Mata

When you type a statement into Mata, Mata compiles what you typed and, if it compiled without error, executes it:

```
: 2 + 2
      4
: _
```

We typed  $2 + 2$ , a particular example from the general class of expressions. Mata responded with 4, the evaluation of the expression.

Often what you type are expressions, although you will probably choose more complicated examples. When an expression is not assigned to a variable, the result of the expression is displayed. Assignment is performed by the `=` operator:

```
: x = 2 + 2
: x
      4
: _
```

When we type “ $x = 2 + 2$ ”, the expression is evaluated and stored in the variable we just named `x`. The result is not displayed. We can look at the contents of `x`, however, simply by typing “`x`”. From Mata’s perspective, `x` is not only a variable but also an expression, albeit a rather simple one. Just as  $2 + 2$  says to load 2, load another 2, and add them, the expression `x` says to load `x` and stop there.

As an aside, Mata distinguishes uppercase and lowercase. `X` is not the same as `x`:

```
: X = 2 + 3
: x
      4
: X
      5
: _
```

## Making mistakes: Interpreting error messages

If you make a mistake, Mata complains, and then you continue on your way. For instance,

```
: 2,,3
invalid expression
r(3000);
: _
```

`2,,3` makes no sense to Mata, so Mata complained. This is an example of what is called a compile-time error; Mata could not make sense out of what we typed.

The other kind of error is called a run-time error. For example, we have no variable called `y`. Let us ask Mata to show us the contents of `y`:

```
: y
<istmt>: 3499 y not found
r(3499);
: _
```

Here what we typed made perfect sense—show me  $y$ —but  $y$  has never been defined. This ugly message is called a run-time error message—see [M-2] **errors** for a complete description—but all that’s important is to understand the difference between

```
invalid expression
```

and

```
<istmt>: 3499 y not found
```

The run-time message is prefixed by an identity (<istmt> here) and a number (3499 here). Mata is telling us, “I was executing your *istmt* [that’s what everything you type is called] and I got error 3499, the details of which are that I was unable to find  $y$ .”

The compile-time error message is of a simpler form: `invalid expression`. When you get such unprefixed error messages, that means Mata could not understand what you typed. When you get the more complicated error message, that means Mata understood what you typed, but there was a problem performing your request.

Another way to tell the difference between compile-time errors and run-time errors is to look at the return code. Compile-time errors have a return code of 3000:

```
: 2,,3
invalid expression
r(3000);
```

Run-time errors have a return code that might be in the 3000s, but is never 3000 exactly:

```
: y
                                <istmt>: 3499 y not found
r(3499);
```

Whether the error is compile-time or run-time, once the error message is issued, Mata is ready to continue just as if the error never happened.

## Working with real numbers, complex numbers, and strings

As we have seen, Mata works with real numbers:

```
: 2 + 3
5
```

Mata also understands complex numbers; you write the imaginary part by suffixing a lowercase  $i$ :

```
: 1+2i + 4-1i
5 + 1i
```

For imaginary numbers, you can omit the real part:

```
: 1+2i - 2i
1
```

Whether a number is real or complex, you can use the same computer notation for the imaginary part as you would for the real part:

```
: 2.5e+3i
2500i
: 1.25e+2+2.5e+3i          /* i.e., 1.25e+02 + 2.5e+03i */
125 + 2500i
```

We purposely wrote the last example in nearly unreadable form just to emphasize that Mata could interpret it.

Mata also understands strings, which you write enclosed in double quotes:

```
: "Alpha" + "Beta"
AlphaBeta
```

Just like Stata, Mata understands simple and compound double quotes:

```
: 'Alpha' + 'Beta'
AlphaBeta
```

You can add complex and reals,

```
: 1+2i + 3
4+2i
```

but you may not add reals or complex to strings:

```
: 2 + "alpha"
type mismatch: real + string not allowed
r(3000);
```

We got a run-time error. Mata understood `2 + "alpha"` all right; it just could not perform our request.

## Working with scalars, vectors, and matrices

In addition to understanding scalars—be they real, complex, or string—Mata understands vectors and matrices of real, complex, and string elements:

```
: x = (1, 2)
: x
      1  2
1 | 1  2
```

`x` now contains the row vector (1, 2). We can add vectors:

```
: x + (3, 4)
      1  2
1 | 4  6
```

The “,” is the [column-join operator](#); things like (1, 2) are expressions, just as (1 + 2) is an expression:

```
: y = (3, 4)
: z = (x, y)
: z
      1  2  3  4
1 | 1  2  3  4
```

In the above, we could have dispensed with the parentheses and typed “`y = 3, 4`” followed by “`z = x, y`”, just as we could using the `+` operator, although most people find vectors more readable when enclosed in parentheses.

$\backslash$  is the **row-join** operator:

```
: a = (1 \ 2)
```

```
: a
```

```
1
```

1	1
2	2

```
:
```

```
: b = (3 \ 4)
```

```
: c = (a \ b)
```

```
: c
```

```
1
```

1	1
2	2
3	3
4	4

Using the column-join and row-join operators, we can enter matrices:

```
: A = (1, 2 \ 3, 4)
```

```
: A
```

```
1 2
```

1	1	2
2	3	4

The use of these operators is not limited to scalars. Remember,  $x$  is the row vector  $(1, 2)$ ,  $y$  is the row vector  $(3, 4)$ ,  $a$  is the column vector  $(1 \ \backslash \ 2)$ , and  $b$  is the column vector  $(3 \ \backslash \ 4)$ . Therefore,

```
: x \ y
```

```
1 2
```

1	1	2
2	3	4

```
: a, b
```

```
1 2
```

1	1	3
2	2	4

But if we try something nonsensical, we get an error:

```
: a, x
```

```
<istmt>: 3200 conformability error
```

We create complex vectors and matrices just as we create real ones, the only difference being that their elements are complex:

```
: Z = (1 + 1i, 2 + 3i \ 3 - 2i, -1 - 1i)
```

```
: Z
```

```
1 2
```

1	1 + 1i	2 + 3i
2	3 - 2i	-1 - 1i

Similarly, we can create string vectors and matrices, which are vectors and matrices with string elements:

```
: S = ("1st element", "2nd element" \ "another row", "last element")
: S
```

	1	2
1	1st element	2nd element
2	another row	last element

For strings, the individual elements can be up to 2,147,483,647 characters long.

## Working with functions

Mata's expressions also include functions:

```
: sqrt(4)
2
: sqrt(-4)
.
```

When we ask for the square root of  $-4$ , Mata replies “.” Further, . can be stored just like any other number:

```
: findout = sqrt(-4)
: findout
.
```

“.” means missing, that there is no answer to our calculation. Taking the square root of a negative number is not an error; it merely produces missing. To Mata, missing is a number like any other number, and the rules for all the operators have been generalized to understand missing. For instance, the addition rule is generalized such that missing plus anything is missing:

```
: 2 + .
.
```

Still, it should surprise you that Mata produced missing for the `sqrt(-4)`. We said that Mata understands complex numbers, so should not the answer be  $2i$ ? The answer is that it should be if you are working on the complex plane, but otherwise, missing is probably a better answer. Mata attempts to intuit the kind of answer you want by context, and in particular, uses inheritance rules. If you ask for the square root of a real number, you get a real number back. If you ask for the square root of a complex number, you get a complex number back:

```
: sqrt(-4 + 0i)
2i
```

Here complex means multipart:  $-4 + 0i$  is a complex number; it merely happens to have 0 imaginary part. Thus:

```
: areal = -4
: acomplex = -4 + 0i
: sqrt(areal)
.
: sqrt(acomplex)
2i
```

If you ever have a real scalar, vector, or matrix, and want to make it complex, use the `C()` function, which means “convert to complex”:

```
: sqrt(C(areal))
2i
```

`C()` is documented in [M-5] `C()`. `C()` allows one or two arguments. With one argument, it casts to complex. With two arguments, it makes a complex out of the two real arguments. Thus you could type

```
: sqrt(-4 + 2i)
.485868272 + 2.05817103i
```

or you could type

```
: sqrt(C(-4, 2))
.485868272 + 2.05817103i
```

By the way, used with one argument, `C()` also allows complex, and then it does nothing:

```
: sqrt(C(acomplex))
2i
```

## Distinguishing real and complex values

It is virtually impossible to tell the difference between a real value and a complex value with zero imaginary part:

```
: areal = -4
: acomplex = -4 + 0i
: areal
-4
: acomplex
-4
```

Yet, as we have seen, the difference is important: `sqrt(areal)` is missing, `sqrt(acomplex)` is `2i`. One solution is the `eltype()` function:

```
: eltype(areal)
real
: eltype(acomplex)
complex
```

`eltype()` can also be used with strings,

```
: astring = "hello"
: eltype(astring)
string
```

but this is useful mostly in programming contexts.

## Working with matrix and scalar functions

Some functions are matrix functions: they require a matrix and return a matrix. Mata's `invsym(X)` is an example of such a function. It returns the matrix that is the inverse of symmetric, real matrix  $X$ :

```
: X = (76, 53, 48 \ 53, 88, 46 \ 48, 46, 63)
: Xi = invsym(X)
: Xi
[symmetric]
```

	1	2	3
1	.0298458083		
2	-.0098470272	.0216268926	
3	-.0155497706	-.0082885675	.0337724301

```
: Xi * X
```

	1	2	3
1	1	-8.67362e-17	-8.50015e-17
2	-1.38778e-16	1	-1.02349e-16
3	0	1.11022e-16	1

The last matrix,  $\text{Xi} * X$ , differs just a little from the identity matrix because of unavoidable computational roundoff error.

Other functions are, mathematically speaking, scalar functions. `sqrt()` is an example in that it makes no sense to speak of `sqrt(X)`. (That is, it makes no sense to speak of `sqrt(X)` unless we were speaking of the Cholesky square-root decomposition. Mata has such a matrix function; see [M-5] `cholesky(.)`.)

When a function is, mathematically speaking, a scalar function, the corresponding Mata function will usually allow vector and matrix arguments and, then, the Mata function makes the calculation on each element individually:

```
: M = (1, 2 \ 3, 4 \ 5, 6)
: M
```

	1	2
1	1	2
2	3	4
3	5	6

```
:
: S = sqrt(M)
: S
```

	1	2
1	1	1.414213562
2	1.732050808	2
3	2.236067977	2.449489743

```
:
: S[1,2]*S[1,2]
2
: S[2,1]*S[2,1]
3
```

When a function returns a result calculated in this way, it is said to return an element-by-element result.

### Performing element-by-element calculations: Colon operators

Mata's operators, such as + (addition) and \* (multiplication), work as you would expect. In particular, \* performs matrix multiplication:

```
: A = (1, 2 \ 3, 4)
: B = (5, 6 \ 7, 8)
: A*B
```

	1	2
1	19	22
2	43	50

The first element of the result was calculated as  $1 * 5 + 2 * 7 = 19$ .

Sometimes, you really want the element-by-element result. When you do, place a colon in front of the operator: Mata's \*: operator performs element-by-element multiplication:

```
: A:*B
```

	1	2
1	5	12
2	21	32

See [\[M-2\] op\\_colon](#) for more information.

### Writing programs

Mata is a complete programming language; it will allow you to create your own functions:

```
: function add(a,b) return(a+b)
```

That single statement creates a new function, although perhaps you would prefer if we typed it as

```
: function add(a,b)
> {
>     return(a+b)
> }
```

because that makes it obvious that a program can contain many lines. In either case, once defined, we can use the function:

```
: add(1,2)
3
: add(1+2i,4-1i)
5 + 1i
: add( (1,2), (3,4) )
```

	1	2
1	4	6



```

: add(x,y)
      1      2
1  

|   |   |
|---|---|
| 4 | 6 |
|---|---|



: add(A,A)
      1      2
1  

|   |   |
|---|---|
| 2 | 4 |
| 6 | 8 |


2

:
: Z1 = (1+1i, 1+1i \ 2, 2i)
: Z2 = (1+2i, -3+3i \ 6i, -2+2i)
: add(Z1, Z2)
      1      2
1  

|        |         |
|--------|---------|
| 2 + 3i | -2 + 4i |
| 2 + 6i | -2 + 4i |


2

:
: add("Alpha","Beta")
AlphaBeta

:
: S1 = ("one", "two" \ "three", "four")
: S2 = ("abc", "def" \ "ghi", "jkl")
: add(S1, S2)
      1      2
1  

|          |         |
|----------|---------|
| oneabc   | twodef  |
| threeghi | fourjkl |


2

```

Of course, our little function `add()` does not do anything that the `+` operator does not already do, but we could write a program that did do something different. The following program will allow us to make  $n \times n$  identity matrices:

```

: real matrix id(real scalar n)
> {
>   real scalar i
>   real matrix res
>
>   res = J(n, n, 0)
>   for (i=1; i<=n; i++) {
>     res[i,i] = 1
>   }
>   return(res)
> }

:
: I3 = id(3)
: I3
[symmetric]
      1      2      3
1  

|   |   |   |
|---|---|---|
| 1 |   |   |
| 0 | 1 |   |
| 0 | 0 | 1 |


2
3

```

The function `J()` in the program line `res = J(n, n, 0)` is a Mata built-in function that returns an  $n \times n$  matrix containing 0s (`J(r, c, val)` returns an  $r \times c$  matrix, the elements of which are all equal to *val*); see [M-5] `J()`.

`for (i=1; i<=n; i++)` says that starting with `i=1` and so long as `i<=n` do what is inside the braces (set `res[i,i]` equal to 1) and then (we are back to the `for` part again), increment `i`.

The final line—`return(res)`—says to return the matrix we have just created.

Actually, just as with `add()`, we do not need `id()` because Mata has a built-in function `I(n)` that makes identity matrices, but it is interesting to see how the problem could be programmed.

## More functions

Mata has many functions already and much of this manual concerns documenting what those functions do; see [M-4] [intro](#). But right now, what is important is that many of the functions are themselves written in Mata!

One of those functions is `pi()`; it takes no arguments and returns the value of  $\pi$ . The code for it reads

```
real scalar pi() return(3.141592653589793238462643)
```

There is no reason to type the above function because it is already included as part of Mata:

```
: pi()
3.141592654
```

When Mata lists a result, it does not show as many digits, but we could ask to see more:

```
: printf("%17.0g", pi())
3.14159265358979
```

Other Mata functions include the hyperbolic function `tanh(u)`. The code for `tanh(u)` reads

```
numeric matrix tanh(numeric matrix u)
{
    numeric matrix eu, emu
    eu = exp(u)
    emu = exp(-u)
    return( (eu-emu)/(eu+emu) )
}
```

See for yourself: at the Stata dot prompt (not the Mata colon prompt), type

```
. viewsource tanh.mata
```

When the code for a function was written in Mata (as opposed to having been written in C), `viewsource` can show you the code; see [M-1] [source](#).

Returning to the function `tanh()`,

```
numeric matrix tanh(numeric matrix u)
{
    numeric matrix eu, emu

    eu = exp(u)
    emu = exp(-u)
    return( (eu-emu)/(eu+emu) )
}
```

this is the first time we have seen the word `numeric`: it means real or complex. Built-in (previously written) function `exp()` works like `sqrt()` in that it allows a real or complex argument and correspondingly returns a real or complex result. Said in Mata jargon: `exp()` allows a `numeric` argument and correspondingly returns a `numeric` result. `tanh()` will also work like `sqrt()` and `exp()`.

Another characteristic `tanh()` shares with `sqrt()` and `exp()` is element-by-element operation. `tanh()` is element-by-element because `exp()` is element-by-element and because we were careful to use the `:/` (element-by-element) divide operator.

In any case, there is no need to type the above functions because they are already part of Mata. You could learn more about them by seeing their manual entry, [\[M-5\] `sin\(\)`](#).

At the other extreme, Mata functions can become long. Here is Mata's function to solve  $AX = B$  for  $X$  when  $A$  is lower triangular, placing the result  $X$  back into  $A$ :

```
real scalar _solvelower(
    numeric matrix A, numeric matrix b,
    |real scalar usertol, numeric scalar userd)
{
    real scalar          tol, rank, a_t, b_t, d_t
    real scalar          n, m, i, im1, complex_case
    numeric rowvector    sum
    numeric scalar       zero, d

    d = userd

    if ((n=rows(A))!=cols(A)) _error(3205)
    if (n != rows(b))         _error(3200)
    if (isview(b))             _error(3104)
    m = cols(b)
    rank = n

    a_t = iscomplex(A)
    b_t = iscomplex(b)
    d_t = d<. ? iscomplex(d) : 0

    complex_case = a_t | b_t | d_t
```

```

    if (complex_case) {
        if (!a_t) A = C(A)
        if (!b_t) b = C(b)
        if (d<. & !d_t) d = C(d)
        zero = 0i
    }
    else zero = 0

    if (n==0 | m==0) return(0)

    tol = solve_tol(A, usertol)

    if (abs(d) >=. ) {
        if (abs(d=A[1,1])<=tol) {
            b[1,.] = J(1, m, zero)
            --rank
        }
        else {
            b[1,.] = b[1,.] :/ d
            if (missing(d)) rank = .
        }

        for (i=2; i<=n; i++) {
            im1 = i - 1
            sum = A[|i,1i,im1|] * b[|1,1\im1,m|]
            if (abs(d=A[i,i])<=tol) {
                b[i,.] = J(1, m, zero)
                --rank
            }
            else {
                b[i,.] = (b[i,.]-sum) :/ d
                if (missing(d)) rank = .
            }
        }
    }
    else {
        if (abs(d)<=tol) {
            rank = 0
            b = J(rows(b), cols(b), zero)
        }
        else {
            b[1,.] = b[1,.] :/ d

            for (i=2; i<=n; i++) {
                im1 = i - 1
                sum = A[|i,1i,im1|] * b[|1,1\im1,m|]
                b[i,.] = (b[i,.]-sum) :/ d
            }
        }
    }

    return(rank)
}

```

If the function were not already part of Mata and you wanted to use it, you could type it into a do-file or onto the end of an ado-file (especially good if you just want to use `_solvelower()` as a subroutine). In those cases, do not forget to enter and exit Mata:

---

```

program mycommand
...
  ado-file code appears here
...
end
mata:
  _solvelower() code appears here
end

```

---

begin ado-file

end ado-file

Sharp-eyed readers will notice that we put a colon on the end of the Mata command. That's a detail, and why we did that is explained in [M-3] [mata](#).

In addition to loading functions by putting their code in do- and ado-files, you can also save the compiled versions of functions in .mo files (see [M-3] [mata mosave](#)) or into .mlib Mata libraries (see [M-3] [mata mlib](#)).

For `_solvelower()`, it has already been saved into a library, namely, Mata's official library, so you need not do any of this.

## Mata environment commands

When you are using Mata, there is a set of commands that will tell you about and manipulate Mata's environment.

The most useful such command is `mata describe`; see [M-3] [mata describe](#):

```

: mata describe

```

# bytes	type	name and extent
76	transmorphic matrix	add()
200	real matrix	id()
32	real matrix	A[2,2]
32	real matrix	B[2,2]
72	real matrix	I3[3,3]
48	real matrix	M[3,2]
48	real matrix	S[3,2]
47	string matrix	S1[2,2]
44	string matrix	S2[2,2]
72	real matrix	X[3,3]
72	real matrix	Xi[3,3]
64	complex matrix	Z[2,2]
64	complex matrix	Z1[2,2]
64	complex matrix	Z2[2,2]
16	real colvector	a[2]
16	complex scalar	acomplex
8	real scalar	areal
16	real colvector	b[2]
32	real colvector	c[4]
8	real scalar	findout
16	real rowvector	x[2]
16	real rowvector	y[2]
32	real rowvector	z[4]

---

```

: _

```

Another useful command is `mata clear` (see [\[M-3\] mata clear](#)), which will clear Mata without disturbing Stata:

```
: mata clear
: mata describe
      # bytes   type                name and extent
-----
-----
```

There are other useful `mata` commands; see [\[M-3\] intro](#). Do not confuse this command `mata`, which you type at Mata’s colon prompt, with Stata’s command `mata`, which you type at Stata’s dot prompt and which invokes Mata.

## Exiting Mata

When you are done using Mata, type `end` to Mata’s colon prompt:

```
: end
-----
. -
```

Exiting Mata does not clear it:

```
. mata
----- mata (type end to exit) -----
: x = 2
: y = (3 + 2i)
: function add(a,b) return(a+b)
: end
-----
. ...
. mata
----- mata (type end to exit) -----
: mata describe
      # bytes   type                name and extent
-----
      76   transmorphic matrix    add()
       8    real scalar           x
      16   complex scalar         y
-----
: end
```

Exiting Stata clears Mata, as does Stata’s `clear mata` command; see [\[D\] clear](#).

## Also see

[\[M-1\] intro](#) — Introduction and advice

# Title

[M-1] **help** — Obtaining online help

## Syntax

```
help m# entryname
```

```
help mata functionname()
```

The `help` command may be issued at either Stata's dot prompt or Mata's colon prompt.

## Description

Help for Mata is available online in Stata. This entry describes how to access it.

## Remarks

To see this entry online, type

```
. help m1 help
```

at Stata's dot prompt or Mata's colon prompt. You type that because this entry is [\[M-1\] \*\*help\*\*](#).

To see the entry for function `max()`, for example, type

```
. help mata max()
```

`max()` is documented in [\[M-5\] \*\*minmax\(\)\*\*](#), but that will not matter; Stata will find the appropriate entry.

To enter the Mata help system from the top, from whence you can click your way to any section or function, type

```
. help mata
```

To access Mata's PDF manual, click on the title.

## Also see

[\[R\] \*\*help\*\*](#) — Display online help

[\[M-3\] \*\*mata help\*\*](#) — Obtain online help

[\[M-1\] \*\*intro\*\*](#) — Introduction and advice

# Title

[M-1] how — How Mata works

## Description

Below we take away some of the mystery and show you how Mata works. Everyone, we suspect, will find this entertaining, and advanced users will find the description useful for predicting what Mata will do when faced with unusual situations.

## Remarks

Remarks are presented under the following headings:

*What happens when you define a program*  
*What happens when you work interactively*  
*What happens when you type a mata environment command*  
*Working with object code I: .mo files*  
*Working with object code II: .mlib libraries*  
*The Mata environment*

If you are reading the entries in the order suggested in [M-0] [intro](#), browse [M-1] [intro](#) next for sections that interest you, and then see [M-2] [syntax](#).

## What happens when you define a program

Let's say you fire up Mata and type

```
: function tryit()
> {
>     real scalar i
>
>     for (i=1; i<=10; i++) i
> }
```

Mata compiles the program: it reads what you type and produces binary codes that tell Mata exactly what it is to do when the time comes to execute the program. In fact, given the above program, Mata produces the binary code

```
00b4 3608 4000 0000 0100 0000 2000 0000
0000 0000 ffff ffff 0300 0000 0000 0000
0100 7472 7969 7400 1700 0100 1f00 0700
0000 0800 0000 0200 0100 0800 2a00 0300
1e00 0300
```

which looks meaningless to you and me, but Mata knows exactly what to make of it. The compiled version of the program is called object code, and it is the object code, not the original source code, that Mata stores in memory. In fact, the original source is discarded once the object code has been stored.

It is this compilation step—the conversion of text into object code—that makes Mata able to execute programs so quickly.



Later, when the time comes to execute the program, Stata follows the instructions it has previously recorded:

```
: tryit()
1
2
3
4
5
6
7
8
9
10
```

## What happens when you work interactively

Let's say you type

```
: x = 3
```

In the jargon of Mata, that is called an *istmt*—an interactive statement. Obviously, Mata stores 3 in *x*, but how?

Mata first compiles the single statement and stores the resulting object code under the name *<istmt>*. The result is much as if you had typed

```
: function <istmt>()
> {
>     x = 3
> }
```

except, of course, you could not define a program named *<istmt>* because the name is invalid. Mata has ways of getting around that.

At this point, Mata has discarded the source code *x = 3* and has stored the corresponding object code. Next, Mata executes *<istmt>*. The result is much as if you had typed

```
: <istmt>()
```

That done, there is only one thing left to do, which is to discard the object code. The result is much as if you typed

```
: mata drop <istmt>()
```

So there you have it: you type

```
: x = 3
```

and Mata executes

```
: function <istmt>()
> {
>     x = 3
> }
: <istmt>()
: mata drop <istmt>()
```

## □ Technical note

The above story is not exactly true because, as told, variable `x` would be local to function `<istmt>()` so, when `<istmt>()` concluded execution, variable `x` would be discarded. To prevent that from happening, Mata makes all variables defined by `<istmt>()` global. Thus you can type

```
: x = 3
```

followed by

```
: y = x + 2
```

and all works out just as you expect: `y` is set to 5.



## What happens when you type a mata environment command

When you are at a colon prompt and type something that begins with the word `mata`, such as

```
: mata describe
```

or

```
: mata clear
```

something completely different happens: Mata freezes itself and temporarily transfers control to a command interpreter like Stata's. The command interpreter accesses Mata's environment and reports on it or changes it. Once done, the interpreter returns to Mata, which comes back to life, and issues a new colon prompt:

```
: _
```

Once something is typed at the prompt, Mata will examine it to determine if it begins with the word `mata` (in which case the interpretive process repeats), or if it is the beginning of a function definition (in which case the program will be compiled but not executed), or anything else (in which case Mata will try to compile and execute it as an `<istmt>()`).

## Working with object code I: .mo files

Everything hinges on the object code that Mata produces, and, if you wish, you can save the object code on disk. The advantage of doing this is that, at some future date, your program can be executed without compilation, which saves time. If you send the object code to others, they can use your program without ever seeing the source code behind it.

After you type, say,

```
: function tryit()
> {
>     real scalar i
>
>     for (i=1; i<=10; i++) i
> }
```

Mata has created the object code and discarded the source. If you now type

```
: mata mosave tryit()
```

the Mata interpreter will create file `tryit.mo` in the current directory; see [\[M-3\] mata mosave](#). The new file will contain the object code.

At some future date, were you to type

```
: tryit()
```

without having first defined the program, Mata would look along the ado-path (see [P] [sysdir](#) and [U] [17.5 Where does Stata look for ado-files?](#)) for a file named `tryit.mo`. Finding the file, Mata loads it (so Mata now has the object code and executes it in the usual way).

## Working with object code II: .mlib libraries

Object code can be saved in `.mlib` libraries (files) instead of `.mo` files. `.mo` files contain the object code for one program. `.mlib` files contain the object code for a group of files.

The first step is to choose a name (we will choose `lmylib`—library names are required to start with the lowercase letter *l*) and create an empty library of that name:

```
: mata mlib create lmylib
```

Once created, new functions can be added to the library:

```
: mata mlib add lmylib tryit()
```

New functions can be added at any time, either immediately after creation or later—even much later; see [M-3] [mata mlib](#).

We mentioned that when Mata needs to execute a function that it does not find in memory, Mata looks for a `.mo` file of the same name. Before Mata does that, however, Mata thumbs through its libraries to see if it can find the function there.

## The Mata environment

Certain settings of Mata affect how it behaves. You can see those settings by typing `mata query` at the Mata prompt:

```
: mata query
```

---

```

Mata settings
  set matastrict      off
  set matalnum        off
  set mataoptimize    on
  set matafavor       space      may be space or speed
  set matacache       400        kilobytes
  set matalibs        lmatabase;lmataopt;lmataado
  set matamofirst     off
```

```
: _
```

You can change these settings by using `mata set`; see [M-3] [mata set](#). We recommend the default settings, except that we admit to being partial to `mata set matastrict on`.

## Reference

Gould, W. W. 2006. [Mata Matters: Precision](#). *Stata Journal* 6: 550–560.

# Also see

[M-1] [intro](#) — Introduction and advice

## Description

With Mata, you simply type matrix formulas to obtain the desired results. Below we provide guidelines when doing this with statistical formulas.

## Remarks

You have data and statistical formulas that you wish to calculate, such as  $b = (X'X)^{-1}X'y$ . Perform the following nine steps:

1. Start in Stata. Load the data.
2. If you are doing time-series analysis, generate new variables containing any *op.varname* variables you need, such as `l.gnp` or `d.r`.
3. Create a constant variable (`. gen cons = 1`). In most statistical formulas, you will find it useful.
4. Drop variables that you will not need. This saves memory and makes some things easier because you can just refer to all the variables.
5. Drop observations with missing values. Mata understands missing values, but Mata is a matrix language, not a statistical system, so Mata does not always ignore observations with missing values.
6. Put variables on roughly the same numeric scale. This is optional, but we recommend it. We explain what we mean and how to do this below.
7. Enter Mata. Do that by typing `mata` at the Stata command prompt. Do not type a colon after the `mata`. This way, when you make a mistake, you will stay in Mata.
8. Use Mata's `st_view()` function (see [\[M-5\] st\\_view\(\)](#)) to create matrices based on your Stata dataset. Create all the matrices you want or find convenient. The matrices created by `st_view()` are in fact views onto one copy of the data.
9. Perform your matrix calculations.

If you are reading the entries in the order suggested in [\[M-0\] intro](#), see [\[M-1\] how](#) next.

### 1. Start in Stata; load the data

We will use the `auto` dataset and will fit the regression

$$\text{mpg}_j = b_0 + b_1 \text{weight}_j + b_2 \text{foreign}_j + e_j$$

by using the formulas

$$\mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$$

$$\mathbf{V} = s^2(\mathbf{X}'\mathbf{X})^{-1}$$

where

$$s^2 = \mathbf{e}'\mathbf{e}/(n - k)$$

$$\mathbf{e} = \mathbf{y} - \mathbf{X}\mathbf{b}$$

$$n = \text{rows}(\mathbf{X})$$

$$k = \text{cols}(\mathbf{X})$$

We begin by typing

```
. sysuse auto
(1978 Automobile Data)
```

## 2. Create any time-series variables

We do not have any time-series variables but, just for a minute, let's pretend we did. If our model contained lagged gnp, we would type

```
. gen lngp = l.gnp
```

so that we would have a new variable `lngp` that we would use in place of `l.gnp` in the subsequent steps.

## 3. Create a constant variable

```
. gen cons = 1
```

## 4. Drop unnecessary variables

We will need the variables `mpg`, `weight`, `foreign`, and `cons`, so it is easier for us to type `keep` instead of `drop`:

```
. keep mpg weight foreign cons
```

## 5. Drop observations with missing values

We do not have any missing values in our data, but let's pretend we did, or let's pretend we are uncertain. Here is an easy trick for getting rid of observations with missing values:

```
. regress mpg weight foreign cons
(output omitted)
. keep if e(sample)
```

We estimated a regression using all the variables and then kept the observations `regress` chose to use. It does not matter which variable you choose as the dependent variable, nor the order of the independent variables, so we just as well could have typed

```
. regress weight mpg foreign cons
(output omitted)
. keep if e(sample)
```

or even

```
. regress cons mpg weight foreign
(output omitted)
. keep if e(sample)
```

The output produced by `regress` is irrelevant, even if some variables are dropped. We are merely borrowing `regress`'s ability to identify the subsample with no missing values.

Using `regress` causes Stata to make many unnecessary calculations and, if that offends you, here is a more sophisticated alternative:

```
. local 0 "mpg weight foreign cons"
. syntax varlist
. marksample touse
. keep if `touse'
. drop `touse'
```

Using `regress` is easier.

## 6. Put variables on roughly the same numeric scale

This step is optional, but we recommend it. You are about to use formulas that have been derived by people who assumed that the usual rules of arithmetic hold, such as  $(a + b) - c = a + (b - c)$ . Many of the standard rules, such as the one shown, are violated when arithmetic is performed in finite precision, and this leads to roundoff error in the final, calculated results.

You can obtain a lot of protection by making sure that your variables are on roughly the same scale, by which we mean their means and standard deviations are all roughly equal. By roughly equal, we mean equal up to a factor of 1,000 or so. So let's look at our data:

```
. summarize
```

Variable	Obs	Mean	Std. Dev.	Min	Max
mpg	74	21.2973	5.785503	12	41
weight	74	3019.459	777.1936	1760	4840
foreign	74	.2972973	.4601885	0	1
cons	74	1	0	1	1

Nothing we see here bothers us much. Variable `weight` is the largest, with a mean and standard deviation that are 1,000 times larger than those of the smallest variable, `foreign`. We would feel comfortable, but only barely, ignoring scale differences. If `weight` were 10 times larger, we would begin to be concerned, and our concern would grow as `weight` grew.

The easiest way to address our concern is to divide `weight` so that, rather than measuring weight in pounds, it measures weight in thousands of pounds:

```
. replace weight = weight/1000
. summarize
```

Variable	Obs	Mean	Std. Dev.	Min	Max
mpg	74	21.2973	5.785503	12	41
weight	74	3.019459	.7771936	1.76	4.84
foreign	74	.2972973	.4601885	0	1
cons	74	1	0	1	1

What you are supposed to do is make the means and standard deviations of the variables roughly equal. If `weight` had a large mean and reasonable standard deviation, we would have subtracted, so that we would have had a variable measuring weight in excess of some number of pounds. Or we could do both, subtracting, say, 2,000 and then dividing by 100, so we would have weight in excess of 2,000 pounds, measured in 100-pound units.

Remember, the definition of roughly equal allows lots of leeway, so you do not have to give up easy interpretation.

## 7. Enter Mata

We type

```
. mata
_____ mata (type end to exit) _____
: _
```

Mata uses a colon prompt, whereas Stata uses a period.

## 8. Use Mata's `st_view()` function to access your data

Our matrix formulas are

$$\mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$$

$$\mathbf{V} = s^2(\mathbf{X}'\mathbf{X})^{-1}$$

where

$$s^2 = \mathbf{e}'\mathbf{e}/(n - k)$$

$$\mathbf{e} = \mathbf{y} - \mathbf{X}\mathbf{b}$$

$$n = \text{rows}(\mathbf{X})$$

$$k = \text{cols}(\mathbf{X})$$

so we are going to need  $\mathbf{y}$  and  $\mathbf{X}$ .  $\mathbf{y}$  is an  $n \times 1$  column vector of dependent-variable values, and  $\mathbf{X}$  is an  $n \times k$  matrix of the  $k$  independent variables, including the constant. Rows are observations, columns are variables.

We make the vector and matrix as follows:

```
: st_view(y=., ., "mpg")
: st_view(X=., ., ("weight", "foreign", "cons"))
```

Let us explain. We wish we could type

```
: y = st_view(., "mpg")
: X = st_view(., ("weight", "foreign", "cons"))
```

because that is what the functions are really doing. We cannot because `st_view()` (unlike all other Mata functions), returns a special kind of matrix called a view. A view acts like a regular matrix in nearly every respect, but views do not consume nearly as much memory, because they are in fact views onto the underlying Stata dataset!

We could instead create  $\mathbf{y}$  and  $\mathbf{X}$  with Mata's `st_data()` function (see [M-5] `st_data()`), and then we could type the creation of  $\mathbf{y}$  and  $\mathbf{X}$  the natural way,

```
: y = st_data(., "mpg")
: X = st_data(., ("weight", "foreign", "cons"))
```



`st_data()` returns a real matrix, which is a copy of the data Stata has stored in memory.

We could use `st_data()` and be done with the problem. For our automobile-data example, that would be a fine solution. But were the automobile data larger, you might run short of memory, and views can save lots of memory. You can create views willy-nilly—lots and lots of them—and never consume much memory! Views are wonderfully convenient and it is worth mastering the little bit of syntax to use them.

`st_view()` requires three arguments: the name of the view matrix to be created, the observations (rows) the matrix is to contain, and the variables (columns). If we wanted to create a view matrix `Z` containing all the observations and all the variables, we could type

```
: st_view(Z, ., .)
```

`st_view()` understands missing value in the second and third positions to mean all the observations and all the variables. Let's try it:

```
: st_view(Z, ., .)
                                err:<istmt>: 3499 Z not found
r(3499);
: _
```

That did not work because Mata requires `Z` to be predefined. The reasons are technical, but it should not surprise you that function arguments need to be defined before a function can be used. Mata just does not understand that `st_view()` really does not need `Z` defined. The way around Mata's confusion is to define `Z` and then let `st_view()` redefine it:

```
: Z = .
: st_view(Z, ., .)
```

You can, if you wish, combine all that into one statement

```
: st_view(Z=., ., .)
```

and that is what we did when we defined `y` and `X`:

```
: st_view(y=., ., "mpg")
: st_view(X=., ., ("weight", "foreign", "cons"))
```

The second argument `(.)` specified that we wanted all the observations, and the third argument specified the variables we wanted. Be careful not to omit the “extra” parentheses when typing the variables. Were you to type

```
: st_view(X=., ., "weight", "foreign", "cons")
```

you would be told you typed an invalid expression. `st_view()` expects three arguments, and the third argument is a row vector specifying the variables to be selected: `("weight", "foreign", "cons")`.

At this point, we suggest you type

```
: y
(output omitted)
: X
(output omitted)
```

to see that `y` and `X` really do contain our data. In case you have lost track of what we have typed, here is our complete session so far:

```

. sysuse auto
. gen cons = 1
. keep mpg weight foreign cons
. regress mpg weight foreign cons
. keep if e(sample)
. replace weight = weight/1000
. mata
: st_view(y=., ., "mpg")
: st_view(X=., ., ("weight", "foreign", "cons"))

```

## 9. Perform your matrix calculations

To remind you: our matrix calculations are

$$\mathbf{b} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$$

$$\mathbf{V} = s^2(\mathbf{X}'\mathbf{X})^{-1}$$

where

$$s^2 = \mathbf{e}'\mathbf{e}/(n - k)$$

$$\mathbf{e} = \mathbf{y} - \mathbf{X}\mathbf{b}$$

$$n = \text{rows}(\mathbf{X})$$

$$k = \text{cols}(\mathbf{X})$$

Let's get our regression coefficients,

```

: b = invsym(X'X)*X'y
: b

```

1	-6.587886358
2	-1.650029004
3	41.67970227

and let's form the residuals, define  $n$  and  $k$ , and obtain  $s^2$ ,

```

: e = y - X*b
: n = rows(X)
: k = cols(X)
: s2 = (e'e)/(n-k)

```

so we are able to calculate the variance matrix:

```

: V = s2*invsym(X'X)
: V
[symmetric]

```

	1	2	3
1	.4059128628		
2	.4064025078	1.157763273	
3	-1.346459802	-1.57131579	4.689594304

We are done.

We can present the results in more readable fashion by pulling the diagonal of  $V$  and calculating the square root of each element:

```
: se = sqrt(diagonal(V))
: (b, se)
```

	1	2
1	-6.587886358	.6371129122
2	-1.650029004	1.075994086
3	41.67970227	2.165547114

You know that if we were to type

```
: 2+3
5
```

Mata evaluates the expression and shows us the result, and that is exactly what happened when we typed

```
: (b, se)
```

$(b, se)$  is an expression, and Mata evaluated it and displayed the result. The expression means to form the matrix whose first column is  $b$  and second column is  $se$ . We could obtain a listing of the coefficient, standard error, and its  $t$  statistic by asking Mata to display  $(b, se, b:/se)$ ,

```
: (b, se, b:/se)
```

	1	2	3
1	-6.587886358	.6371129122	-10.34021793
2	-1.650029004	1.075994086	-1.533492633
3	41.67970227	2.165547114	19.24673077

In the expression above,  $b:/se$  means to divide the elements of  $b$  by the elements of  $se$ .  $:/$  is called a colon operator and you can learn more about it by seeing [\[M-2\] op\\_colon](#).

We could add the significance level by typing

```
: (b, se, b:/se, 2*ttail(n-k, abs(b:/se)))
```

	1	2	3	4
1	-6.587886358	.6371129122	-10.34021793	8.28286e-16
2	-1.650029004	1.075994086	-1.533492633	.1295987129
3	41.67970227	2.165547114	19.24673077	6.89556e-30

Those are the same results reported by `regress`; type

```
. sysuse auto
. replace weight = weight/1000
. regress mpg weight foreign
```

and compare results.

## Review

Our complete session was

```
. sysuse auto
. gen cons = 1
. keep mpg weight foreign cons
. regress mpg weight foreign cons
. keep if e(sample)
. replace weight = weight/1000

. mata
: st_view(y=., ., "mpg")
: st_view(X=., ., ("weight", "foreign", "cons"))
: b = invsym(X'X)*X'y
: b
: e = y - X*b
: n = rows(X)
: k = cols(X)
: s2= (e'e)/(n-k)
: V = s2*invsym(X'X)
: V
: se = sqrt(diagonal(V))
: (b, se)
: (b, se, b:/se)
: (b, se, b:/se, 2*ttail(n-k, abs(b:/se)))
: end
```

## Reference

Gould, W. W. 2006. [Mata Matters: Interactive use](#). *Stata Journal* 6: 387–396.

## Also see

[\[M-1\] intro](#) — Introduction and advice

# Title

[M-1] **LAPACK** — The LAPACK linear-algebra routines

## Description

LAPACK stands for Linear Algebra PACKage and is a freely available set of FORTRAN 90 routines for solving systems of simultaneous equations, eigenvalue problems, and singular value problems. Many of the LAPACK routines are based on older EISPACK and LINPACK routines, and the more modern LAPACK does much of its computation by using BLAS (Basic Linear Algebra Subprogram).

## Remarks

The LAPACK and BLAS routines form the basis for many of Mata's linear-algebra capabilities. Individual functions of Mata that use LAPACK routines always make note of that fact.

For up-to-date information on LAPACK, see <http://www.netlib.org/lapack/>.

Advanced programmers can directly access the LAPACK functions; see [M-5] **lapack()**.

## Acknowledgments

We thank the authors of LAPACK for their excellent work:

E. Anderson, Z. Bai, C. Bischof, S. Blackford, J. Demmel, J. Dongarra, J. Du Croz, A. Greenbaum, S. Hammarling, A. McKenney, and D. Sorensen.

## Reference

Anderson, E., Z. Bai, C. Bischof, S. Blackford, J. Demmel, J. J. Dongarra, J. Du Croz, A. Greenbaum, S. Hammarling, A. McKenney, and D. Sorensen. 1999. *LAPACK Users' Guide*. 3rd ed. Philadelphia: Society for Industrial and Applied Mathematics.

## Also see

[R] **copyright lapack** — LAPACK copyright notification

[M-5] **lapack()** — LAPACK linear-algebra functions

[M-1] **intro** — Introduction and advice

# Title

[M-1] limits — Limits and memory utilization

## Summary

Limits:

	Minimum	Maximum
Scalars, vectors, matrices		
rows	0	2,147,483,647
columns	0	2,147,483,647
String elements, length	0	2,147,483,647

Stata's `matsize` plays no role in these limits.

Size approximations:

	Memory requirements
real matrices	$oh + r*c*8$
complex matrices	$oh + r*c*16$
pointer matrices	$oh + r*c*8$
string matrices	$oh + r*c*8 + total\_length\_of\_strings$

where  $r$  and  $c$  represent the number of rows and columns and  
where  $oh$  is overhead and is approximately 64 bytes

## Description

Mata imposes limits, but those limits are of little importance compared with the memory requirements. Mata stores matrices in memory and requests the memory for them from the operating system.

## Remarks

Mata requests (and returns) memory from the operating system as it needs it, and if the operating system cannot provide it, Mata issues the following error:

```
: x = foo(A, B)
      foo(): 3900  unable to allocate ...
      <istmt>:    -  function returned error
r(3900);
```

Stata's `matsize` (see [R] [matsize](#)) and Stata's `set min_memory` and `set max_memory` values (see [D] [memory](#)) play no role in Mata or, at least, they play no direct role.

## Also see

- [M-3] [mata memory](#) — Report on Mata's memory usage
- [M-5] [mindouble\(\)](#) — Minimum and maximum nonmissing value
- [M-1] [intro](#) — Introduction and advice

# Title

[M-1] **naming** — Advice on naming functions and variables

## Syntax

A *name* is 1–32 characters long, the first character of which must be

A–Z      a–z      \_

and the remaining characters of which may be

A–Z      a–z      \_      0–9

except that names may not be a word reserved by Mata (see [M-2] **reswords** for a list).

Examples of names include

x	x2	alpha
logarithm_of_x	LogOfX	

Case matters: alpha, Alpha, and ALPHA are different names.

Variables and functions have separate name spaces, which means a variable and a function can have the same name, such as `value` and `value()`, and Mata will not confuse them.

## Description

Advice is offered on how to name variables and functions.

## Remarks

Remarks are presented under the following headings:

*[Interactive use](#)*

*[Naming variables](#)*

*[Naming functions](#)*

*[What happens when functions have the same names](#)*

*[How to determine if a function name has been taken](#)*

## Interactive use

Use whatever names you find convenient: Mata will tell you if there is a problem.

The following sections are for programmers who want to write code that will require the minimum amount of maintenance in the future.

## Naming variables

Mostly, you can name variables however you please. Variables are local to the program in which they appear, so one function can have a variable or argument named `x` and another function can have a variable or argument of the same name, and there is no confusion.

If you are writing a large system of programs that has global variables, on the other hand, we recommend that you give the global variables long names, preferably with a common prefix identifying your system. For instance,

```
multeq_no_of_equations
multeq_eq
multeq_inuse
```

This way, your variables will not become confused with variables from other systems.

## Naming functions

Our first recommendation is that, for the most part, you give functions all-lowercase names: `foo()` rather than `Foo()` or `F00()`. If you have a function with one or more capital letters in the name, and if you want to save the function's object code, you must do so in `.mlib` libraries; `.mo` files are not sufficient. `.mo` files require that filename be the same as the function name, which means `Foo.mo` for `Foo()`. Not all operating systems respect case in filenames. Even if your operating system does respect case, you will not be able to share your `.mo` files with others whose operating systems do not.

We have no strong recommendation against mixed case; we merely remind you to use `.mlib` library files if you use it.

Of greater importance is the name you choose. Mata provides many functions and more will be added over time. You will find it more convenient if you choose names that StataCorp and other users do not choose.

That means to avoid words that appear in the English-language dictionary and to avoid short names, say, those four characters or fewer. You might have guessed that `svd()` would be taken, but who would have guessed `lud()`? Or `qrd()`? Or `e()`?

Your best defense against new official functions, and other user-written functions, is to choose long function names.

## What happens when functions have the same names

There are two kinds of official functions: built-in functions and library functions. User-written functions are invariably library functions (here we draw no distinction between functions supplied in `.mo` files and those supplied in `.mlib` files).

Mata will issue an error message if you attempt to define a function with the same name as a built-in function.

Mata will let you define a new function with the same name as a library function if the library function is not currently in memory. If you store your function in a `.mo` file or a `.mlib` library, however, in the future the official Mata library function will take precedence over your function: your function will never be loaded. This feature works nicely for interactive users, but for long-term programming, you will want to avoid naming your functions after Mata functions.



A similar result is obtained if you name your function after a user-written function that is installed on your computer. You can do so if the user-written function is not currently in memory. In the future, however, one or the other function will take precedence and, no matter which, something will break.

## How to determine if a function name has been taken

Use `mata which` (see [M-3] [mata which](#)):

```
: mata which det_of_triangular()
function det_of_triangular() not found
r(111);
: mata which det()
det(): lmatbase
```

## Also see

[M-2] [reswords](#) — Reserved words

[M-1] [intro](#) — Introduction and advice

Syntax

Action	Permutation matrix notation	Permutation vector notation
permute rows	$B = P * A$	$B = A[p, .]$
permute columns	$B = A * P$	$B = A[. , p]$
unpermute rows	$B = P' A$	$B = A ; B[p, .] = A$ or $B = A[\text{invorder}(p), .]$
unpermute columns	$B = A * P'$	$B = A ; B[. , p] = A$ or $B = A[. , \text{invorder}(p)]$

A *permutation matrix* is an  $n \times n$  matrix that is a row (or column) permutation of the identity matrix.

A *permutation vector* is a  $1 \times n$  or  $n \times 1$  vector of the integers 1 through  $n$ .

The following permutation matrix and permutation vector are equivalent:

$$P = \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & 0 \end{bmatrix} \iff p = \begin{bmatrix} 2 \\ 3 \\ 1 \end{bmatrix}$$

Either can be used to permute the rows of

$$A = \begin{bmatrix} a & b & c & d \\ e & f & g & h \\ i & j & k & l \end{bmatrix} \text{ to produce } \begin{bmatrix} e & f & g & h \\ i & j & k & l \\ a & b & c & d \end{bmatrix}$$

and to permute the columns of

$$P = \begin{bmatrix} m & n & o \\ p & q & r \\ s & t & u \\ v & w & x \end{bmatrix} \text{ to produce } \begin{bmatrix} n & o & m \\ q & r & p \\ t & u & s \\ w & x & v \end{bmatrix}$$

Description

Permutation matrices are a special kind of orthogonal matrix that, via multiplication, reorder the rows or columns of another matrix. Permutation matrices cast the reordering in terms of multiplication.

Permutation vectors also reorder the rows or columns of another matrix, but they do it via subscripting. This alternative method of achieving the same end is, computerwise, more efficient, in that it uses less memory and less computer time.

The relationship between the two is shown below.

## Remarks

Remarks are presented under the following headings:

*Permutation matrices*

*How permutation matrices arise*

*Permutation vectors*

## Permutation matrices

A permutation matrix is a square matrix whose rows are a permutation of the identity matrix. The following are the full set of all  $2 \times 2$  permutation matrices:

$$\begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \quad (1)$$

$$\begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \quad (2)$$

Let  $P$  be an  $n \times n$  permutation matrix. If  $n \times m$  matrix  $A$  is premultiplied by  $P$ , the result is to reorder its rows. For example,

$$\begin{array}{ccccc} P & * & A & = & PA \\ \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & 0 \end{bmatrix} & * & \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix} & = & \begin{bmatrix} 4 & 5 & 6 \\ 7 & 8 & 9 \\ 1 & 2 & 3 \end{bmatrix} \end{array} \quad (3)$$

Above, we illustrated the reordering using square matrix  $A$ , but  $A$  did not have to be square.

If  $m \times n$  matrix  $B$  is postmultiplied by  $P$ , the result is to reorder its columns. We illustrate using square matrix  $A$  again:

$$\begin{array}{ccccc} A & * & P & = & AP \\ \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix} & * & \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & 0 \end{bmatrix} & = & \begin{bmatrix} 3 & 1 & 2 \\ 6 & 4 & 5 \\ 9 & 7 & 8 \end{bmatrix} \end{array} \quad (4)$$

Say that we reorder the rows of  $A$  by forming  $PA$ . Obviously, we can unreorder the rows by forming  $P^{-1}PA$ . Because permutation matrices are orthogonal, their inverses are equal to their transpose. Thus the inverse of the permutation matrix  $(0, 1, 0 \backslash 0, 0, 1 \backslash 1, 0, 0)$  we have been using is  $(0, 0, 1 \backslash 1, 0, 0 \backslash 0, 1, 0)$ . For instance, taking our results from (3)

$$\begin{array}{ccccc}
 P' & * & PA & = & A \\
 \begin{bmatrix} 0 & 0 & 1 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \end{bmatrix} & * & \begin{bmatrix} 4 & 5 & 6 \\ 7 & 8 & 9 \\ 1 & 2 & 3 \end{bmatrix} & = & \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}
 \end{array} \tag{3'}$$

Allow us to summarize:

1. A permutation matrix  $P$  is a square matrix whose rows are a permutation of the identity matrix.
2.  $PA$  = a row permutation of  $A$ .
3.  $AP$  = a column permutation of  $A$ .
4. The inverse permutation is given by  $P'$ .
5.  $P'PA = A$ .
6.  $APP' = A$ .

## How permutation matrices arise

Some of Mata's matrix functions implicitly permute the rows (or columns) of a matrix. For instance, the LU decomposition of matrix  $A$  is defined as

$$A = LU$$

where  $L$  is lower triangular and  $U$  is upper triangular. For any matrix  $A$ , one can solve for  $L$  and  $U$ , and Mata has a function that will do that (see [M-5] `lud()`). However, Mata's function does not solve the problem as stated. Instead, it solves

$$P'A = LU$$

where  $P'$  is a permutation matrix that Mata makes up! Just to be clear; Mata's function solves for  $L$  and  $U$ , but for a row permutation of  $A$ , not  $A$  itself, although the function does tell you what permutation it chose (the function returns  $L$ ,  $U$ , and  $P$ ). The function permutes the rows because, that way, it can produce a more accurate answer.

You will sometimes read that a function engages in pivoting. What that means is that, rather than solving the problem for the matrix as given, it solves the problem for a permutation of the original matrix, and the function chooses the permutation in a way to minimize numerical roundoff error. Functions that do this invariably return the permutation matrix along with the other results, because you are going to need it.

For instance, one use of LU decomposition is to calculate inverses. If  $A = LU$  then  $A^{-1} = U^{-1}L^{-1}$ . Calculating the inverses of triangular matrices is an easy problem, so one recipe for calculating inverses is

1. decompose  $A$  into  $L$  and  $U$ ,
2. calculate  $U^{-1}$ ,
3. calculate  $L^{-1}$ , and
4. multiply the results together.

That would be the solution except that the LU decomposition function does not decompose  $A$  into  $L$  and  $U$ ; it decomposes  $P'A$ , although the function does tell us  $P$ . Thus we can write,

$$\begin{aligned} P'A &= LU \\ A &= PLU && \text{(remember } P'^{-1} = P) \\ A^{-1} &= U^{-1}L^{-1}P' \end{aligned}$$

Thus the solution to our problem is to use the  $U$  and  $L$  just as we planned—calculate  $U^{-1}L^{-1}$ —and then make a column permutation of that, which we can do by postmultiplying by  $P'$ .

There is, however, a detail that we have yet to reveal to you: Mata's LU decomposition function does not return  $P$ , the permutation matrix. It instead returns  $p$ , a permutation vector equivalent to  $P$ , and so the last step—forming the column permutation by postmultiplying by  $P'$ —is done differently. That is the subject of the next section.

Permutation vectors are more efficient than permutation matrices, but you are going to discover that they are not as mathematically transparent. Thus when working with a function that returns a permutation vector—when working with a function that permutes the rows or columns—think in terms of permutation matrices and then translate back into permutation vectors.

## Permutation vectors

Permutation vectors are used with Mata's subscripting operator, so before explaining permutation vectors, let's understand subscripting.

Not surprisingly, Mata allows subscripting. Given matrix

$$A = \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}$$

Mata understands that

$$A[2,3] = 6$$

Mata also understands that if one or the other subscript is specified as `.` (missing value), the entire column or row is to be selected:

$$A[.,3] = \begin{bmatrix} 3 \\ 6 \\ 9 \end{bmatrix}$$

$$A[2,.] = \begin{bmatrix} 4 & 5 & 6 \end{bmatrix}$$

Mata also understands that if a vector is specified for either subscript

$$A \left[ \begin{bmatrix} 2 \\ 3 \\ 2 \end{bmatrix}, . \right] = \begin{bmatrix} 4 & 5 & 6 \\ 7 & 8 & 9 \\ 4 & 5 & 6 \end{bmatrix}$$

In Mata, we would actually write the above as  $A[(2\backslash 3\backslash 2),.]$ , and Mata would understand that we want the matrix made up of rows 2, 3, and 2 of  $A$ , all columns. Similarly, we can request all rows, columns 2, 3, and 2:

$$A[.,(2,3,2)] = \begin{bmatrix} 2 & 3 & 2 \\ 5 & 6 & 5 \\ 8 & 9 & 8 \end{bmatrix}$$

In the above, we wrote (2,3,2) as a row vector because it seems more logical that way, but we could just as well have written  $A[.,(2\backslash 3\backslash 2)]$ . In subscripting, Mata does not care whether the vectors are rows or columns.

In any case, we can use a vector of indices inside Mata's subscripts to select rows and columns of a matrix, and that means we can permute them. All that is required is that the vector we specify contain a permutation of the integers 1 through  $n$  because, otherwise, we would repeat some rows or columns and omit others.

A permutation vector is an  $n \times 1$  or  $1 \times n$  vector containing a permutation of the integers 1 through  $n$ . For example, the permutation vector equivalent to the permutation matrix

$$P = \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & 0 \end{bmatrix}$$

is

$$p = \begin{bmatrix} 2 \\ 3 \\ 1 \end{bmatrix}$$

$p$  can be used with subscripting to permute the rows of  $A$

$$A[p,.] = \begin{bmatrix} 4 & 5 & 6 \\ 7 & 8 & 9 \\ 1 & 2 & 3 \end{bmatrix}$$

and similarly,  $A[.,p]$  would permute the columns.

Also subscripting can be used on the left-hand side of the equal-sign assignment operator. So far, we have assumed that the subscripts are on the right-hand side of the assignment operator, such as

$$B = A[p,.]$$

We have learned that if  $p = (2\backslash 3\backslash 1)$  (or  $p = (2,3,1)$ ), the result is to copy the second row of  $A$  to the first row of  $B$ , the third row of  $A$  to the second row of  $B$ , and the first row of  $A$  to the third row of  $B$ . Coding

$$B[p,.] = A$$

does the inverse: it copies the first row of  $A$  to the second row of  $B$ , the second row of  $A$  to the third row of  $B$ , and the third row of  $A$  to the first row of  $B$ .  $B[p,.] = A$  really is the inverse of  $C = A[p,.]$  in that, if we code

$$\begin{aligned} C &= A[p,.] \\ B[p,.] &= C \end{aligned}$$

$B$  will be equal to  $C$ , and if we code

```
C[p,.] = A
B = C[p,.]
```

$B$  will also be equal to  $C$ .

There is, however, one pitfall that you must watch for when using subscripts on the left-hand side: the matrix on the left-hand side must already exist and it must be of the appropriate (here same) dimension. Thus when performing the inverse copy, it is common to code

```
B = C
B[p,.] = C
```

The first line is not unnecessary; it is what ensures that  $B$  exists and is of the proper dimension, although we could just as well code

```
B = J(rows(C), cols(C), .)
B[p,.] = C
```

The first construction is preferred because it ensures that  $B$  is of the same type as  $C$ . If you really like the second form, you should code

```
B = J(rows(C), cols(C), missingof(C))
B[p,.] = C
```

Going back to the preferred code

```
B = C
B[p,.] = C
```

some programmers combine it into one statement:

```
(B=C)[p,.] = C
```

Also Mata provides an `invorder( $p$ )` (see [M-5] `invorder()`) that will return an inverted  $p$  appropriate for use on the right-hand side, so you can also code

```
B = C[invorder(p),.]
```

## Also see

[M-5] `invorder()` — Permutation vector manipulation

[M-1] `intro` — Introduction and advice

# Title

[M-1] **returnedargs** — Function arguments used to return results

## Syntax

$y = f(x, \dots)$       (function returns result the usual way)  
 $g(x, \dots, y)$       (function returns result in argument  $y$ )

## Description

Most Mata functions leave their arguments unchanged and return a result:

```
: y = f(x, ...)
```

Some Mata functions, however, return nothing and instead return results in one or more arguments:

```
: g(x, ..., y)
```

If you use such functions interactively and the arguments that are to receive results are not already defined ( $y$  in the above example), you will get a variable-not-found error. The solution is to define the arguments to contain something—anything—before calling the function:

```
: y = .  
: g(x, ..., y)
```

You can combine this into one statement:

```
: g(x, ..., y=.)
```

## Remarks

`sqrt( $a$ )`—see [M-5] **sqrt()**—calculates the (element-by-element) square root of  $a$  and returns the result:

```
: x = 4  
: y = sqrt(x)  
: y           // y now contains 2  
2  
: x           // x is unchanged  
4
```

Most functions work like `sqrt()`, although many take more than one argument.

On the other hand, `polydiv( $c_a$ ,  $c_b$ ,  $c_q$ ,  $c_r$ )`—see [M-5] **polyeval()**—takes the polynomial stored in  $c_a$  and the polynomial stored in  $c_b$  and divides them. It returns the quotient in the third argument ( $c_q$ ) and the remainder in the fourth ( $c_r$ ).  $c_a$  and  $c_b$  are left unchanged. The function itself returns nothing:

```
: A = (1,2,3)  
: B = (0,1)  
: polydiv(A, B, Q, R)
```



```

: Q                                // Q has been redefined
      1      2
1  

|   |   |
|---|---|
| 2 | 3 |
|---|---|



: R                                // as has R
      1

: A                                // while A and B are unchanged
      1      2      3
1  

|   |   |   |
|---|---|---|
| 1 | 2 | 3 |
|---|---|---|



: B
      1      2
1  

|   |   |
|---|---|
| 0 | 1 |
|---|---|


```

As another example, `st_view(V, i, j)`—see [M-5] `st_view()`—creates a view onto the Stata dataset. Views are like matrices but consume less memory. Arguments *i* and *j* specify the observations and variables to be selected. Rather than returning the matrix, however, the result is returned in the first argument (*V*).

```

: st_view(V, (1\5), ("mpg", "weight"))
: V
      1      2
1  

|    |      |
|----|------|
| 22 | 2930 |
| 15 | 4080 |


2

```

If you try to use these functions interactively, you will probably get an error:

```

: polydiv(A, B, Q, R)
      <istmt>: 3499 Q not found
r(3499);
: st_view(V, (1\5), ("mpg", "weight"))
      <istmt>: 3499 V not found
r(3499);

```

Arguments must be defined before they are used, even if their only purpose is to receive a newly calculated result. In such cases, it does not matter how the argument is defined because its contents will be replaced. Easiest is to fill in a missing value:

```

: Q = .
: R = .
: polydiv(A, B, Q, R)
: V = .
: st_view(V, (1\5), ("mpg", "weight"))

```

You can also define the argument inside the function:

```

: polydiv(A, B, Q=., R=.)
: st_view(V=., (1\5), ("mpg", "weight"))

```

When you use functions like these inside a program, however, you need not worry about defining the arguments, because they are defined by virtue of appearing in your program:

```
function foo()
{
    ...
    polydiv(A, B, Q, R)
    st_view(V, (1\5), ("mpg", "weight"))
    ...
}
```

When Mata compiles your program, however, you may see warning messages:

```
: function foo()
> {
>   ...
>   polydiv(A, B, Q, R)
>   st_view(V, (1\5), ("mpg", "weight"))
>   ...
> }
note: variable Q may be used before set
note: variable R may be used before set
note: variable V may be used before set
```

If the warning messages bother you, either define the variables before they are used just as you would interactively or use `pragma` to suppress the warning messages; see [M-2] [pragma](#).

## Also see

[M-1] [intro](#) — Introduction and advice

# Title

[M-1] **source** — Viewing the source code

## Syntax

```
viewsource functionname.mata
```

## Description

Many Mata functions are written in Mata. `viewsource` will allow you to examine their source code.

## Remarks

Some Mata functions are implemented in C (they are part of Mata itself), and others are written in Mata.

`viewsource` allows you to look at the official source code written in Mata. Reviewing this code is a great way to learn Mata.

The official source code is stored in `.mata` files. To see the source code for `diag()` (see [M-5] [diag\(\)](#)), for instance, type

```
. viewsource diag.mata
```

You type this at Stata's dot prompt, not at Mata's colon prompt.

If a function is built in, such as `abs()` (see [M-5] [abs\(\)](#)), here is what will happen when you attempt to view the source code:

```
. viewsource abs.mata
file "abs.mata" not found
r(601);
```

You can verify that `abs()` is built in by using the `mata which` (see [M-3] [mata which](#)) command:

```
. mata: mata which abs()
abs():  built-in
```

`viewsource` can be also used to look at source code of user-written functions if the distribution included the source code (it might not).

## Also see

[P] [viewsource](#) — View source code

[M-1] [intro](#) — Introduction and advice

# Title

[M-1] tolerance — Use and specification of tolerances

## Syntax

*somefunction*(... , *real scalar tol* , ...)

where, concerning argument *tol*,

- optional     Argument *tol* is usually optional; not specifying *tol* is equivalent to specifying  $tol = 1$ .
- $tol > 0$      Specifying  $tol > 0$  specifies the amount by which the usual tolerance is to be multiplied:  $tol = 2$  means twice the usual tolerance;  $tol = 0.5$  means half the usual tolerance.
- $tol < 0$      Specifying  $tol < 0$  specifies the negative of the value to be used for the tolerance:  $tol = -1e-14$  means  $1e-14$  is to be used.
- $tol = 0$      Specifying  $tol = 0$  means all numbers are to be taken at face value, no matter how close to 0 they are. The single exception is when *tol* is applied to values that, mathematically, must be greater than or equal to zero. Then negative values (which arise from roundoff error) are treated as if they were zero.

The default tolerance is given by formula, such as

$eta = 1e-14$

or

$eta = \text{epsilon}(1)$      (see [M-5] [epsilon\(\)](#))

or

$eta = 1000 * \text{epsilon}(\text{trace}(\text{abs}(A)) / \text{rows}(A))$

Specifying  $tol > 0$  specifies a value to be used to multiply *eta*. Specifying  $tol < 0$  specifies that  $-tol$  be used in place of *eta*. Specifying  $tol = 0$  specifies that *eta* be set to 0.

The formula for *eta* and how *eta* is used are found under *Remarks*. For instance, the *Remarks* might say that *A* is declared to be singular if any diagonal element of *U* of its LU decomposition is less than or equal to *eta*.

## Description

The results provided by many of the numerical routines in Mata depend on tolerances. Mata provides default tolerances, but those can be overridden.

## Remarks

Remarks are presented under the following headings:

*The problem*  
*Absolute versus relative tolerances*  
*Specifying tolerances*

## The problem

In many formulas, zero is a special number in that, when the number arises, sometimes the result cannot be calculated or, other times, something special needs to be done.

The problem is that zero—0.00000000000—seldom arises in numerical calculation. Because of roundoff error, what would be zero were the calculation performed in infinite precision in fact is  $1.03948\text{e-}15$ , or  $-4.4376\text{e-}16$ , etc.

If one behaves as if these small numbers are exactly what they seem to be ( $1.03948\text{e-}15$  is taken to mean  $1.03948\text{e-}15$  and not zero), some formulas produce wildly inaccurate results; see [M-5] `lusolve()` for an example.

Thus routines use *tolerances*—preset numbers—to determine when a number is small enough to be considered to be zero.

The problem with tolerances is determining what they ought to be.

## Absolute versus relative tolerances

Tolerances come in two varieties: absolute and relative.

An absolute tolerance is a fixed number that is used to make direct comparisons. If the tolerance for a particular routine were  $1\text{e-}14$ , then  $8.99\text{e-}15$  in some calculation would be considered to be close enough to zero to act as if it were, in fact, zero, and  $1.000001\text{e-}14$  would be considered a valid, nonzero number.

But is  $1\text{e-}14$  small? The number may look small to you, but whether  $1\text{e-}14$  is small depends on what is being measured and the units in which it is measured. If all the numbers in a certain problem were around  $1\text{e-}12$ , you might suspect that  $1\text{e-}14$  is a reasonable number.

That leads to relative measures of tolerance. Rather than treating, say, a predetermined quantity as being so small as to be zero, one specifies a value (for example,  $1\text{e-}14$ ) multiplied by something and uses that as the definition of small.

Consider the following matrix:

$$\begin{bmatrix} 5.5\text{e-}15 & 1.2\text{e-}16 \\ 1.3\text{e-}16 & 6.4\text{e-}15 \end{bmatrix}$$

What is the rank of the matrix? One way to answer that question would be to take the LU decomposition of the matrix and then count the number of diagonal elements of  $U$  that are greater than zero. Here, however, we will just look at the matrix.

The absolutist view is that the matrix is full of roundoff error and that the matrix is really indistinguishable from the matrix

$$\begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix}$$

The matrix has rank 0. The relativist view is that the matrix has rank 2 because, other than a scale factor of  $1\text{e-}16$ , the matrix is indistinguishable from

$$\begin{bmatrix} 55.0 & 1.2 \\ 1.3 & 64.0 \end{bmatrix}$$

There is no way this question can be answered until someone tells you how the matrix arose and the units in which it is measured.

Nevertheless, most Mata routines would (by default) adopt the relativist view: the matrix is of full rank. That is because most Mata routines are implemented using relative measures of tolerance, chosen because Mata routines are mostly used by people performing statistics, who tend to make calculations such as  $X'X$  and  $X'Z$  on data matrices, and those resulting matrices can contain very large numbers. Such a matrix might contain

$$\begin{bmatrix} 5.5\text{e}+14 & 1.2\text{e}+12 \\ 1.3\text{e}+13 & 2.4\text{e}+13 \end{bmatrix}$$

Given a matrix with such large elements, one is tempted to change one's view as to what is small. Calculate the rank of the following matrix:

$$\begin{bmatrix} 5.5\text{e}+14 & 1.2\text{e}+12 & 1.5\text{e}-04 \\ 1.3\text{e}+13 & 2.4\text{e}+13 & 2.8\text{e}-05 \\ 1.3\text{e}-04 & 2.4\text{e}-05 & 8.7\text{e}-05 \end{bmatrix}$$

This time, we will do the problem correctly: we will take the LU decomposition and count the number of nonzero entries along the diagonal of  $U$ . For the above matrix, the diagonal of  $U$  turns out to be (5.5e+14, 2.4e+13, 0.000087).

An absolutist would tell you that the matrix is of full rank; the smallest number along the diagonal of  $U$  is 0.000087 (8.7e-5), and that is still a respectable number, at least when compared with computer precision, which is about 2.22e-16 (see [M-5] [epsilon\(\)](#)).

Most Mata routines would tell you that the matrix has rank 2. Numbers such as 0.000087 may seem respectable when compared with machine precision, but 0.000087 is, relatively speaking, a very small number, being about 4.6e-19 relative to the average value of the diagonal elements.

## Specifying tolerances

Most Mata routines use relative tolerances, but there is no rule. You must read the documentation for the function you are using.

When the tolerance entry for a function directs you here, [M-1] [tolerance](#), then the tolerance works as summarized under [Syntax](#) above. Specify a positive number, and that number multiplies the default; specify a negative number, and the corresponding positive number is used in place of the default.

## Also see

[M-5] [epsilon\(\)](#) — Unit roundoff error (machine precision)

[M-5] [solve\\_tol\(\)](#) — Tolerance used by solvers and inverters

[M-1] [intro](#) — Introduction and advice

## **[M-2] Language definition**





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## Description

This section defines the Mata programming language.

## Remarks

[M-2] [syntax](#) provides an overview, dense and brief, and the other sections expand on it.

Also see [M-1] [intro](#) for an introduction to Mata.

## Also see

[M-0] [intro](#) — Introduction to the Mata manual

# Title

[M-2] **break** — Break out of for, while, or do loop

## Syntax

```
for, while, or do {  
    ...  
    if (...) {  
        ...  
        break  
    }  
}  
stmt                               ← break jumps here  
...
```

## Description

**break** exits the innermost **for**, **while**, or **do** loop. Execution continues with the statement immediately following the close of the loop, just as if the loop had terminated normally.

**break** nearly always occurs following an **if**.

## Remarks

In the following code,

```
for (i=1; i<=rows(A); i++) {  
    for (j=1; j<=cols(A); j++) {  
        ...  
        if (A[i,j]==0) break  
    }  
    printf("j = %g\n", j)  
}
```

the **break** statement will be executed if any element of  $A[i,j]$  is zero. Assume that the statement is executed for  $i=2$  and  $j=3$ . Execution will continue with the **printf()** statement, which is to say, the  $j$  loop will be canceled but the  $i$  loop will continue. The value of  $j$  upon exiting the loop will be 3; when you break out of the loop, the  $j++$  is not executed.

## Also see

[M-2] **do** — do ... while (exp)

[M-2] **for** — for (exp1; exp2; exp3) stmt

[M-2] **while** — while (exp) stmt

[M-2] **continue** — Continue with next iteration of for, while, or do loop

[M-2] **intro** — Language definition

## Syntax

```
class classname [extends classname] {  
    declaration(s)  
}
```

Syntax is presented under the following headings:

[Introduction](#)

[Example](#)

[Declaration of member variables](#)

[Declaration and definition of methods \(member functions\)](#)

[Default exposure in declarations](#)

[Description](#) and [Remarks](#) follow that.

## Introduction

Stata's two programming languages, ado and Mata, each support object-oriented programming. This manual entry explains object-oriented programming in Mata. Most users interested in object-oriented programming will wish to program in Mata. See [\[P\] class](#) to learn about object-oriented programming in ado.

## Example

The following example is explained in detail in [Description](#).

```
class coord {  
    real scalar    x, y  
    real scalar    length(), angle()  
}  
real scalar coord::length()  
{  
    return(sqrt(x^2 + y^2))  
}  
real scalar coord::angle()  
{  
    return(atan2(y, x)*360/(2*pi()))  
}  
class rotated_coord extends coord {  
    real scalar    theta  
    real scalar    angle()  
    void          new()  
}
```

```

real scalar rotated_coord::angle()
{
    return(super.angle() - theta)
}
void rotated_coord::new()
{
    theta = 0
}

```

One could use the class interactively:

```

: b = rotated_coord()
: b.x = b.y = 1
: b.angle()                // displayed will be 45
: b.theta = 30
: b.angle()                // displayed will be 15

```

Note that executing the class as if it were a function creates an instance of the class. When using the class inside other functions, it is not necessary to create the instance explicitly as long as you declare the member instance variable to be a scalar:

```

void myfunc()
{
    class rotated_coord scalar    b
    b.x = b.y = 1
    b.angle()                    // displayed will be 45
    b.theta = 30
    b.angle()                    // displayed will be 15
}

```

## Declaration of member variables

Declarations are of the form

```
[exposure] [static] [final] matatype name [, name [, ...]]
```

where

```

exposure := {public|protected|private}
matatype := {eltype orgtype | eltype | orgtype}
eltype :=  transmorphic      orgtype := matrix
           numeric            vector
           real               rowvector
           complex            colvector
           string             scalar
           pointer
           class classname
           struct structname

```

For example,

```
class S {
    real matrix                M
    private real scalar        type
    static real scalar         count
    class coord scalar         c
}
```

## Declaration and definition of methods (member functions)

Declarations are of the form

```
[exposure] [static] [final] [virtual]
matatype name() [, name() [...]]
```

For example,

```
class T {
    ...
    real matrix                inverse()
    protected real scalar      type()
    class coord scalar         c()
}
```

Note that function arguments, even if allowed, are not declared.

Member functions (methods) and member variables may share the same names and no special meaning is attached to the fact. `type` and `c` below are variables, and `type()` and `c()` are functions:

```
class U {
    real matrix                M
    private real scalar        type
    static real scalar         count
    class coord scalar         c

    real matrix                inverse()
    protected real scalar      type()
    class coord scalar         c()
}
```

Member functions are defined separately, after the class is defined. For example,

```
class V {
    real matrix                M
    private real scalar        type
    static real scalar         count
    class coord scalar         c

    real matrix                inverse()
    protected real scalar      type()
    class coord scalar         c()
}
```

```

real matrix V::inverse(...)
{
    ...
}
real scalar V::type(...)
{
    ...
}
class coord scalar V::c(...)
{
    ...
}

```

When you define member functions, they must be of the same *matatype* as they were previously declared to be, but you omit *exposure* (as well as `static`, `final`, and `virtual`).

## Default exposure in declarations

Variables and functions are `public` unless explicitly declared otherwise. (They are also not `static`, not `final`, and not `virtual`, but that is not part of exposure and so has nothing to do with this subsection.)

You may use any of the exposure modifiers `public`, `protected`, and `private`, followed by a colon, to create blocks with a different default:

```

class V {
    public:
        real matrix          M
        static real scalar   count
        class coord scalar   c
        real matrix          inverse()
        class coord scalar   c()
    private:
        real scalar          type
    protected:
        real scalar          type()
}

```

## Description

`class` provides object-oriented programming, also known as class programming, to Mata.

A class is a set of variables or related functions (methods) (or both) tied together under one name. Classes may be derived from other classes according to [inheritance](#).

Let's look at the details of the example from the first page of this entry (under the heading [Example](#)):

1. First, we created a class called `coord`. When we coded

```

class coord {
    real scalar   x, y
    real scalar   length(), angle()
}

```

we specified that each element of a `coord` stores two real values, which we called `x` and `y`. `coord` also contains two functions, which we called `length()` and `angle()`. `length()` and `angle()` are functions because of the open and close parentheses at the end of the names, and `x` and `y` are variables because of the absence of parentheses. In the jargon, `x` and `y` are called member variables, and `length()` and `angle()` are called member functions.

The above, called the class's definition, defines a blueprint for the type `coord`.

A variable that is of type `coord` is called an instance of a `coord`. Say variables `b` and `c` are instances of `coord`, although we have not yet explained how you might arrange that. Then `b.x` and `b.y` would be `b`'s values of `x` and `y`, and `c.x` and `c.y` would be `c`'s values. We could run the functions on the values in `b` by coding `b.length()` and `b.angle()`, or on the values in `c` by coding `c.length()` and `c.angle()`.

- Next we defined `coord`'s `length()` and `angle()` functions. The definitions were

```
real scalar coord::length()
{
    return(sqrt(x^2 + y^2))
}
real scalar coord::angle()
{
    return(atan2(y, x)*360/(2*pi()))
}
```

These functions are similar to regular Mata functions. These functions do not happen to take any arguments, but that is not a requirement. It often happens that member functions do not require arguments because the functions are defined in terms of the class and its concepts. That is exactly what is occurring here. The first function says that when someone has an instance of a `coord` and they ask for its length, return the square root of the sum of the `x` and `y` values, individually squared. The second function says that when someone asks for the angle, return the arctangent of `y` and `x`, multiplied by `360/(2*pi())` to convert the result from radians to degrees.

- Next we defined another new concept, a `rotated_coord`, by extending (inheriting from) class `coord`:

```
class rotated_coord extends coord {
    real scalar    theta
    real scalar    angle()
    void           new()
}
```

So think of a `coord` and read the above as the additions. In addition to `x` and `y` of a regular `coord`, a `rotated_coord` has new variable `theta`. We also declared that we were adding two functions, `angle()` and `new()`. But wait, `angle()` already existed! What we are actually saying by explicitly mentioning `angle()` is that we are going to change the definition of `angle()`. Function `new()` is indeed new.

Notice that we did not mention previously existing function `length()`. Our silence indicates that the concept of `length()` remains unchanged.

So what is a `rotated_coord`? It is a `coord` with the addition of `theta`, a redefinition of how `angle()` is calculated, and the addition of a function called `new()`.



This is an example of inheritance. `class rotated_coord` is an extension of `class coord`. In object-oriented programming, we would say this as “`class rotated_coord` inherits from `class coord`”. A class that inherits from another class is known as a “subclass”.

4. Next we defined our replacement and new functions. The definitions are

```
real scalar rotated_coord::angle()
{
    return(super.angle() - theta)
}
void rotated_coord::new()
{
    theta = 0
}
```

Concerning `angle()`, we stated that it is calculated by taking the result of `super.angle()` and subtracting `theta` from it. `super.angle()` is how one refers to the parent’s definition of a function. If you are unfamiliar with object-oriented programming, parent may seem like an odd word in this context. We are inheriting concepts from `coord` to define `rotated_coord`, and in that sense, `coord` is the parent concept. Anyway, the new definition of an angle is the old definition, minus `theta`, the angle of rotation.

`new()` is a special function and is given a special name in the sense that the name `new()` is reserved. The `new()` function, if it exists, is a function that is called automatically whenever a new instance of the class is created. Our `new()` function says that when a new instance of a `rotated_coord` is created, initialize the instance’s `theta` to 0.

Well, that seems like a good idea. But we did not have a `new()` function when we defined our previous class, `coord`. Did we forget something? Maybe. When you do not specify a `new()` function—when you do not specify how variables are to be initialized—they are initialized in the usual Mata way: missing values. `x` and `y` will be initialized to contain missing. Given our `new()` function for `rotated_coord`, however, `theta` will be initialized to 0.

`new()` is called a “constructor”, because it is used to construct, or initialize, the class when a new instance of the class is created.

And that completes the definition of our two, related classes.

There are two ways to create instances of a `coord` or a `rotated_coord`. One is mostly for interactive use and the other for programming use.

If you interactively type `a=coord()` (note the parentheses), you will create a `coord` and store it in `a`. If you interactively type `b=rotated_coord()`, you will create a `rotated_coord` and store it in `b`. In the first example, typing `b=rotated_coord()` is exactly what we chose to do:

```
: b = rotated_coord()
```

Recall that a `rotated_coord` contains an `x`, `y`, and `theta`. At this stage, `x` and `y` equal missing, and `theta` is 0. In the example, we set `b`’s `x` and `y` values to 1, and then asked for the resulting `angle()`:

```
: b.x = b.y = 1
: b.angle()
45
```

`b-dot-x` is how one refers to `b`'s value of `x`. One can use `b.x` (and `b.y`) just as one would use any real scalar variable in Mata.

If we reset `b`'s `theta` to be 30 degrees, then `b`'s angle ought to change to being 15 degrees. That is exactly what happens:

```
: b.theta = 30
: b.angle()
15
```

`b-dot-angle()` is how one specifies that member function `angle()` is to be run on instance `b`. Now you know why member functions so seldom have additional arguments: they are, in effect, passed an instance of a class and so have access to all the values of member variables of that class. We repeat, however, that a member function could take additional arguments. Had we coded

```
real scalar rotated_coord::angle(real scalar base)
{
    return(super.angle() - theta - base)
}
```

then `angle()` would have taken an argument and returned the result measured from `base`.

The difference between using `rotated_coord` interactively and using it inside a Mata program is that if we declare a variable (say, `b`) to be a class `rotated_coord` scalar, with the emphasis on scalar, then we do not need to bother coding `b=rotated_coord()` to fill in `b` initially. Coding `class rotated_coord scalar b` implies that `b` needs to be initialized because it is a scalar, and so that happens automatically. It would not hurt if we also coded `b=rotated_coord()`, but it would be a waste of our time and of the computer's once it got around to executing our program.

Now let's show you something we did not show in the first example. Remember when we defined `length()` for a `coord`? Remember how we did not define `length()` for a `rotated_coord`? Remember how we did not even mention `length()`? Even so, `length()` is a concept of a `rotated_coord`, because part of the definition of `rotated_coord` was inherited from `coord`, and that happened because when we declared `rotated_coord`, we said

```
class rotated_coord extends coord
```

The inheritance happened because we said `extends`. Let's test that `length()` works with our `rotated_coord` class instance, `b`:

```
: b.length()
1.414213562
```

In the above, inheritance is what saved us from having to write additional, repeated code.

Let's review. First, we defined a `coord`. From that, we defined a `rotated_coord`. You might now define `translated_and_rotated_coord` using `rotated_coord` as a starting point. It would not be difficult.

Classes have lots of properties, features, and details, but it is the property of inheritance that is at the heart of object-oriented programming.

## Remarks

Remarks are presented under the following headings:

- Notation and jargon*
- Declaring and defining a class*
- Saving classes in files*
- Workflow recommendation*
- When you need to recompile*
- Obtaining instances of a class*
- Constructors and destructors*
- Setting member variable and member function exposure*
- Making a member final*
- Making a member static*
- Virtual functions*
- Referring to the current class using this*
- Using super to access the parent's concept*
- Casting back to a parent*
- Accessing external functions from member functions*
- Pointers to classes*

## Notation and jargon

*:: (double colon)*

The double-colon notation is used as a shorthand in documentation to indicate that a variable or function is a [member](#) of a class and, in two cases, the double-colon notation is also syntax that is understood by Mata.

`S::s` indicates that the variable `s` is a member of class `S`. `S::s` is documentation shorthand.

`S::f()` indicates that function `f()` is a member of class `S`. `S::f()` is documentation shorthand. `S::f()` is something Mata itself understands in two cases:

1. Notation `S::f()` is used when defining member functions.
2. Notation `S::f()` can be used as a way of calling [static](#) member functions.

*be an instance of*

When we write “let `s` be an instance of `S`” we are saying that `s` is an [instance](#) of class `S`, or equivalently, that in some Mata code, `s` is declared as a `class S scalar`.

*class definition, class declaration*

A class definition or declaration is the definition of a class, such as the following:

```
class S {
    private real scalar    a, b
    real scalar            f(), g()
}
```

Note well that the declaration of a Mata variable to be of type `class S`, such as the line `class S scalar s` in

```
void myfunction()
{
    class S scalar s
    ...
}
```

is not a class definition. It is a declaration of an [instance](#) of a class.

*class instance, instance, class A instance*

A class instance is a variable defined according to a class definition. In the code

```
void myfunction()
{
    class S scalar    s
    real scalar      b
    ...
}
```

`s` is a class instance, or, more formally, an instance of class `S`. The term “instance” can be used with all element types, not just with classes. Thus `b` is an instance of a `real`. The term “instance” is more often used with classes and structures, however, because one needs to distinguish definitions of variables containing classes or structures from the definitions of the classes and structures themselves.

*inheritance, extends, subclass, parent, child*

Inheritance is the property of one class definition using the variables and functions of another just as if they were its own. When a class does this, it is said to extend the other class. `T` extending `S` means the same thing as `T` inheriting from `S`. `T` is also said to be a subclass of `S`.

Consider the following definitions:

```
class S {
    real scalar    a, b
    real scalar    f()
}
class T extends S {
    real scalar    c
    real scalar    g()
}
```

Let `s` be an instance of `S` and `t` be an instance of `T`. It is hardly surprising that `s.a`, `s.b`, `s.f()`, `t.c`, and `t.g()` exist, because each is explicitly declared. It is because of inheritance that `t.a`, `t.b`, and `t.f()` also exist, and they exist with no additional code being written.

If `U` extends `T` extends `S`, then `U` is said to be a child of `T` and to be a child of `S`, although formally `U` is the grandchild of `S`. Similarly, both `S` and `T` are said to be parents of `U`, although formally `S` is the grandparent of `U`. It is usually sufficient to label the relationship parent/child without going into details.

*external functions*

An external function is a regular function, such as `sqrt()`, `sin()`, and `myfcn()`, defined outside of the class and, as a matter of fact, outside of all classes. The function could be a function provided by Mata (such as `sqrt()` or `sin()`) or it could be a function you have written, such as `myfcn()`.

An issue arises when calling external functions from inside the code of a member function. When coding a [member function](#), references such as `sqrt()`, `sin()`, and `myfcn()` are assumed to be references to the class’s member functions if a member function of the name exists. If one wants to ensure that the external-function interpretation is made, one codes `::sqrt()`, `::sin()`, and `::myfcn()`. See [Accessing external functions from member functions](#) below.

*member, member variable, member function, method*

A variable or function declared within a class is said to be a member variable, or member function, of the class. Member functions are also known as methods. In what follows, let `ex1` be an instance of `S`, and assume class `S` contains member function `f()` and member variable `v`.

When member variables and functions are used inside member functions, they can simply be referred to by their names, `v` and `f()`. Thus, if we were writing the code for `S::g()`, we could code

```
real scalar S::g()
{
    return(f()*v)
}
```

When member variables and functions are used outside of member functions, which is to say, are used in regular functions, the references must be prefixed with a class instance and a period. Thus one codes

```
real scalar myg(class S scalar ex1)
{
    return(ex1.f()*ex1.v)
}
```

*variable and method overriding*

Generically, a second variable or function overrides a first variable or function when they share the same name and the second causes the first to be hidden, which causes the second variable to be accessed or the second function to be executed in preference to the first. This arises in two ways in Mata's implementation of classes.

In the first way, a variable or function in a parent class is said to be overridden if a child defines a variable or function of the same name. For instance, let `U` extend `T` extend `S`, and assume `S::f()` and `T::f()` are defined. By the rules of [inheritance](#), instances of `U` and `T` that call `f()` will cause `T::f()` to execute. Instances of `S` will cause `S::f()` to be executed. Here `S.f()` is said to be overridden by `T.f()`. Because `T::f()` will usually be implemented in terms of `S::f()`, `T::f()` will find it necessary to call `S::f()`. This is done by using the `super` prefix; see [Using super to access the parent's concept](#).

The second way has to do with stack variables having precedence over member variables in member functions. For example,

```
class S {
    real scalar    a, b
    void          f()
    ...
}
void S::f()
{
    real scalar    a
    a = 0
    b = 0
}
```

Let `s` be an instance of `S`. Then execution of `s.f()` sets `s.b` to be 0; it does not change `s.a`, although it was probably the programmer's intent to set `s.a` to zero, too. Because the programmer

declared a variable named `a` within the program, however, the program’s variable took precedence over the member variable `a`. One solution to this problem would be to change the `a = 0` line to read `this.a = 0`; see [Referring to the current class using this](#).

## Declaring and defining a class

Let’s say we declared a class by executing the code below:

```
class coord {
    real scalar    x, y
    real scalar    length(), angle()
}
```

At this point, class `coord` is said to be declared but not yet fully defined because the code for its member functions `length()` and `angle()` has not yet been entered. Even so, the class is partially functional. In particular,

- 1. The class will work. Obviously, if an attempt to execute `length()` or `angle()` is made, an error message will be issued.
- 2. The class definition can be saved in an `.mo` or `.mlib` file for use later, with the same proviso as in 1).

Member functions of the class are defined in the same way regular Mata functions are defined but the name of the function is specified as `classname::functionname()`.

```
real scalar coord::length()
{
    return(sqrt(x^2 + y^2))
}
real scalar coord::angle()
{
    return(atan2(y, x)*360/(2*pi()))
}
```

The other difference is that member functions have direct access to the class’s member variables and functions. `x` and `y` in the above are the `x` and `y` values from an instance of class `coord`.

Class `coord` is now fully defined.

## Saving classes in files

The notation `coord()`—classname-open parenthesis-close parenthesis—is used to refer to the entire class definition by Mata’s interactive commands such as [mata describe](#), [mata mosave](#), and [mata mlib](#).

`mata describe coord()` would show

: mata describe coord()			
# bytes	type		name and extent
344	classdef scalar		coord()
176	real scalar		::angle()
136	real scalar		::length()

The entire class definition—the compiled `coord()` and all its compiled member functions—can be stored in a `.mo` file by typing

```
: mata mosave coord()
```

The entire class definition could be stored in an already existing `.mlib` library, `lpersonal`, by typing

```
: mata mlib add lpersonal coord()
```

When saving a class definition, both commands allow the additional option `complete`. The option specifies that the class is to be saved only if the class definition is complete and that, otherwise, an error message is to be issued.

```
: mata mlib add lpersonal coord(), complete
```

## Workflow recommendation

Our recommendation is that the source code for classes be kept in separate files, and that the files include the complete definition. At StataCorp, we would save `coord` in file `coord.mata`:

---

```
begin coord.mata
*! version 1.0.0 class coord
version 12
mata:
class coord {
    real scalar    x, y
    real scalar    length(), angle()
}
real scalar coord::length()
{
    return(sqrt(x^2 + y^2))
}
real scalar coord::angle()
{
    return(atan2(y, x)*360/(2*pi()))
}
end
```

---

end coord.mata

Note that the file does not clear Mata, nor does it save the class in a `.mo` or `.mlib` file; the file merely records the source code. With this file, to save `coord` in a `.mo` file, we need only to type

```
. clear mata
. do coord.mata
. mata mata mosave coord(), replace
```

(Note that although file `coord.mata` does not have the extension `.do`, we can execute it just like we would any other do-file by using the `do` command.) Actually, we would put those three lines in yet another do-file called, perhaps, `cr_coord.do`.

We similarly use files for creating libraries, such as

---

```

begin cr_lourlib.do
    version 12
    clear mata
    do coord.mata
    do anotherfile.mata
    do yetanotherfile.mata
    /* etc. */

    mata:
    mata mlib create lourlib, replace
    mata mlib add *()
    end
end cr_lourlib.do

```

---

With the above, it is easy to rebuild libraries after updating code.

## When you need to recompile

If you change a class declaration, you need to recompile all programs that use the class. That includes other class declarations that inherit from the class. For instance, in the opening example, if you change anything in the `coord` declaration,

```

class coord {
    real scalar    x, y
    real scalar    length(), angle()
}

```

even if the change is so minor as to reverse the order of `x` and `y`, or to add a new variable or function, you must not only recompile all of `coord`'s member functions, you must also recompile all functions that use `coord`, and you must recompile `rotated_coord` as well, because `rotated_coord` inherited from `coord`.

You must do this because Mata seriously compiles your code. The Mata compiler deconstructs all use of classes and substitutes low-level, execution-time-efficient constructs that record the address of every member variable and member function. The advantage is that you do not have to sacrifice run-time efficiency to have more readable and easily maintainable code. The disadvantage is that you must recompile when you make changes in the definition.

You do not need to recompile outside functions that use a class if you only change the code of member functions. You do need to recompile if you add or remove member variables or member functions.

To minimize the need for recompiling, especially if you are distributing your code to others physically removed from you, you may want to adopt the pass-through approach.

Assume you have written large, wonderful systems in Mata that all hinge on object-oriented programming. One class inherits from another and that one from another, but the one class you need to tell your users about is class `wonderful`. Rather than doing that, however, merely tell your users that they should declare a `transmorphic` named `wonderfulhandle`—and then just have them pass `wonderfulhandle` around. They begin using your system by making an initialization call:

```
wonderfulhandle = wonderful_init()
```



After that, they use regular functions you provide that require `wonderful` as an argument. From their perspective, `wonderfulhandle` is a mystery. From your perspective, `wonderful_init()` returned an instance of a class `wonderful`, and the other functions you provided receive an instance of a `wonderful`. Sadly, this means that you cannot reveal to your users the beauty of your underlying class system, but it also means that they will not have to recompile their programs when you distribute an update. If the Mata compiler can compile their code knowing only that `wonderfulhandle` is a transmorphic, then it is certain their code does not depend on how `wonderfulhandle` is constructed.

## Obtaining instances of a class

Declaring a class, such as

```
class coord {
    real scalar    x, y
    real scalar    length(), angle()
}
```

in fact creates a function, `coord()`. Function `coord()` will create and return instances of the class:

- `coord()`—`coord()` without arguments—returns a scalar instance of the class.
- `coord(3)`—`coord()` with one argument, here 3—returns a  $1 \times 3$  vector, each element of which is a separate instance of the class.
- `coord(2,3)`—`coord()` with two arguments, here 2 and 3—returns a  $2 \times 3$  matrix, each element of which is a separate instance of the class.

Function `coord()` is useful when using Mata interactively. It is the only way to create instances of a class interactively.

In functions, you can create scalar instances by coding `class coord scalar name`, but in all other cases, you will also need to use function `coord()`. In the following example, the programmer wants a vector of coords:

```
void myfunction()
{
    real scalar          i
    class coord vector    v
    ...
    v = coord(3)
    for (i=1; i<=3; i++) v[i].x = v[i].y = 1
    ...
}
```

This program would have generated a compile-time error had the programmer omitted the line `class coord vector v`. Variable declarations are usually optional in Mata, but variable declarations of type class are not optional.

This program would have generated a run-time error had the programmer omitted the line `v = coord(3)`. Because `v` was declared to be `vector`, `v` began life as  $1 \times 0$ . The first thing the programmer needed to do was to expand `v` to be  $1 \times 3$ .

In practice, one seldom needs class vectors or matrices. A more typical program would read

```
void myfunction()
{
    real scalar          i
    class coord scalar    v
    ...
    v.x = v.y = 1
    ...
}
```

Note particularly the line `class coord scalar v`. The most common error programmers make is to omit the word `scalar` from the declaration. If you do that, then `matrix` is assumed, and then, rather than `v` being  $1 \times 1$ , it will be  $0 \times 0$ . If you did omit the word `scalar`, you have two alternatives. Either go back and put the word back in, or add the line `v = coord()` before initializing `v.x` and `v.y`. It does not matter which you do. In fact, when you specify `scalar`, the compiler merely inserts the line `v = coord()` for you.

## Constructors and destructors

You can specify how variables are initialized, and more, each time a new instance of a class is created, but before we get to that, let's understand the default initialization. In

```
class coord {
    real scalar    x, y
    real scalar    length(), angle()
}
```

there are two variables. When an instance of a `coord` is created, say, by coding `b = coord()`, the values are filled in the usual Mata way. Here, because `x` and `y` are scalars, `b.x` and `b.y` will be filled in with missing. Had they been vectors, row vectors, column vectors, or matrices, they would have been dimensioned  $1 \times 0$  (vectors and row vectors),  $0 \times 1$  (column vectors), and  $0 \times 0$  (matrices).

If you want to control initialization, you may declare a constructor function named `new()` in your class:

```
class coord {
    real scalar    x, y
    real scalar    length(), angle()
    void          new()
}
```

Function `new()` must be declared to be `void` and must take no arguments. You never bother to call `new()` yourself—in fact, you are not allowed to do that. Instead, function `new()` is called automatically each time a new instance of the class is created.

If you wanted every new instance of a `coord` to begin life with `x` and `y` equal to 0, you could code

```
void coord::new()
{
    x = y = 0
}
```

Let's assume we do that and we then inherit from `coord` when creating the new class `rotated_coord`, just as shown in the first example.

```
class rotated_coord extends coord {
    real scalar theta
    real scalar    angle()
    void           new()
}

void rotated_coord::new()
{
    theta = 0
}
```

When we create an instance of a `rotated_coord`, all the variables were initialized to 0. That is, function `rotated_coord()` will know to call both `coord::new()` and `rotated_coord::new()`, and it will call them in that order.

In your `new()` function, you are not limited to initializing values. `new()` is a function and you can code whatever you want, so if you want to set up a link to a radio telescope and check for any incoming messages, you can do that. Closer to home, if you were implementing a file system, you might use a `static` variable to count the number of open files. (We will explain static member variables later.)

On the other side of instance creation—instance destruction—you can declare and create `void destroy()`, which also takes no arguments, to be called each time an instance is destroyed. `destroy()` is known as a destructor. `destroy()` works like `new()` in the sense that you are not allowed to call the function directly. Mata calls `destroy()` for you whenever Mata is releasing the memory associated with an instance. `destroy()` does not serve much use in Mata because, in Mata, memory management is automatic and the freeing of memory is not your responsibility. In some other object-oriented programming languages, such as C++, you are responsible for memory management, and the destructor is where you free the memory associated with the instance.

Still, there is an occasional use for `destroy()` in Mata. Let's consider a system that needs to count the number of instances of itself that exists, perhaps so it can release a resource when the instance count goes to 0. Such a system might, in part, read

```
class bigsystem {
    static real scalar counter
    ...
    void           new(), destroy()
    ...
}

void bigsystem::new()
{
    counter = (counter==. ? 1 : counter+1)
}

void bigsystem::destroy()
{
    counter--
}
```

Note that `bigsystem::new()` must deal with two initializations, first and subsequent. The first time it is called, `counter` is `.` and `new()` must set `counter` to be `1`. After that, `new()` must increment `counter`.

If this system needed to obtain a resource, such as access to a radio telescope, on first initialization and release it on last destruction, and be ready to repeat the process, the code could read

```
void bigsystem::new()
{
    if (counter==.) {
        get_resource()
        counter = 1
    }
    else {
        ++counter
    }
}

void bigsystem::destroy()
{
    if (--counter == 0) {
        release_resource()
        counter = .
    }
}
```

Note that `destroy()`s are run whenever an instance is destroyed, even if that destruction is due to an abort-with-error in the user's program or even in the member functions. The radio telescope will be released when the last instance of `bigsystem` is destroyed, except in one case. The exception is when there is an error in `destroy()` or any of the subroutines `destroy()` calls.

For inheritance, child `destroy()`s are run before parent `destroy()`s.

## Setting member variable and member function exposure

Exposure specifies who may access the member variables and member functions of your class. There are three levels of exposure: from least restrictive to most restrictive, `public`, `protected`, and `private`.

- Public variables and functions may be accessed by anyone, including callers from outside a class.
- Protected variables and functions may be accessed only by [member functions](#) of a class and its [children](#).
- Private variables and functions may be accessed only by member functions of a class (excluding children).

When you do not specify otherwise, variables and functions are `public`. For code readability, you may declare member variables and functions to be `public` with `public`, but this is not necessary.

In programming large systems, it is usually considered good style to make member variables `private` and to provide public functions to set and to access any variables that users might need. This way, the internal design of the class can be subsequently modified and other classes that inherit from the class will not need to be modified, they will just need to be recompiled.

You make member variables and functions protected or private by preceding their declaration with `protected` or `private`:

```
class myclass {
    private real matrix      X
    private real vector      y
    void                    setup()
    real matrix              invert()
    protected real matrix    invert_subroutine()
}
```

Alternatively, you can create blocks with different defaults:

```
class myclass {
    public:
        void                    setup()
        real matrix              invert()
    protected:
        real matrix              invert_subroutine()
    private:
        real matrix              X
        real vector              y
}
```

You may combine `public`, `private`, and `protected`, with or without colons, freely.

## Making a member final

A member variable or function is said to be `final` if no [children](#) define a variable or function of the same name. Ensuring that a definition is the final definition can be enforced by including `final` in the declaration of the member, such as

```
class myclass {
    public:
        void                    setup()
        real matrix              invert()
    protected:
        final real matrix        invert_subroutine()
    private:
        real matrix              X
        real vector              y
}
```

In the above, no class that inherits from `myclass` can redefine `invert_subroutine()`.

## Making a member static

Being static is an optional property of a class member variable function. In what follows, let `ex1` and `ex2` both be instances of `S`, assume `c` is a static variable of the class, and assume `f()` is a static function of the class.

A static variable is a variable whose value is shared across all instances of the class. Thus, `ex1.c` and `ex2.c` will always be equal. If `ex1.c` is changed, say, by `ex1.c=5`, then `ex2.c` will also be 5.

A static function is a function that does not use the values of nonstatic member variables of the class. Thus `ex1.f(3)` and `ex2.f(3)` will be equal regardless of how `ex1` and `ex2` differ.

Outside of member functions, static functions may also be invoked by coding the class name, two colons, and the function name, and thus used even if you do not have a class instance. For instance, equivalent to coding `ex1.f(3)` or `ex2.f(3)` is `S::f(3)`.

## Virtual functions

When a function is declared to be virtual, the rules of [inheritance](#) act as though they were reversed. Usually, when one class inherits from another, if the child class does not define a function, then it inherits the parent's function. For virtual functions, however, the parent has access to the child's definition of the function, which is why we say that it is, in some sense, reverse inheritance.

The canonical motivational example of this deals with animals. Without loss of generality, we will use barnyard animals to illustrate. A class, `animal`, is defined. Classes `cow` and `sheep` are defined that extend (inherit from) `animal`. One of the functions defined in `animal` needs to make the sound of the specific animal, and it calls virtual function `sound()` to do that. At the `animal` level, function `animal::sound()` is defined to display "Squeak!". Because function `sound()` is virtual, however, each of the specific-animal classes is supposed to define their own sound. Class `cow` defines `cow::sound()` that displays "Moo!" and class `sheep` defines `sheep::sound()` that displays "Baa!". The result is that when a routine in `animal` calls `sound()`, it behaves as if the inheritance is reversed, the appropriate `sound()` routine is called, and the user sees "Moo!" or "Baa!". If a new specific animal is added, and if the programmer forgets to define `sound()`, then `animal::sound()` will run, and the user sees "Squeak!". In this case, that is supposed to be the sound of the system in trouble, but it is really just the default action.

Let's code this example. First, we will code the usual case:

```
class animal {
    ...
    void      sound()
    void      poke()
    ...
}
void animal::sound() { "Squeak!" }
void animal::poke()
{
    ...
    sound()
    ...
}
class cow extends animal {
    ...
}
class sheep extends animal {
    ...
}
```

In the above example, when an animal, cow, or sheep is poked, among other things, it emits a sound. Poking is defined at the animal level because poking is mostly generic across animals, except for

the sound they emit. If `c` is an instance of `cow`, then one can poke that particular cow by coding `c.poke()`. `poke()`, however, is inherited from `animal`, and the generic action is taken, along with the cow emitting a generic squeak.

Now we make `sound()` virtual:

```
class animal {
    ...
    virtual void  sound()
    void          poke()
    ...
}
void animal::sound() { "Squeak!" }
void animal::poke()
{
    ...
    sound()
    ...
}
class cow extends animal {
    ...
    virtual void  sound()
}
void cow::sound() { "Moo!" }
class sheep extends animal {
    ...
    virtual void  sound()
}
void sheep::sound() { "Baa!" }
```

Now let's trace through what happens when we poke a particular cow by coding `c.poke()`. `c`, to remind you, is a `cow`. There is no `cow::poke()` function; however, `c.poke()` executes `animal::poke()`. `animal::poke()` calls `sound()`, which, were `sound()` not a virtual function, would be `animal::sound()`, which would emit a squeak. Poke a cow, get a "Squeak!" Because `sound()` is virtual, `animal::poke()` called with a cow calls `cow::sound()`, and we hear a "Moo!"

Focusing on syntax, it is important that both `cow` and `sheep` repeated `virtual void sound()` in their declarations. If you define function  $S:f()$ , then  $f()$  must be declared in  $S$ . Consider a case, however, where we have two breeds of cows, `Angus` and `Holstein`, and they emit slightly different sounds. The `Holstein`, being of Dutch origin, gets a bit of a U sound into the moo in a way no native English speaker can duplicate. Thus we would declare two new classes, `angus extends cow` and `holstein extends cow`, and we would define `angus::sound()` and `holstein::sound()`. Perhaps we would not bother to define `angus::sound()`; then an `Angus` would get the generic cow sound.

But let's pretend that, instead, we defined `angus::sound()` and removed the function for `cow::sound()`. Then it does not matter whether we include the line `virtual void sound()` in `cow`'s declaration. Formally, it should be included, because the line of reverse declaration should not be broken, but Mata does not care one way or the other.

A common use of virtual functions is to allow you to process a list of objects without any knowledge of the specific type of object, as long as all the objects are subclasses of the same base class:

```

class animal      animals
animals = animal(3,1)
animals[1] = cow()
animals[2] = sheep()
animals[3] = holstein()
for(i=1; i<=length(animals); i++) {
    animals[i].sound()
}

```

Note the use of `animals = animal(3,1)` to initialize the vector of animals. This is an example of how to create a nonscalar class instance, as discussed in [Obtaining instances of a class](#).

When the code above is executed, the appropriate sound for each animal will be displayed even though `animals` is a vector declared to be of type `class animal`. Because `sound()` is virtual, the appropriate sound from each specific animal child class is called.

## Referring to the current class using this

`this` is used within member functions to refer to the class as a whole. For instance, consider the following:

```

class S {
    real scalar      n
    real matrix      M
    void             make_M()
}
void S::make_M(real scalar n)
{
    real scalar      i, j
    this.n = n
    M = J(n, n, 0)
    for (i=1; i<=n; i++) {
        for (j=1; j<=i; j++) M[i,j] = 1
    }
}

```

In the above program, references to `M` are understood to be the class instance definition of `M`. References to `n`, however, refer to the function's definition of `n` because the program's `n` has precedence over the class's definition of it. `this.n` is how one refers to the class's variable in such cases. `M` also could have been referred to as `this.M`, but that was not necessary.

If, in function `S::f()`, it was necessary to call a function outside of the class, and if that function required that we pass the class instance as a whole as an argument, we could code `this` for the class instance. The line might read `outside_function(this)`.



## Using super to access the parent’s concept

The `super` modifier is a way of dealing with variables and functions that have been [overridden](#) in subclasses or, said differently, is a way of accessing the parent’s concept of a variable or function.

Let `T` extend `S` and let `t` be an instance of `T`. Then `t.f()` refers to `T::f()` if the function exists, and otherwise to `S::f()`. `t.super.f()` always refers to `S::f()`.

More generally, in a series of inheritances, `z.super.f()` refers to the parent’s concept of `f()`—the `f()` the parent would call if no `super` were specified—`z.super.super.f()` refers to the grandparent’s concept of `f()`, and so on. For example, let `W` extend `V` extend `U` extend `T` extend `S`. Furthermore, assume

<code>S::f()</code>	exists	
<code>T::f()</code>		does not exist
<code>U::f()</code>	exists	
<code>V::f()</code>		does not exist
<code>W::f()</code>		does not exist

Finally, let `s` be an instance of `S`, `t` be an instance of `T`, and so on. Then calls of the form `w.f()`, `w.super.f()`, ..., `w.super.super.super.super.f()`, `v.f()`, `v.super.f()`, and so on, result in execution of

		Number of supers specified				
		0	1	2	3	4
<code>s.</code>	<code>S::f()</code>					
<code>t.</code>	<code>S::f()</code>		<code>S::f()</code>			
<code>u.</code>	<code>U::f()</code>		<code>S::f()</code>	<code>S::f()</code>		
<code>v.</code>	<code>U::f()</code>		<code>U::f()</code>	<code>S::f()</code>	<code>S::f()</code>	
<code>w.</code>	<code>U::f()</code>		<code>U::f()</code>	<code>U::f()</code>	<code>S::f()</code>	<code>S::f()</code>

## Casting back to a parent

A [class instance](#) may be treated as a class instance of a parent by casting. Assume `U` extends `T` extends `S`, and let `u` be an instance of `U`. Then

```
((class T) u
```

is treated as an instance of `T`, and

```
((class S) u
```

is treated as an instance of `S`. `((class T) u)` could be used anywhere an instance of `T` is required and `((class S) u)` could be used anywhere an instance of `S` is required.

For instance, assume `S::f()` is [overridden](#) by `T::f()`. If an instance of `U` found it necessary to call `S::f()`, one way it could do that would be `u.super.super.f()`. Another would be

```
((class S) u).f()
```

## Accessing external functions from member functions

In the opening example, class `coord` contained a member function named `length()`. Had we been coding a regular (nonclass) function, we could not have created that function with the name `length()`. The name `length()` is reserved by Mata for its own function that returns the length (number of elements) of a vector; see [M-5] [rows\(\)](#). We are allowed to create a function named `length()` inside classes. Outside of member functions, `c.length()` is an unambiguous reference to `c`'s definition of `length()`, and `length()` by itself is an unambiguous reference to `length()`'s usual definition.

There is, however, possible confusion when coding member functions. Let's add another member function to `coord`: `ortho()`, the code for which reads

```
class coord scalar coord::ortho()
{
    class coord scalar      newcoord
    real scalar             r, t
    r = length()
    t = angle()
    newcoord.x = r*cos(t)
    newcoord.y = r*sin(t)
    return(newcoord)
}
```

Note that in the above code, we call `length()`. Because we are writing a member function, and because the class defines a member function of that name, `length()` is interpreted as being a call to `coord`'s definition of `length()`. If `coord` did not define such a member function, then `length()` would have been interpreted as a call to Mata's `length()`.

So what do we do if the class provides a function, externally there is a function of the same name, and we want the external function? We prefix the function's name with double colons. If we wanted `length()` to be interpreted as Mata's function and not the class's, we would have coded

```
r = ::length()
```

There is nothing special about the name `length()` here. If the class provided member function `myfcn()`, and there was an external definition as well, and we wanted the externally defined function, we would code `::myfcn()`. In fact, it is not even necessary that the class provide a definition. If we cannot remember whether class `coord` includes `myfcn()`, but we know we want the external `myfcn()` no matter what, we can code `::myfcn()`.

Placing double-colons in front of external function names used inside the definitions of member functions is considered good style. This way, if you ever go back and add another member function to the class, you do not have to worry that some already written member function is assuming that the name goes straight through to an externally defined function.

## Pointers to classes

Just as you can obtain a pointer to any other Mata variable, you can obtain a pointer to an instance of a class. Recall our [example](#) in [Virtual functions](#) of animals that make various sounds when poked.

For the sake of example, we will obtain a pointer to an instance of a `cow()` and access the `poke()` function through the pointer:

```
void pokeacow() {  
    class cow scalar          c  
    pointer(class cow scalar) scalar  p  
    p = &c  
    p->poke()  
}
```

You access the member variables and functions through a pointer to a class with the operator `->` just like you would for structures. See [Pointers to structures](#) in [M-2] **struct** for more information.

## Also see

[M-2] **declarations** — Declarations and types

[M-2] **struct** — Structures

[P] **class** — Class programming

[M-2] **intro** — Language definition

## Syntax

```
/* enclosed comment */
```

```
// rest-of-line comment
```

Notes:

1. Comments may appear in do-files and ado-files; they are not allowed interactively.
2. Stata's beginning-of-the-line asterisk comment is not allowed in Mata:

```
. * valid in Stata but not in Mata
```

## Description

*/\** and *\*/* and *//* are how you place comments in Mata programs.

## Remarks

There are two comment styles: */\** and *\*/* and *//*. You may use one, the other, or both.

Remarks are presented under the following headings:

*The */\* \*/* enclosed comment*

*The *//* rest-of-line comment*

### The */\* \*/* enclosed comment

Enclosed comments may appear on a line:

```
/* What follows uses an approximation formula: */
```

Enclosed comments may appear within a line and even in the middle of a Mata expression:

```
x = x + /*left-single quote*/ char(96)
```

Enclosed comments may themselves contain multiple lines:

```
/*  
    We use the approximation based on sin(x) approximately  
    equaling x for small x; x measure in radians  
*/
```

Enclosed comments may be nested, which is useful for commenting out code that itself contains comments:

```
/*  
for (i=1; i<=rows(x); i++) {           /* normalization */  
    x[i] = x[i] :/ value[i]  
}  
*/
```

## The `//` rest-of-line comment

The rest-of-line comment may appear by itself on a line

```
// What follows uses an approximation formula:
```

or it may appear at the end of a line:

```
x = x + char(96)      // append single quote
```

In either case, the comment concludes when the line ends.

## Also see

[\[M-2\] intro](#) — Language definition

# Title

[M-2] **continue** — Continue with next iteration of for, while, or do loop

## Syntax

```
for, while, or do {  
    ...  
    if (...) {  
        ...  
        continue  
    }  
    ...  
}  
...
```

## Description

`continue` restarts the innermost `for`, `while`, or `do` loop. Execution continues just as if the loop had reached its logical end.

`continue` nearly always occurs following an `if`.

## Remarks

The following two code fragments are equivalent:

```
for (i=1; i<=rows(A); i++) {  
    for (j=1; j<=cols(A); j++) {  
        if (i==j) continue  
        ... action to be performed on A[i,j] ...  
    }  
}
```

and

```
for (i=1; i<=rows(A); i++) {  
    for (j=1; j<=cols(A); j++) {  
        if (i!=j) {  
            ... action to be performed on A[i,j] ...  
        }  
    }  
}
```

`continue` operates on the innermost `for` or `while` loop, and even when the `continue` action is taken, standard end-of-loop processing takes place (which is `j++` here).

## Also see

[M-2] **do** — do ... while (exp)

[M-2] **for** — for (exp1; exp2; exp3) stmt

[M-2] **while** — while (exp) stmt

[M-2] **break** — Break out of for, while, or do loop

[M-2] **intro** — Language definition

Syntax

```
declaration1 fcname(declaration2)
{
    declaration3
    ...
}
```

such as

```
real matrix myfunction(real matrix I, real scalar i)
{
    real scalar      j, k
    real vector      v
    ...
}
```

declaration<sub>1</sub> is one of

```
function
type [function]
void [function]
```

declaration<sub>2</sub> is

```
[type] argname [, [type] argname [, ... ]]
```

where *argname* is the name you wish to assign to the argument.

declaration<sub>3</sub> are lines of the form of either of

```
type          varname [, varname [, ... ]]
```

```
external [type] varname [, varname [, ... ]]
```

*type* is defined as one of

<i>eltype</i>	<i>orgtype</i>	such as real vector
<i>eltype</i>		such as real
<i>orgtype</i>		such as vector

*eltype* and *orgtype* are each one of

<i>eltype</i>	<i>orgtype</i>
transmorphic	matrix
numeric	vector
real	rowvector
complex	colvector
string	scalar
pointer	



If *eltype* is not specified, *transmorphic* is assumed. If *orgtype* is not specified, *matrix* is assumed.

## Description

Types and the use of declarations is explained. Also discussed is the calling convention (functions are called by address, not by value, and so may change the caller's arguments), and the use of external globals.

Mata also has structures—the *eltype* is `struct name`—but these are not discussed here. For a discussion of structures, see [M-2] [struct](#).

Mata also has classes—the *eltype* is `class name`—but these are not discussed here. For a discussion of classes, see [M-2] [class](#).

Declarations are optional but, for careful work, their use is recommended.

## Remarks

Remarks are presented under the following headings:

- The purpose of declarations*
- Types, element types, and organizational types*
- Implicit declarations*
- Element types*
- Organizational types*
- Function declarations*
- Argument declarations*
- The by-address calling convention*
- Variable declarations*
- Linking to external globals*

## The purpose of declarations

Declarations occur in three places: in front of function definitions, inside the parentheses defining the function's arguments, and at the top of the body of the function, defining private variables the function will use. For instance, consider the function

```
real matrix swaprows(real matrix A, real scalar i1, real scalar i2)
{
    real matrix      B
    real rowvector   v
    B = A
    v = B[i1, .]
    B[i1, .] = B[i2, .]
    B[i2, .] = v
    return(B)
}
```

This function returns a copy of matrix A with rows i1 and i2 swapped.

There are three sets of declarations in the above function. First, there is a declaration in front of the function name:

```
real matrix swaprows(...)  
{  
    ...  
}
```

That declaration states that this function will return a real matrix.

The second set of declarations occur inside the parentheses:

```
... swaprows(real matrix A, real scalar i1, real scalar i2)  
{  
    ...  
}
```

Those declarations state that this function expects to receive three arguments, which we chose to call A, i1, and i2, and which we expect to be a real matrix, a real scalar, and a real scalar, respectively.

The third set of declarations occur at the top of the body of the function:

```
... swaprows(...)  
{  
    real matrix      B  
    real rowvector v  
    ...  
}
```

Those declarations state that we will use variables B and v inside our function and that, as a matter of fact, B will be a real matrix and v a real row vector.

We could have omitted all those declarations. Our function could have read

```
function swaprows(A, i1, i2)  
{  
    B = A  
    v = B[i1, .]  
    B[i1, .] = B[i2, .]  
    B[i2, .] = v  
    return(B)  
}
```

and it would have worked just fine. So why include the declarations?

1. By including the outside declaration, we announced to other programs what to expect. They can depend on `swaprows()` returning a real matrix because, when `swaprows()` is done, Mata will verify that the function really is returning a real matrix and, if it is not, abort execution.

Without the outside declaration, anything goes. Our function could return a real scalar in one case, a complex row vector in another, and nothing at all in yet another case.

Including the outside declaration makes debugging easier.

2. By including the argument declaration, we announced to other programmers what they are expected to pass to our function. We have made it easier to understand our function.

We have also told Mata what to expect and, if some other program attempts to use our function incorrectly, Mata will stop execution.

Just as in (1), we have made debugging easier.

3. By including the inside declaration, we have told Mata what variables we will need and how we will be using them. Mata can do two things with that information: first, it can make sure that we are using the variables correctly (making debugging easier again), and second, Mata can produce more efficient code (making our function run faster).

Interactively, we admit that we sometimes define functions without declarations. For more careful work, however, we include them.

## Types, element types, and organizational types

When you use Mata interactively, you just willy-nilly create new variables:

```
: n = 2
: A = (1,2 \ 3,4)
: z = (sqrt(-4+0i), sqrt(4))
```

When you create a variable, you may not think about the type, but Mata does. `n` above is, to Mata, a real scalar. `A` is a real matrix. `z` is a complex row vector.

Mata thinks of the type of a variable as having two parts:

1. the type of the elements the variable contains (such as real or complex) and
2. how those elements are organized (such as a row vector or a matrix).

We call those two types the *eltype*—element type—and *orgtype*—organizational type. The *eltypes* and *orgtypes* are

<i>eltype</i>	<i>orgtype</i>
transmorphic	matrix
numeric	vector
real	rowvector
complex	colvector
string	scalar
pointer	

You may choose one of each and so describe all the types Mata understands.

## Implicit declarations

When you do not declare an object, Mata behaves as if you declared it to be `transmorphic matrix`:

1. `transmorphic` means that the matrix can be `real`, `complex`, `string`, or `pointer`.
2. `matrix` means that the organization is to be  $r \times c$ ,  $r \geq 0$  and  $c \geq 0$ .

At one point in your function, a transmorphic matrix might be a real scalar (real being a special case of transmorphic and scalar being a special case of a matrix when  $r = c = 1$ ), and at another point, it might be a string colvector (string being a special case of transmorphic, and colvector being a special case of a matrix when  $c = 1$ ).

Consider our `swaprows()` function without declarations,

```
function swaprows(A, i1, i2)
{
    B = A
    v = B[i1, .]
    B[i1, .] = B[i2, .]
    B[i2, .] = v
    return(B)
}
```

The result of compiling this function is just as if the function read

```
transmorphic matrix swaprows(transmorphic matrix A,
                              transmorphic matrix i1,
                              transmorphic matrix i2)
{
    transmorphic matrix    B
    transmorphic matrix    v
    B = A
    v = B[i1, .]
    B[i1, .] = B[i2, .]
    B[i2, .] = v
    return(B)
}
```

When we declare a variable, we put restrictions on it.

## Element types

There are six *eltypes*, or element types:

1. **transmorphic**, which means real, complex, string, or pointer.
2. **numeric**, which means real or complex.
3. **real**, which means that the elements are real numbers, such as 1, 3,  $-50$ , and 3.14159.
4. **complex**, which means that each element is a pair of numbers, which are given the interpretation  $a + bi$ . **complex** is a storage type; the number stored in a complex might be real, such as  $2 + 0i$ .
5. **string**, which means the elements are strings of text. Each element may contain up to 2,147,483,647 characters and strings may (need not) contain binary 0; that is, strings may be binary strings or text strings.
6. **pointer** means the elements are pointers to (addresses of) other Mata matrices, vectors, scalars, or even functions; see [M-2] **pointers**.

## Organizational types

There are five *orgtypes*, or organizational types:

1. `matrix`, which means  $r \times c$ ,  $r \geq 0$  and  $c \geq 0$ .
2. `vector`, which means  $1 \times n$  or  $n \times 1$ ,  $n \geq 0$ .
3. `rowvector`, which means  $1 \times n$ ,  $n \geq 0$ .
4. `colvector`, which means  $n \times 1$ ,  $n \geq 0$ .
5. `scalar`, which means  $1 \times 1$ .

Sharp-eyed readers will note that vectors and matrices can have zero rows or columns! See [\[M-2\] void](#) for more information.

## Function declarations

Function declarations are the declarations that appear in front of the function name, such as

```
real matrix swaprows(...)
{
    ...
}
```

The syntax for what may appear there is

```
function
type [function]
void [function]
```

Something must appear in front of the name, and if you do not want to declare the type (which makes the type `transmorphic matrix`), you just put the word `function`:

```
function swaprows(...)
{
    ...
}
```

You may also declare the type and include the word `function` if you wish,

```
real matrix function swaprows(...)
{
    ...
}
```

but most programmers omit the word `function`; it makes no difference.

In addition to all the usual types, `void` is a type allowed only with functions—it states that the function returns nothing:

```

void _swaprows(real matrix A, real scalar i1, real scalar i2)
{
    real rowvector v
    v = A[i1, .]
    A[i1, .] = A[i2, .]
    A[i2, .] = v
}

```

The function above returns nothing; it instead modifies the matrix it is passed. That might be useful to save memory, especially if every use of the original `swaprows()` was going to be

```
A = swaprows(A, i1, i2)
```

In any case, we named this new function `_swaprows()` (note the underscore), to flag the user that there is something odd and deserving caution concerning the use of this function.

`void`, that is to say, returning nothing, is also considered a special case of a *transmorphic matrix* because Mata secretly returns a  $0 \times 0$  real matrix, which the caller just discards.

## Argument declarations

Argument declarations are the declarations that appear inside the parentheses, such as

```

... swaprows(real matrix A, real scalar i1, real scalar i2)
{
    ...
}

```

The syntax for what may appear there is

```
[type] argname [, [type] argname [, ...]]
```

The names are required—they specify how we will refer to the argument—and the types are optional. Omit the type and *transmorphic matrix* is assumed. Specify the type, and it will be checked when your function is called. If the caller attempts to use your function incorrectly, Mata will stop the execution and complain.

## The by-address calling convention

Arguments are passed to functions by address, not by value. If you change the value of an argument, you will change the caller's argument. That is what made `_swaprows()` (above) work. The caller passed us `A` and we changed it. And that is why in the original version of `swaprows()`, the first line read

```
B = A
```

we did our work on `B`, and returned `B`. We did not want to modify the caller's original matrix.

You do not ordinarily have to make copies of the caller's arguments, but you do have to be careful if you do not want to change the argument. That is why in all the official functions (with the single exception of `st_view()`—see [M-5] `st_view()`), if a function changes the caller's argument, the function's name starts with an underscore. The reverse logic does not hold: some functions start with an underscore and do not change the caller's argument. The underscore signifies caution, and you need to read the function's documentation to find out what it is you need to be cautious about.

## Variable declarations

The variable declarations are the declarations that appear at the top of the body of a function:

```
... swaprows(...)
{
    real matrix      B
    real rowvector  v
    ...
}
```

These declarations are optional. If you omit them, Mata will observe that you are using `B` and `v` in your code, and then Mata will compile your code just as if you had declared the variables to be **transmorphic matrix**, meaning that the resulting compiled code might be a little more inefficient than it could be, but that is all.

The variable declarations are optional as long as you have not mata set `matastrict` on; see [M-3] **mata set**. Some programmers believe so strongly that variables really ought to be declared that Mata provides a provision to issue an error when they forget.

In any case, these declarations—explicit or implicit—define the variables we will use. The variables we use in our function are private—it does not matter if there are other variables named `B` and `v` floating around somewhere. Private variables are created when a function is invoked and destroyed when the function ends. The variables are private but, as explained above, if we pass our variables to another function, that function may change their values. Most functions do not.

The syntax for declaring variables is

```
type           varname [ , varname [ , ... ] ]
external [ type ] varname [ , varname [ , ... ] ]
```

`real matrix B` and `real rowvector v` match the first syntax.

## Linking to external globals

The second syntax has to do with linking to global variables. When you use Mata interactively and type

```
: n = 2
```

you create a variable named `n`. That variable is global. When you code inside a function

```
... myfunction(...)
{
    external n
    ...
}
```

The `n` variable your function will use is the global variable named `n`. If your function were to examine the value of `n` right now, it would discover that it contained 2.

If the variable does not already exist, the statement `external n` will create it. Pretend that we had not previously defined `n`. If `myfunction()` were to examine the contents of `n`, it would discover that `n` is a  $0 \times 0$  matrix. That is because we coded

```
external n
```

and Mata behaved as if we had coded

```
external transmorphic matrix n
```

Let's modify `myfunction()` to read:

```
... myfunction(...)
{
    external real scalar n
    ...
}
```

Let's consider the possibilities:

1. `n` does not exist. Here `external real scalar n` will create `n`—as a real scalar, of course—and set its value to `missing`.  
If `n` had been declared a `rowvector`, a  $1 \times 0$  vector would have been created.  
If `n` had been declared a `colvector`, a  $0 \times 1$  vector would have been created.  
If `n` had been declared a `vector`, a  $0 \times 1$  vector would have been created. Mata could just as well have created a  $1 \times 0$  vector, but it creates a  $0 \times 1$ .  
If `n` had been declared a `matrix`, a  $0 \times 0$  matrix would have been created.
2. `n` exists, and it is a `real scalar`. Our function executes, using the global `n`.
3. `n` exists, and it is a `real  $1 \times 1$  rowvector, colvector, or matrix`. The important thing is that it is  $1 \times 1$ ; our function executes, using the global `n`.
4. `n` exists, but it is `complex` or `string` or `pointer`, or it is `real` but not  $1 \times 1$ . Mata issues an error message and aborts execution of our function.

Complicated systems of programs sometimes find it convenient to communicate via globals. Because globals are globals, we recommend that you give your globals long names. A good approach is to put the name of your system as a prefix:

```
... myfunction(...)
{
    external real scalar mysystem_n
    ...
}
```

For another approach to globals, see [\[M-5\] `findexternal\(\)`](#) and [\[M-5\] `valofexternal\(\)`](#).

## Also see

[\[M-2\] `intro`](#) — Language definition



# Title

[M-2] **do** — do ... while (exp)

## Syntax

```
do stmt while (exp)
```

```
do {  
    stmts  
} while (exp)
```

where *exp* must evaluate to a real scalar.

## Description

do executes *stmt* or *stmts* one or more times, until *exp* is zero.

## Remarks

One common use of do is to loop until convergence:

```
do {  
    lasta = a  
    a = get_new_a(lasta)  
} while (mreldif(a, lasta)>1e-10)
```

The loop is executed at least once, and the conditioning expression is not executed until the loop's body has been executed.

## Also see

[M-2] **for** — for (exp1; exp2; exp3) stmt

[M-2] **while** — while (exp) stmt

[M-2] **break** — Break out of for, while, or do loop

[M-2] **continue** — Continue with next iteration of for, while, or do loop

[M-2] **intro** — Language definition

## Description

When an error occurs, Mata presents a number as well as text describing the problem. The codes are presented below.

Also the error codes can be used as an argument with `_error()`, see [\[M-5\] `error\(\)`](#).

Mata's error codes are a special case of Stata's return codes. In particular, they are the return codes in the range 3000–3999. In addition to the 3000-level codes, it is possible for Mata functions to generate any Stata error message and return code.

## Remarks

Error messages in Mata break into two classes: errors that occur when the function is compiled (code 3000) and errors that occur when the function is executed (codes 3001–3999).

Compile-time error messages look like this:

```
: 2,,3
invalid expression
r(3000);

: "this" + "that
mismatched quotes
r(3000);
```

The text of the message varies according to the error made, but the error code is always 3000.

Run-time errors look like this:

```
: myfunction(2,3)
      solve(): 3200 conformability error
      mysub(): - function returned error
myfunction(): - function returned error
      <istmt>: - function returned error
r(3200);
```

The output is called a traceback log. Read from bottom to top, it says that what we typed (the `<istmt>`) called `myfunction()`, which called `mysub()`, which called `solve()` and, at that point, things went wrong. The error is possibly in `solve()`, but because `solve()` is a library function, it is more likely that the error is in how `mysub()` called `solve()`. Moreover, the error is seldom in the program listed at the top of the traceback log because the log lists the identity of the program that detected the error. Say `solve()` did have an error. Then the traceback log would probably have read something like

```
      *: 3200 conformability error
      solve(): - function returned error
      mysub(): - function returned error
myfunction(): - function returned error
      <istmt>: - function returned error
```

The above log says the problem was detected by `*` (the multiplication operator), and at that point, `solve()` would be suspect, because one must ask, why did `solve()` use the multiplication operator incorrectly?

In any case, let's assume that the problem is not with `solve()`. Then you would guess the problem lies with `mysub()`. If you have used `mysub()` in many previous programs without problems, however, you might now shift your suspicion to `myfunction()`. If `myfunction()` is always trustworthy, perhaps you should not have typed `myfunction(2,3)`. That is, perhaps you are using `myfunction()` incorrectly.

## The error codes

3000. (message varies)

There is an error in what you have typed. Mata cannot interpret what you mean.

3001. incorrect number of arguments

The function expected, say, three arguments and received two, or five. Or the function allows between three and five arguments, but you supplied too many or too few. Fix the calling program.

3002. identical arguments not allowed

You have called a function specifying the same variable more than once. Usually this would not be a problem, but here, it is, usually because the supplied arguments are matrices that the function wishes to overwrite with different results. For instance, say function  $f(A, B, C)$  examines matrix  $A$  and returns a calculation based on  $A$  in  $B$  and  $C$ . The function might well complain that you specified the same matrix for  $B$  and  $C$ .

3010. attempt to dereference NULL pointer

The program made reference to `*s`, and `s` contains NULL; see [\[M-2\] pointers](#).

3011. invalid lval

In an assignment, what appears on the left-hand side of the equals is not something to which a value can be assigned; see [\[M-2\] op\\_assignment](#).

3012. undefined operation on a pointer

You have, for instance, attempted to add two pointers; see [\[M-2\] pointers](#).

3020. class child/parent compiled at different times

One class is being used to extend another class, and the parent has been updated since the class was compiled.

3021. class compiled at different times

A function using a predefined class has not been recompiled since the class has last been changed.

3101. matrix found where function required

A particular argument to a function is required to be a function, and a matrix was found instead.

3102. function found where matrix required

A particular argument to a function is required to be a matrix, vector, or scalar, and a function was found instead.

3103. `view` found where `array` required

In general, view matrices can be used wherever a matrix is required, but there are a few exceptions, both in low-level routines and in routines that wish to write results back to the argument. Here a view is not acceptable. If  $V$  is the view variable, simply code  $X = V$  and then pass  $X$  in its stead. See [M-5] [st\\_view\(\)](#).

3104. `array` found where `view` required

A function argument was specified with a matrix that was not a view, and a view was required. See [M-5] [st\\_view\(\)](#).

## 3200. conformability error

A matrix, vector, or scalar has the wrong number of rows and/or columns for what is required. Adding a  $2 \times 3$  matrix to a  $1 \times 4$  would result in this error.

3201. `vector` required

An argument is required to be  $r \times 1$  or  $1 \times c$ , and a matrix was found instead.

3202. `rowvector` required

An argument is required to be  $1 \times c$  and it is not.

3203. `colvector` required

An argument is required to be  $r \times 1$  and it is not.

3204. `matrix` found where `scalar` required

An argument is required to be  $1 \times 1$  and it is not.

3205. `square matrix` required

An argument is required to be  $n \times n$  and it is not.

3206. invalid use of `view` containing `op.vars`

Factor variables have been used in a view, and the view is now being used in a context that does not support the use of factor variables.

3250. `type` mismatch

The *eltype* of an argument does not match what is required. For instance, perhaps a real was expected and a string was received. See *eltype* in [M-6] [Glossary](#).

3251. `nonnumeric` found where `numeric` required

An argument was expected to be real or complex and it is not.

3252. `noncomplex` found where `complex` required

An argument was expected to be complex and it is not.

3253. `nonreal` found where `real` required

An argument was expected to be real and it is not.

3254. `nonstring` found where `string` required

An argument was expected to be string and it is not.

3255. `real` or `string` required

An argument was expected to be real or string and it is not.

3256. `numeric` or `string` required

An argument was expected to be real, complex, or string and it is not.

3257. `nonpointer` found where `pointer` required

An argument was expected to be a pointer and it is not.

3258. **nonvoid found where void required**  
An argument was expected to be void and it is not.
3259. **nonstruct found where struct required**  
A variable that is not a structure was passed to a function that expected the variable to be a structure.
3260. **nonclass found where class required**  
A variable that is not a class was passed to a function that expected the variable to be a class.
3261. **non class/struct found where class/struct required**  
A variable that is not a class or a structure was passed to a function that expected the variable to be a class or a structure.
3300. **argument out of range**  
The *eltype* and *orgtype* of the argument are correct, but the argument contains an invalid value, such as if you had asked for the 20th row of a  $4 \times 5$  matrix. See *eltype* and *orgtype* in [M-6] **Glossary**.
3301. **subscript invalid**  
The subscript is out of range (refers to a row or column that does not exist) or contains the wrong number of elements. See [M-2] **subscripts**.
3302. **invalid %fmt**  
The %fmt for formatting data is invalid. See [M-5] **printf()** and see [U] **12.5 Formats: Controlling how data are displayed**.
3303. **invalid permutation vector**  
The vector specified does not meet the requirements of a permutation vector, namely, that an  $n$ -element vector contain a permutation of the integers 1 through  $n$ . See [M-1] **permutation**.
3304. **struct nested too deeply**  
Structures may contain structures that contain structures, and so on, but only to a depth of 500.
3305. **class nested too deeply**  
Classes may contain classes that contain classes, and so on, but only to a depth of 500.
3351. **argument has missing values**  
In general, Mata is tolerant of missing values, but there are exceptions. This function does not allow the matrix, vector, or scalar to have missing values.
3352. **singular matrix**  
The matrix is singular and the requested result cannot be carried out. If singular matrices are a possibility, then you are probably using the wrong function.
3353. **matrix not positive definite**  
The matrix is non-positive definite and the requested results cannot be computed. If non-positive-definite matrices are a possibility, then you are probably using the wrong function.
3360. **failure to converge**  
The function that issued this message used an algorithm that the function expected would converge but did not, probably because the input matrix was extreme in some way.

3492. `resulting string too long`

A string that the function was attempting to produce became too long. Because the maximum length of strings in Mata is 2,147,483,647 characters, it is unlikely that Mata imposed the limit. Review the documentation on the function for the source of the limit that was imposed (for example, perhaps a string was being produced for use by Stata). In any case, this error does not arise because of an out-of-memory situation. It arises because some limit was imposed.

3498. `(message varies)`

An error specific to this function arose. The text of the message should describe the problem.

3499. `_____ not found`

The specified variable or function could not be found. For a function, it was not already loaded, it is not in the libraries, and there is no `.mo` file with its name.

3500. `invalid Stata variable name`

A variable name—which name is contained in a Mata string variable—is not appropriate for use with Stata.

3598. `Stata returned error`

You are using a Stata interface function and have asked Stata to perform a task. Stata could not or refused.

3601. `invalid file handle`

The number specified does not correspond to an open file handle; see [M-5] [fopen\(\)](#).

3602. `invalid filename`

The filename specified is invalid.

3603. `invalid file mode`

The file mode (whether read, write, read-write, etc.) specified is invalid; see [M-5] [fopen\(\)](#).

3611. `too many open files`

The maximum number of files that may be open simultaneously is 50, although your operating system may not allow that many.

3612. `file too large for 32-bit Stata`

You are running 32-bit Stata on a 64-bit computer, and the file you wish to process is larger than 4 gigabytes.

3621. `attempt to write read-only file`

The file was opened read-only and an attempt was made to write into it.

3622. `attempt to read write-only file`

The file was opened write-only and an attempt was made to read it.

3623. `attempt to seek append-only file`

The file was opened append-only and then an attempt was made to seek into the file; see [M-5] [fopen\(\)](#).

3698. `file seek error`

An attempt was made to seek to an invalid part of the file, or the seek failed for other reasons; see [M-5] [fopen\(\)](#).

## 3900. out of memory

Mata is out of memory; the operating system refused to supply what Mata requested. There is no Mata or Stata setting that affects this, and so nothing in Mata or Stata to reset in order to get more memory. You must take up the problem with your operating system.

## 3901. macro memory in use

This error message should not occur; please notify StataCorp if it does.

## 3930. error in LAPACK routine

The linear-algebra LAPACK routines—see [M-1] **LAPACK**—generated an error that Mata did not expect. Please notify StataCorp if you should receive this error.

## 3995. unallocated function

This error message should not occur; please notify StataCorp if it does.

## 3996. built-in unallocated

This error message should not occur; please notify StataCorp if it does.

## 3997. unimplemented opcode

This error message should not occur; please notify StataCorp if it does.

## 3998. stack overflow

Your program nested too deeply. For instance, imagine calculating the factorial of  $n$  by recursively calling yourself and then requesting the factorial of  $1e+100$ . Functions that call themselves in an infinite loop inevitably cause this error.

## 3999. system assertion false

Something unexpected happened in the internal Mata code. Please contact Technical Services and report the problem. You do not need to exit Stata or Mata. Mata stopped doing what was leading to a problem.

## Also see

[M-5] **error()** — Issue error message

[M-2] **intro** — Language definition

# Title

[M-2] **exp** — Expressions

## Syntax

*exp*

## Description

*exp* is used in syntax diagrams to mean “any valid expression may appear here”. Expressions can range from being simple constants

```
2
"this"
3+2i
```

to being names of variables

```
A
beta
varwithverylongname
```

to being a full-fledged scalar, string, or matrix expression:

```
sqrt(2)/2
substr(userinput, 15, strlen(otherstr))
conj(X)'X
```

## Remarks

Remarks are presented under the following headings:

- What’s an expression*
- Assignment suppresses display, as does (void)*
- The pieces of an expression*
- Numeric literals*
- String literals*
- Variable names*
- Operators*
- Functions*

## What’s an expression

Everybody knows what an expression is: expressions are things like 2+3 and `invsym(X’X)*X’y`. Simpler things are also expressions, such as numeric constants

2 is an expression

and string literals

"hi there" is an expression



and function calls:

`sqrt(2)` is an expression

Even when functions do not return anything (the function is void), the code that causes the function to run is an expression. For instance, the function `swap()` (see [M-5] `swap()`) interchanges the contents of its arguments and returns nothing. Even so,

`swap(A, B)` is an expression

## Assignment suppresses display, as does (void)

The equal sign assigns the result of an expression to a variable. For instance,

```
a = 2 + 3
```

assigns 5 to `a`. When the result of an expression is not assigned to a variable, the result is displayed at the terminal. This is true of expressions entered interactively and of expressions coded in programs. For instance, given the program

```
function example(a, b)
{
    "the answer is"
    a+b
}
```

executing `example()` produces

```
: example(2, 3)
  the answer is
  5
```

The fact that 5 appeared is easy enough to understand; we coded the expression `a+b` without assigning it to another variable. The fact that “the answer is” also appeared may surprise you. Nevertheless, we coded “the answer is” in our program, and that is an example of an expression, and because we did not assign the expression to a variable, it was displayed.

In programming situations, there will be times when you want to execute a function—call it `setup()`—but do not care what the function returns, even though the function itself is not void (that is, it returns something). If you code

```
function example(...)
{
    ...
    setup(...)
    ...
}
```

the result will be to display what `setup()` returns. You have two alternatives. You could assign the result of `setup` to a variable even though you will subsequently not use the variable

```
function example(...)
{
    ...
    result = setup(...)
    ...
}
```

or you could cast the result of the function to be void:

```
function example(...)
{
    ...
    (void) setup(...)
    ...
}
```

Placing (void) in front of an expression prevents the result from being displayed.

## The pieces of an expression

Expressions comprise

- numeric literals
- string literals
- variable names
- operators
- functions

## Numeric literals

Numeric literals are just numbers

```
2
3.14159
-7.2
5i
1.213e+32
1.213E+32
1.921fb54442d18X+001
1.921fb54442d18x+001
.
.a
.b
```

but you can suffix an *i* onto the end to mean imaginary, such as *5i* above. To create complex numbers, you combine real and imaginary numbers using the + operator, as in *2+5i*. In any case, you can put the *i* on the end of any literal, so *1.213e+32i* is valid, as is *1.921fb54442d18X+001i*.

*1.921fb54442d18X+001i* is a formidable-looking beast, with or without the *i*.

*1.921fb54442d18X+001* is a way of writing floating-point numbers in binary; it is described in [U] 12.5 **Formats: Controlling how data are displayed**. Most people never use it.

Also, numeric literals include Stata's missing values, *.*, *.a*, *.b*, ..., *.z*.

Complex variables may contain missing just as real variables may, but they get only one: *.a+.bi* is not allowed. A complex variable contains a valid complex value, or it contains *.*, *.a*, *.b*, ..., *.z*.

## String literals

String literals are enclosed in double quotes or in compound double quotes:

```
"the answer is"
"a string"
'also a string'
"The "factor" of a matrix"
""
'""'
```

Strings in Mata contain between 0 and 2,147,483,647 characters. "" or '""' is how one writes the 0-length string.

Any ASCII character may appear in the string, but no provision is provided for typing the unprintable ASCII characters into the string literal. Instead, you use the `char()` function; see [M-5] [ascii\(\)](#). For instance, `char(13)` is carriage return, so the expression

```
"my string" + char(13)
```

produces “my string” followed by a carriage return.

No character is given a special interpretation. In particular, backslash (\) is given no special meaning by Mata. The string literal `"my string\n"` is just that: the characters “my string” followed by a backslash followed by an “n”. Some functions, such as `printf()` (see [M-5] [printf\(\)](#)), give a special meaning to the two-character sequence `\n`, but that special interpretation is a property of the function, not Mata, and is noted in the function’s documentation.

Strings are not zero (null) terminated in Mata. Mata knows that the string `"hello"` is of length 5, but it does not achieve that knowledge by padding a binary 0 as the string’s fifth character. Thus strings may be used to hold binary information.

Although Mata gives no special interpretation to binary 0, some Mata functions do. For instance, `strmatch(s, pattern)` returns 1 if *s* matches *pattern* and 0 otherwise; see [M-5] [strmatch\(\)](#). For this function, both strings are considered to end at the point they contain a binary 0, if they contain a binary 0. Most strings do not, and then the function considers the entire string. In any case, if there is special treatment of binary 0, that is on a function-by-function basis, and a note of that is made in the function’s documentation.

## Variable names

Variable names are just that. Names are case sensitive and no abbreviations are allowed:

```
X
x
MyVar
VeryLongVariableNameForUseInMata
MyVariable
```

The maximum length of a variable name is 32 characters.

# Operators

## Operators, listed by precedence, low to high

Operator	Operator name	Documentation
$a = b$	assignment	[M-2] <a href="#">op_assignment</a>
$a ? b : c$	conditional	[M-2] <a href="#">op_conditional</a>
$a \setminus b$	column join	[M-2] <a href="#">op_join</a>
$a :: b$	column to	[M-2] <a href="#">op_range</a>
$a , b$	row join	[M-2] <a href="#">op_join</a>
$a . . b$	row to	[M-2] <a href="#">op_range</a>
$a :  b$	e.w. or	[M-2] <a href="#">op_colon</a>
$a   b$	or	[M-2] <a href="#">op_logical</a>
$a :& b$	e.w. and	[M-2] <a href="#">op_colon</a>
$a \& b$	and	[M-2] <a href="#">op_logical</a>
$a :== b$	e.w. equal	[M-2] <a href="#">op_colon</a>
$a == b$	equal	[M-2] <a href="#">op_logical</a>
$a :>= b$	e.w. greater than or equal	[M-2] <a href="#">op_colon</a>
$a >= b$	greater than or equal	[M-2] <a href="#">op_logical</a>
$a :<= b$	e.w. less than or equal	[M-2] <a href="#">op_colon</a>
$a <= b$	less than or equal	[M-2] <a href="#">op_logical</a>
$a :< b$	e.w. less than	[M-2] <a href="#">op_colon</a>
$a < b$	less than	[M-2] <a href="#">op_logical</a>
$a :> b$	e.w. greater than	[M-2] <a href="#">op_colon</a>
$a > b$	greater than	[M-2] <a href="#">op_logical</a>
$a :!= b$	e.w. not equal	[M-2] <a href="#">op_colon</a>
$a != b$	not equal	[M-2] <a href="#">op_logical</a>
$a :+ b$	e.w. addition	[M-2] <a href="#">op_colon</a>
$a + b$	addition	[M-2] <a href="#">op_arith</a>
$a :- b$	e.w. subtraction	[M-2] <a href="#">op_colon</a>
$a - b$	subtraction	[M-2] <a href="#">op_arith</a>
$a :* b$	e.w. multiplication	[M-2] <a href="#">op_colon</a>
$a * b$	multiplication	[M-2] <a href="#">op_arith</a>
$a \# b$	Kronecker	[M-2] <a href="#">op_kronecker</a>
$a :/ b$	e.w. division	[M-2] <a href="#">op_colon</a>
$a / b$	division	[M-2] <a href="#">op_arith</a>
$-a$	negation	[M-2] <a href="#">op_arith</a>
$a :^ b$	e.w. power	[M-2] <a href="#">op_colon</a>
$a ^ b$	power	[M-2] <a href="#">op_arith</a>
$a'$	transposition	[M-2] <a href="#">op_transpose</a>
$*a$	contents of	[M-2] <a href="#">pointers</a>
$\&a$	address of	[M-2] <a href="#">pointers</a>
$!a$	not	[M-2] <a href="#">op_logical</a>
$a[exp]$	subscript	[M-2] <a href="#">subscripts</a>
$a[ exp ]$	range subscript	[M-2] <a href="#">subscripts</a>
$a++$	increment	[M-2] <a href="#">op_increment</a>
$a--$	decrement	[M-2] <a href="#">op_increment</a>
$++a$	increment	[M-2] <a href="#">op_increment</a>
$--a$	decrement	[M-2] <a href="#">op_increment</a>

(e.w. = elementwise)

---

## Functions

Functions supplied with Mata are documented in [\[M-5\]](#). An index to the functions can be found in [\[M-4\] intro](#).

## Reference

Gould, W. W. 2006. [Mata Matters: Precision](#). *Stata Journal* 6: 550–560.

## Also see

[\[M-2\] intro](#) — Language definition

# Title

[M-2] **for** — for (*exp1*; *exp2*; *exp3*) *stmt*

## Syntax

```
for (exp1; exp2; exp3) stmt
```

```
for (exp1; exp2; exp3) {  
    stmts  
}
```

where *exp1* and *exp3* are optional, and *exp2* must evaluate to a real scalar.

## Description

for is equivalent to

```
    exp1  
while (exp2) {  
    stmt(s)  
    exp3  
}
```

*stmt(s)* is executed zero or more times. The loop continues as long as *exp2* is not equal to zero.

## Remarks

To understand for, enter the following program

```
function example(n)  
{  
    for (i=1; i<=n; i++) {  
        printf("i=%g\n", i)  
    }  
    printf("done\n")  
}
```

and run `example(3)`, `example(2)`, `example(1)`, `example(0)`, and `example(-1)`.

Common uses of for include

```
for (i=1; i<=rows(A); i++) {  
    for (j=1; j<=cols(A); j++) {  
        ...  
    }  
}
```

## Also see

[M-2] [semicolons](#) — Use of semicolons

[M-2] [do](#) — do ... while (exp)

[M-2] [while](#) — while (exp) stmt

[M-2] [break](#) — Break out of for, while, or do loop

[M-2] [continue](#) — Continue with next iteration of for, while, or do loop

[M-2] [intro](#) — Language definition

# Title

[M-2] **ftof** — Passing functions to functions

## Syntax

*example(..., &somefunction(), ...)*

where *example()* is coded

```
function example(..., f, ...)
{
    ...
    (*f)(...)
    ...
}
```

## Description

Functions can receive other functions as arguments.

Below is described (1) how to call a function that receives a function as an argument and (2) how to write a function that receives a function as an argument.

## Remarks

Remarks are presented under the following headings:

[\*Passing functions to functions\*](#)

[\*Writing functions that receive functions, the simplified convention\*](#)

[\*Passing built-in functions\*](#)

## Passing functions to functions

Someone has written a program that receives a function as an argument. We will imagine that function is

*real scalar* `fderiv(function(), x)`

and that `fderiv()` numerically evaluates the derivative of *function()* at *x*. The documentation for `fderiv()` tells you to write a function that takes one argument and returns the evaluation of the function at that argument, such as

```
real scalar expratio(real scalar x)
{
    return(exp(x)/exp(-x))
}
```

To call `fderiv()` and have it evaluate the derivative of `expratio()` at 3, you code

```
fderiv(&expratio(), 3)
```

To pass a function to a function, you code `&` in front of the function's name and `()` after. Coding `&expratio()` passes the address of the function `expratio()` to `fderiv()`.



## Writing functions that receive functions, the simplified convention

To receive a function, you include a variable among the program arguments to receive the function—we will use *f*—and you then code `(*f)(...)` to call the passed function. The code for `fderiv()` might read

```
function fderiv(f, x)
{
    return( ((*f)(x+1e-6) - (*f)(x)) / 1e-6 )
}
```

or, if you prefer to be explicit about your declarations,

```
real scalar fderiv(pointer scalar f, real scalar x)
{
    return( ((*f)(x+1e-6) - (*f)(x)) / 1e-6 )
}
```

or, if you prefer to be even more explicit:

```
real scalar fderiv(pointer(real scalar function) scalar f,
                      real scalar x)
{
    return( ((*f)(x+1e-6) - (*f)(x)) / 1e-6 )
}
```

In any case, using pointers, you type `(*f)(...)` to execute the function passed. See [\[M-2\] pointers](#) for more information.

Aside: the function `fderiv()` would work but, because of the formula it uses, would return very inaccurate results.

## Passing built-in functions

You cannot pass built-in functions to other functions. For instance, [\[M-5\] `exp\(\)`](#) is built in, which is revealed by [\[M-3\] mata which](#):

```
: mata which exp()
exp(): built-in
```

Not all official functions are built in. Many are implemented in Mata as library functions, but `exp()` is built in and coding `&exp()` will result in an error. If you wanted to pass `exp()` to a function, create your own version of it

```
: function myexp(x) return(exp(x))
```

and then pass `&myexp()`.

## Also see

[\[M-2\] intro](#) — Language definition

# Title

[M-2] goto — goto label

## Syntax

```
label:  ...  
      ...  
      goto label
```

where *label*: may occur before or after the *goto*.

## Description

*goto label* causes control to pass to the statement following *label*:. *label* may be any name up to eight characters long.

## Remarks

These days, good style is to avoid using *goto*.

*goto* is useful when translating a FORTRAN program, such as

```
A = 4.0e0/3.0e0  
10 B = A - 1.0e0  
   C = B + B + B  
   EPS = DABS(C - 1.0e0)  
   if (EPS.EQ.0.0e0) GOTO 10
```

The Mata translation is

```
      a = 4/3  
s10:  b = a - 1  
      c = b + b + b  
      eps = abs(c-1)  
      if (eps==0) goto s10
```

although

```
a = 4/3  
do {  
    b = a - 1  
    c = b + b + b  
    eps = abs(c - 1)  
} while (eps==0)
```

is more readable.

## Reference

Gould, W. W. 2005. [Mata Matters: Translating Fortran](#). *Stata Journal* 5: 421–441.

## Also see

[M-2] **do** — do ... while (exp)

[M-2] **for** — for (exp1; exp2; exp3) stmt

[M-2] **while** — while (exp) stmt

[M-2] **break** — Break out of for, while, or do loop

[M-2] **continue** — Continue with next iteration of for, while, or do loop

[M-2] **intro** — Language definition

# Title

[M-2] **if** — `if (exp) ... else ...`

## Syntax

```
if (exp) stmt1

if (exp) stmt1
else stmt2

if (exp) {
    stmts1
}
else {
    stmts2
}
if (exp1) ...
else if (exp2) ...
else if (exp3) ...
...
else ...
```

where *exp*, *exp*<sub>1</sub>, *exp*<sub>2</sub>, *exp*<sub>3</sub>, ... must evaluate to real scalars.

## Description

`if` evaluates the expression, and if it is true (evaluates to a nonzero number), `if` executes the statement or statement block that immediately follows it; otherwise, `if` skips the statement or block.

`if ... else` evaluates the expression, and if it is true (evaluates to a nonzero number), `if` executes the statement or statement block that immediately follows it and skips the statement or statement block following the `else`; otherwise, it skips the statement or statement block immediately following it and executes the statement or statement block following the `else`.

## Remarks

if followed by multiple `elses` is interpreted as being nested, that is,

```
if (exp1) ...  
else if (exp2) ...  
else if (exp3) ...  
...  
else ...
```

is equivalent to

```
if (exp1) ...  
else {  
    if (exp2) ...  
    else {  
        if (exp3) ...  
        else {  
            ...  
        }  
    }  
}
```

## Also see

[\[M-2\] intro](#) — Language definition

# Title

[M-2] op\_arith — Arithmetic operators

## Syntax

$a + b$	addition
$a - b$	subtraction
$a * b$	multiplication
$a / b$	division
$a ^ b$	power
$-a$	negation

where  $a$  and  $b$  may be numeric scalars, vectors, or matrices.

## Description

The above operators perform basic arithmetic.

## Remarks

Also see [M-2] op\_colon for the `:+`, `:-`, `:*`, and `:/` operators. Colon operators have relaxed conformability restrictions.

The `*` and `:*` multiplication operators can also perform string duplication—`3*"a" = "aaa"`—see [M-5] strdup().

## Conformability

$a + b, a - b:$				
	$a:$	$r \times c$		
	$b:$	$r \times c$		
	$result:$	$r \times c$		
$a * b:$				
	$a:$	$k \times n$	$k \times n$	$1 \times 1$
	$b:$	$n \times m$	$1 \times 1$	$n \times m$
	$result:$	$k \times m$	$k \times n$	$n \times m$
$a / b:$				
	$a:$	$r \times c$		
	$b:$	$1 \times 1$		
	$result:$	$r \times c$		
$a ^ b:$				
	$a:$	$1 \times 1$		
	$b:$	$1 \times 1$		
	$result:$	$1 \times 1$		

$-a$ :

$a$ :	$r \times c$
$result$ :	$r \times c$

## Diagnostics

All operators return missing when arguments are missing.

$a*b$  with  $a$ :  $k \times 0$  and  $b$ :  $0 \times m$  returns a  $k \times m$  matrix of zeros.

$a/b$  returns missing when  $b==0$  or when  $a/b$  would result in overflow.

$a^b$  returns a real when both  $a$  and  $b$  are real; thus,  $(-4)^.5$  evaluates to missing, whereas  $(-4+0i)^.5$  evaluates to  $2i$ .

$a^b$  returns missing on overflow.

## Also see

[\[M-2\] exp](#) — Expressions

[\[M-2\] intro](#) — Language definition

# Title

[M-2] **op\_assignment** — Assignment operator

## Syntax

*lval* = *exp*

where *exp* is any valid expression and where *lval* is

*name*

*name*[*exp*]

*name*[*exp*, *exp*]

*name*[|*exp*|]

In pointer use (advanced), *name* may be

\**lval*

\*(*lval*)

\*(*lval*[*exp*])

\*(*lval*[*exp*, *exp*])

\*(*lval*[|*exp*|])

in addition to being a variable name.

## Description

= assigns the evaluation of *exp* to *lval*.

Do not confuse the = assignment operator with the == equality operator. Coding

x = y

assigns the value of y to x. Coding

if (x==y) ...

(note doubled equal signs)

performs the action if the value of x is equal to the value of y. See [M-2] **op\_logical** for a description of the == equality operator.

If the result of an expression is not assigned to a variable, then the result is displayed at the terminal; see [M-2] **exp**.

## Remarks

Remarks are presented under the following headings:

*Assignment suppresses display*

*The equal-assignment operator*

*lvals, what appears on the left-hand side*

*Row, column, and element lvals*

*Pointer lvals*



## Assignment suppresses display

When you interactively enter an expression or code an expression in a program without the equal-assignment operator, the result of the expression is displayed at the terminal:

```
: 2 + 3
5
```

When you assign the expression to a variable, the result is not displayed:

```
: x = 2 + 3
```

## The equal-assignment operator

Equals is an operator, so in addition to coding

```
a = 2 + 3
```

you can code

```
a = b = 2 + 3
```

or

```
y = x / (denominator = sqrt(a+b))
```

or even

```
y1 = y2 = x / (denominator = sqrt(sum=a+b))
```

This last is equivalent to

```
sum = a + b
denominator = sqrt(sum)
y2 = x / denominator
y1 = y2
```

Equals binds weakly, so

```
a = b = 2 + 3
```

is interpreted as

```
a = b = (2 + 3)
```

and not

```
a = (b=2) + 3
```

## lvals, what appears on the left-hand side

What appears to the left of the equals is called an *lval*, short for left-hand-side value. It would make no sense, for instance, to code

```
sqrt(4) = 3
```

and, as a matter of fact, you are not allowed to code that because `sqrt(4)` is not an *lval*:

```
: sqrt(4) = 3
invalid lval
r(3000);
```

An *lval* is anything that can hold values. A scalar can hold values

```
a = 3
x = sqrt(4)
```

a matrix can hold values

```
A = (1, 2 \ 3, 4)
B = invsym(C)
```

a matrix row can hold values

```
A[1,.] = (7, 8)
```

a matrix column can hold values

```
A[:,2] = (9 \ 10)
```

and finally, a matrix element can hold a value

```
A[1,2] = 7
```

*lvals* are usually one of the above forms. The other forms have to do with pointer variables, which most programmers never use; they are discussed under [Pointer lvals](#) below.

## Row, column, and element lvals

When you assign to a row, column, or element of a matrix,

```
A[1,.] = (7, 8)
A[:,2] = (9 \ 10)
A[1,2] = 7
```

the row, column, or element must already exist:

```
: A = (1, 2 \ 3, 4)
: A[3,4] = 4
<istmt>: 3301 subscript invalid
r(3301);
```

This is usually not an issue because, by the time you are assigning to a row, column, or element, the matrix has already been created, but in the event you need to create it first, use the `J()` function; see [\[M-5\] J\(\)](#). The following code fragment creates a  $3 \times 4$  matrix containing the sum of its indices:

```
A = J(3, 4, .)
for (i=1; i<=3; i++) {
    for (j=1; j<=4; j++) A[i,j] = i + j
}
```

## Pointer lvals

In addition to the standard *lvals*

```
A = (1, 2 \ 3, 4)
A[1, .] = (7, 8)
A[. , 2] = (9 \ 10)
A[1, 2] = 7
```

pointer *lvals* are allowed. For instance,

```
*p = 3
```

stores 3 in the address pointed to by pointer scalar *p*.

```
(*q)[1, 2] = 4
```

stores 4 in the (1,2) element of the address pointed to by pointer scalar *q*, whereas

```
*Q[1, 2] = 4
```

stores 4 in the address pointed to by the (1,2) element of pointer matrix *Q*.

```
*Q[2, 1][1, 3] = 5
```

is equivalent to

```
*(Q[2, 1])[1, 3] = 5
```

and stores 5 in the (1,3) element of the address pointed to by the (2,1) element of pointer matrix *Q*.

Pointers to pointers, pointers to pointers to pointers, etc., are also allowed. For instance,

```
**r = 3
```

stores 3 in the address pointed to by the address pointed to by pointer scalar *r*, whereas

```
*((*(Q[1, 2]))[2, 1])[3, 4] = 7
```

stores 7 in the (3,4) address pointed to by the (2,1) address pointed to by the (1,2) address of pointer matrix *Q*.

## Conformability

$a = b$ :

*input*:

$b: \quad r \times c$

*output*:

$a: \quad r \times c$

## Diagnostics

$a = b$  aborts with error if there is insufficient memory to store a copy of *b* in *a*.

## Also see

[M-5] `swap()` — Interchange contents of variables

[M-2] `exp` — Expressions

[M-2] `intro` — Language definition

## Syntax

$a : + b$	addition
$a : - b$	subtraction
$a : * b$	multiplication
$a : / b$	division
$a : ^ b$	power
$a : == b$	equality
$a : != b$	inequality
$a : > b$	greater than
$a : >= b$	greater than or equal to
$a : < b$	less than
$a : <= b$	less than or equal to
$a : \& b$	and
$a :   b$	or

## Description

Colon operators perform element-by-element operations.

## Remarks

Remarks are presented under the following headings:

*C-conformability: element by element*  
*Usefulness of colon logical operators*  
*Use parentheses*

## C-conformability: element by element

The colon operators perform the indicated operation on each pair of elements of  $a$  and  $b$ . For instance,

$$\begin{bmatrix} c & d \\ f & g \\ h & i \end{bmatrix} : * \begin{bmatrix} j & k \\ l & m \\ n & o \end{bmatrix} = \begin{bmatrix} c * j & d * k \\ f * l & g * m \\ h * n & i * o \end{bmatrix}$$

Also colon operators have a relaxed definition of conformability:

$$\begin{aligned} \begin{bmatrix} c \\ f \\ g \end{bmatrix} : * \begin{bmatrix} j & k \\ l & m \\ n & o \end{bmatrix} &= \begin{bmatrix} c * j & c * k \\ f * l & f * m \\ g * n & g * o \end{bmatrix} \\ \begin{bmatrix} c & d \\ f & g \\ h & i \end{bmatrix} : * \begin{bmatrix} j \\ l \\ n \end{bmatrix} &= \begin{bmatrix} c * j & d * j \\ f * l & g * l \\ h * n & i * n \end{bmatrix} \\ [c \ d] : * \begin{bmatrix} j & k \\ l & m \\ n & o \end{bmatrix} &= \begin{bmatrix} c * j & d * k \\ c * l & d * m \\ c * n & d * o \end{bmatrix} \\ \begin{bmatrix} c & d \\ f & g \\ h & i \end{bmatrix} : * [l \ m] &= \begin{bmatrix} c * l & d * m \\ f * l & g * m \\ h * l & i * m \end{bmatrix} \\ c : * \begin{bmatrix} j & k \\ l & m \\ n & o \end{bmatrix} &= \begin{bmatrix} c * j & c * k \\ c * l & c * m \\ c * n & c * o \end{bmatrix} \\ \begin{bmatrix} c & d \\ f & g \\ h & i \end{bmatrix} : * j &= \begin{bmatrix} c * j & d * j \\ f * j & g * j \\ h * j & i * j \end{bmatrix} \end{aligned}$$

The matrices above are said to be c-conformable; the *c* stands for colon. The matrices have the same number of rows and columns, or one or the other is a vector with the same number of rows or columns as the matrix, or one or the other is a scalar.

C-conformability is relaxed, but not everything is allowed. The following is an error:

$$(c \ d \ e) : * \begin{bmatrix} f \\ g \\ h \end{bmatrix}$$

Usefulness of colon logical operators

It is worth paying particular attention to the colon logical operators because they can produce pattern vectors and matrices. Consider the matrix

: x = (5, 0 \ 0, 2 \ 3, 8)

: x

12

150

202

338

Which elements of  $x$  contain 0?

```

: x==0
      1  2
1  [ 0  1
2  [ 1  0
3  [ 0  0

```

How many zeros are there in  $x$ ?

```

: sum(x==0)
      2

```

## Use parentheses

Because of their relaxed conformability requirements, colon operators are not associative even when the underlying operator is. For instance, you expect  $(a+b)+c == a+(b+c)$ , at least ignoring numerical roundoff error. Nevertheless,  $(a:+b):+c == a:+(b:+c)$  does not necessarily hold. Consider what happens when

```

a:      1 × 4
b:      5 × 1
c:      5 × 4

```

Then  $(a:+b):+c$  is an error because  $a:+b$  is not  $c$ -conformable.

Nevertheless,  $a:+(b:+c)$  is not an error and in fact produces a  $5 \times 4$  matrix because  $b:+c$  is  $5 \times 4$ , which is  $c$ -conformable with  $a$ .

## Conformability

```

a :op b:
a:      r1 × c1
b:      r2 × c2,  a and b c-conformable
result:  max(r1,r2) × max(c1,c2)

```

## Diagnostics

The colon operators return missing and abort with error under the same conditions that the underlying operator returns missing and aborts with error.

## Also see

[\[M-2\] exp](#) — Expressions

[\[M-2\] intro](#) — Language definition

# Title

[M-2] op\_conditional — Conditional operator

## Syntax

$a \text{ ? } b : c$

where  $a$  must evaluate to a real scalar, and  $b$  and  $c$  may be of any type whatsoever.

## Description

The conditional operator returns  $b$  if  $a$  is true ( $a$  is not equal to 0) and  $c$  otherwise.

## Remarks

Conditional operators

$\text{dof} = (\text{k}==0 \text{ ? } \text{n}-1 : \text{n}-\text{k})$

are more compact than the `if-else` alternative

```
if (k==0) dof = n-1
else      dof = n-k
```

and they can be used as parts of expressions:

$\text{mse} = \text{ess}/(\text{k}==0 \text{ ? } \text{n}-1 : \text{n}-\text{k})$

## Conformability

$a \text{ ? } b : c$ :

$a$ :	$1 \times 1$	
$b$ :	$r_1 \times c_1$	
$c$ :	$r_2 \times c_2$	
$result$ :	$r_1 \times c_1$	or $r_2 \times c_2$

## Diagnostics

In  $a \text{ ? } b : c$ , only the necessary parts are evaluated:  $a$  and  $b$  if  $a$  is true, or  $a$  and  $c$  if  $a$  is false. However, the `++` and `--` operators are always evaluated:

$(\text{k}==0 \text{ ? } i++ : j++)$

increments both  $i$  and  $j$ , regardless of the value of  $k$ .



## Also see

[M-2] [exp](#) — Expressions

[M-2] [intro](#) — Language definition

## Syntax

$++i$	increment before
$--i$	decrement before
$i++$	increment after
$i--$	decrement after

where  $i$  must be a real scalar.

## Description

$++i$  and  $i++$  increment  $i$ ; they perform the operation  $i=i+1$ .  $++i$  performs the operation before the evaluation of the expression in which it appears, whereas  $i++$  performs the operation afterward.

$--i$  and  $i--$  decrement  $i$ ; they perform the operation  $i=i-1$ .  $--i$  performs the operation before the evaluation of the expression in which it appears, whereas  $i--$  performs the operation afterward.

## Remarks

These operators are used in code, such as

```
x[i++] = 2
x[--i] = 3
for (i=0; i<100; i++) {
    ...
}
if (++n > 10) {
    ...
}
```

Where these expressions appear, results are as if the current value of  $i$  were substituted, and in addition,  $i$  is incremented, either before or after the expression is evaluated. For instance,

```
x[i++] = 2
```

is equivalent to

```
x[i] = 2 ; i = i + 1
```

and

```
x[++i] = 3
```

is equivalent to

```
i = i + 1 ; x[i] = 3
```

Coding

```
for (i=0; i<100; i++) {
    ...
}
```

or

```
for (i=0; i<100; ++i) {
    ...
}
```

is equivalent to

```
for (i=0; i<100; i=i+1) {
    ...
}
```

because it does not matter whether the incrementation is performed before or after the otherwise null expression.

```
if (++n > 10) {
    ...
}
```

is equivalent to

```
n = n + 1
if (n > 10) {
    ...
}
```

whereas

```
if (n++ > 10) {
    ...
}
```

is equivalent to

```
if (n > 10) {
    n = n + 1
    ...
}
else    n = n + 1
```

The ++ and -- operators may be used only with real scalars and are usually associated with indexing or counting. They result in fast and readable code.

## Conformability

*++i*, *--i*, *i++*, and *i--*:

<i>i</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

`++` and `--` are allowed with real scalars only. That is, `++i` or `i++` is valid, assuming `i` is a real scalar, but `x[i,j]++` is not valid.

`++` and `--` abort with error if applied to a variable that is not a real scalar.

`++i`, `i++`, `--i`, and `i--` should be the only reference to `i` in the expression. Do not code, for instance,

```
x[i++] = y[i]
x[++i] = y[i]
x[i] = y[i++]
x[i] = y[++i]
```

The value of `i` in the above expressions is formally undefined; whatever is its value, you cannot depend on that value being obtained by earlier or later versions of the compiler. Instead code

```
i++ ; x[i] = y[i]
```

or code

```
x[i] = y[i] ; i++
```

according to the desired outcome.

It is, however, perfectly reasonable to code

```
x[i++] = y[j++]
```

That is, multiple `++` and `--` operators may occur in the same expression; it is multiple references to the target of the `++` and `--` that must be avoided.

## Also see

[\[M-2\] exp](#) — Expressions

[\[M-2\] intro](#) — Language definition

# Title

[M-2] op\_join — Row- and column-join operators

## Syntax

$$a \ , \ b$$

$$a \ \backslash \ b$$

## Description

, and \ are Mata’s row-join and column-join operators.

## Remarks

Remarks are presented under the following headings:

- Comma and backslash are operators
- Comma as a separator
- Warning about the misuse of comma and backslash operators

### Comma and backslash are operators

That , and \ are operators cannot be emphasized enough. When one types

: (1, 2 \ 3, 4)

	1	2
1	1	2
2	3	4

one is tempted to think, “Ah, comma and backslash are how you separate elements when you enter a matrix.” If you think like that, you will not appreciate the power of , and \.

, and \ are operators in the same way that \* and + are operators.

, is the operator that takes a  $r \times c_1$  matrix and a  $r \times c_2$  matrix, and returns a  $r \times (c_1 + c_2)$  matrix.

\ is the operator that takes a  $r_1 \times c$  matrix and a  $r_2 \times c$  matrix, and returns a  $(r_1 + r_2) \times c$  matrix.

, and \ may be used with scalars, vectors, or matrices:

: a = (1 \ 2)

: b = (3 \ 4)

: a, b

	1	2
1	1	3
2	2	4

: c = (1, 2)

```
: d = (3, 4)
: c \ d
      1  2
1     1  2
2     3  4
```

, binds more tightly than \, meaning that  $e, f \backslash g, h$  is interpreted as  $(e, f) \backslash (g, h)$ . In this, , and \ are no different from \* and + operators: \* binds more tightly than + and  $e*f + g*h$  is interpreted as  $(e*f)+(g*h)$ .

Just as it sometimes makes sense to type  $e*(f+g)*h$ , it can make sense to type  $e, (f \backslash g), h$ :

```
: e = 1 \ 2
: f = 5 \ 6
: g = 3
: h = 4
: e, (g\h), f
      1  2  3
1     1  3  5
2     2  4  6
```

Comma as a separator

, has a second meaning in Mata: it is the argument separator for functions. When you type

```
: myfunc(a, b)
```

the comma that appears inside the parentheses is not the comma row-join operator; it is the comma argument separator. If you wanted to call myfunc() with second argument equal to row vector (1,2), you must type

```
: myfunc(a, (1,2))
```

and not

```
: myfunc(a, 1, 2)
```

because otherwise Mata will think you are trying to pass three arguments to myfunc(). When you open another set of parentheses inside a function's argument list, comma reverts back to its usual row-join meaning.

Warning about the misuse of comma and backslash operators

Misuse or mere overuse of , and \ can substantially reduce the speed with which your code executes. Consider the actions Mata must take when you code, say,

```
a \ b
```

First, Mata must allocate a matrix or vector containing rows(a)+rows(b) rows, then it must copy a into the new matrix or vector, and then it must copy b. Nothing inefficient has happened yet, but now consider

```
(a \ b) \ c
```

Picking up where we left off, Mata must allocate a matrix or vector containing  $\text{rows}(a) + \text{rows}(b) + \text{rows}(c)$  rows, then it must copy  $(a \setminus b)$  into the new matrix or vector, and then it must copy  $c$ . Something inefficient just happened:  $a$  was copied twice!

Coding

$$res = (a \setminus b) \setminus c$$

is convenient, but execution would be quicker if we coded

```
res = J(rows(a)+rows(b)+rows(c), cols(a), .)
res[1,.] = a
res[2,.] = b
res[3,.] = c
```

We do not want to cause you concern where none is due. In general, you would not be able to measure the difference between the more efficient code and coding  $res = (a \setminus b) \setminus c$ . But as the number of row or column operators stack up, the combined result becomes more and more inefficient. Even that is not much of a concern. If the inefficient construction itself is buried in a loop, however, and that loop is executed thousands of times, the inefficiency can become important.

With a little thought, you can always substitute predeclaration using `J()` (see [M-5] `J()`) and assignment via subscripting.

## Conformability

$a, b$ :

$a$ :	$r \times c_1$
$b$ :	$r \times c_2$
$result$ :	$r \times (c_1 + c_2)$

$a \setminus b$ :

$a$ :	$r_1 \times c$
$b$ :	$r_2 \times c$
$result$ :	$(r_1 + r_2) \times c$

## Diagnostics

, and  $\setminus$  abort with error if  $a$  and  $b$  are not of the same broad type.

## Also see

[M-2] [exp](#) — Expressions

[M-2] [intro](#) — Language definition

# Title

[M-2] op\_kronecker — Kronecker direct-product operator

## Syntax

$$A\#B$$

where  $A$  and  $B$  may be real or complex.

## Description

$A\#B$  returns the Kronecker direct product.

$\#$  binds tightly:  $X\#A\#B\#Y$  is interpreted as  $X\#(A\#B)\#Y$ .

## Remarks

The Kronecker direct product is also known as the Kronecker product, the direct product, the tensor product, and the outer product.

The Kronecker product  $A\#B$  is the matrix  $||a_{ij}\#B||$ .

## Conformability

$A\#B$ :	$A$ :	$r_1 \times c_1$
	$B$ :	$r_2 \times c_2$
	result:	$r_1\#r_2 \times c_1\#c_2$

## Diagnostics

None.

Leopold Kronecker (1823–1891) was born in Liegnitz, Prussia (now Legnica, Poland), to a well-off family. He attended the universities of Berlin, Bonn, and Breslau before completing a doctorate on the complex roots of unity. For several years, Kronecker devoted himself to business interests while working on mathematics in his spare time, publishing particularly in number theory, elliptic functions, and the theory of equations. He later started giving lectures at the university in Berlin, as was his right as a member of the Academy of Science. In 1883, he was appointed as the chair. Kronecker came to believe that mathematical arguments should involve only finite numbers and a finite number of operations, which led to increasing mathematical and personal disagreements with those who worked on irrational numbers or nonconstructive existence proofs.



## Reference

James, I. 2002. *Remarkable Mathematicians: From Euler to von Neumann*. Cambridge: Cambridge University Press.

## Also see

[M-2] [exp](#) — Expressions

[M-2] [intro](#) — Language definition

## Syntax

$a == b$	true if $a$ equals $b$
$a != b$	true if $a$ not equal to $b$
$a > b$	true if $a$ greater than $b$
$a >= b$	true if $a$ greater than or equal to $b$
$a < b$	true if $a$ less than $b$
$a <= b$	true if $a$ less than or equal to $b$
$!a$	logical negation; true if $a==0$ and false otherwise
$a \& b$	true if $a!=0$ and $b!=0$
$a   b$	true if $a!=0$ or $b!=0$
$a \&\& b$	synonym for $a \& b$
$a    b$	synonym for $a   b$

## Description

The operators above perform logical comparisons, and operator `!` performs logical negation. All operators evaluate to 1 or 0, meaning true or false.

## Remarks

Remarks are presented under the following headings:

[Introduction](#)

[Use of logical operators with pointers](#)

### Introduction

The operators above work as you would expect when used with scalars, and the comparison operators and the not operator have been generalized for use with matrices.

$a==b$  evaluates to true if  $a$  and  $b$  are p-conformable, of the same type, and the corresponding elements are equal. Of the same type means  $a$  and  $b$  are both numeric, both strings, or both pointers. Thus it is not an error to ask if a  $2 \times 2$  matrix is equal to a  $4 \times 1$  vector or if a string variable is equal to a real variable; they are not. Also  $a==b$  is declared to be true if  $a$  or  $b$  are p-conformable and the number of rows or columns is zero.

$a!=b$  is equivalent to  $!(a==b)$ .  $a!=b$  evaluates to true when  $a==b$  would evaluate to false and evaluates to true otherwise.

The remaining comparison operators  $>$ ,  $>=$ ,  $<$ , and  $<=$  work differently from  $==$  and  $!=$  in that they require  $a$  and  $b$  be p-conformable; if they are not, they abort with error. They return true if the corresponding elements have the stated relationship, and return false otherwise. If  $a$  or  $b$  is complex, the comparison is made in terms of the length of the complex vector; for instance,  $a > b$  is equivalent to  $\text{abs}(a) > \text{abs}(b)$ , and so  $-3 > 2+0i$  is true.

$!a$ , when  $a$  is a scalar, evaluates to 0 if  $a$  is not equal to zero and 1 otherwise. Applied to a vector or matrix, the same operation is carried out, element by element:  $!(-1, 0, 1, 2, \dots)$  evaluates to  $(0, 1, 0, 0, 0, \dots)$ .

$\&$  and  $|$  (*and* and *or*) may be used with scalars only. Because so many people are familiar with programming in the C language, Mata provides  $\&\&$  as a synonym for  $\&$  and  $||$  as a synonym for  $|$ .

## Use of logical operators with pointers

In a pointer expression, NULL is treated as false and all other pointer values (address values) are treated as true. Thus the following code is equivalent

<pre>pointer x ... if (x) {     ... }</pre>	<pre>pointer x ... if (x!=NULL) {     ... }</pre>
---	---

The logical operators  $a==b$ ,  $a!=b$ ,  $a\&b$ , and  $a|b$  may be used with pointers.

## Conformability

$a==b$ ,  $a!=b$ :

$a$ :	$r_1 \times c_1$
$b$ :	$r_2 \times c_2$
<i>result</i> :	$1 \times 1$

$a > b$ ,  $a \geq b$ ,  $a < b$ ,  $a \leq b$ :

$a$ :	$r \times c$
$b$ :	$r \times c$
<i>result</i> :	$1 \times 1$

$!a$ :

$a$ :	$r \times c$
<i>result</i> :	$r \times c$

$a\&b$ ,  $a|b$ :

$a$ :	$1 \times 1$
$b$ :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

$a==b$  and  $a!=b$  cannot fail.

$a > b$ ,  $a \geq b$ ,  $a < b$ ,  $a \leq b$  abort with error if  $a$  and  $b$  are not p-conformable, if  $a$  and  $b$  are not of the same general type (numeric and numeric or string and string), or if  $a$  or  $b$  are pointers.

`!a` aborts with error if  $a$  is not real.

`a&b` and `a|b` abort with error if  $a$  and  $b$  are not both real or not both pointers. If  $a$  and  $b$  are pointers, pointer value `NULL` is treated as false and all other pointer values are treated as true. In all cases, a real equal to 0 or 1 is returned.

## Also see

[M-2] [exp](#) — Expressions

[M-2] [intro](#) — Language definition

## Syntax

$a..b$	row range
$a::b$	column range

## Description

The range operators create vectors that count from  $a$  to  $b$ .

$a..b$  returns a row vector.

$a::b$  returns a column vector.

## Remarks

$a..b$  and  $a::b$  count from  $a$  up to but not exceeding  $b$ , incrementing by 1 if  $b \geq a$  and by  $-1$  if  $b < a$ .

$1..4$  creates row vector  $(1,2,3,4)$ .

$1::4$  creates column vector  $(1\backslash 2\backslash 3\backslash 4)$ .

$-1..-4$  creates row vector  $(-1,-2,-3,-4)$ .

$-1::-4$  creates column vector  $(-1\backslash -2\backslash -3\backslash -4)$ .

$1.5..4.5$  creates row vector  $(1.5, 2.5, 3.5, 4.5)$ .

$1.5::4.5$  creates column vector  $(1.5\backslash 2.5\backslash 3.5\backslash 4.5)$ .

$1.5..4.4$  creates row vector  $(1.5, 2.5, 3.5)$ .

$1.5::4.4$  creates column vector  $(1.5\backslash 2.5\backslash 3.5)$ .

$-1.5..-4.4$  creates row vector  $(-1.5, -2.5, -3.5)$ .

$-1.5::-4.4$  creates column vector  $(-1.5\backslash -2.5\backslash -3.5)$ .

$1..1$  and  $1::1$  both return  $(1)$ .

# Conformability

$a \dots b$		
	$a$ :	$1 \times 1$
	$b$ :	$1 \times 1$
	<i>result</i> :	$1 \times \text{trunc}(\text{abs}(b - a)) + 1$
$a : : b$		
	$a$ :	$1 \times 1$
	$b$ :	$1 \times 1$
	<i>result</i> :	$\text{trunc}(\text{abs}(b - a)) + 1 \times 1$

# Diagnostics

$a \dots b$  and  $a : : b$  return missing if  $a \geq .$  or  $b \geq ..$

# Also see

- [M-2] [exp](#) — Expressions
- [M-2] [intro](#) — Language definition

# Title

[M-2] op\_transpose — Conjugate transpose operator

## Syntax

$A'$

## Description

$A'$  returns the [transpose](#) of  $A$  or, if  $A$  is complex, the conjugate transpose.

## Remarks

The  $'$  postfix operator may be used on any type of matrix or vector: real, complex, string, or pointer:

: a

123

123

456

: a'

12

14

25

36

: s

12

alpha

beta

: s'

1

alpha

beta

: p

1

0x1d20e58

0x1b9a368

: p'

12

0x1d20e58

0x1b9a368

```
: z
      1      2
1  1 + 2i  3 + 4i
2  5 + 6i  7 + 8i

: z'
      1      2
1  1 - 2i  5 - 6i
2  3 - 4i  7 - 8i
```

When `'` is applied to a complex, returned is the conjugate transpose. If you do not want this, code `conj(z')` or `conj(z)'`—it makes no difference; see [M-5] `conj()`,

```
: conj(z')
      1      2
1  1 + 2i  5 + 6i
2  3 + 4i  7 + 8i
```

Or use the `transposeonly()` function; see [M-5] `transposeonly()`:

```
: transposeonly(z)
      1      2
1  1 + 2i  5 + 6i
2  3 + 4i  7 + 8i
```

`transposeonly()` executes slightly faster than `conj(z')`.

For real and complex `A`, also see [M-5] `_transpose()`, which provides a way to transpose a matrix in place and so saves memory.

## Conformability

$A'$ :

$A$ :	$r \times c$
result:	$c \times r$

## Diagnostics

The transpose operator cannot fail, but it is easy to use it incorrectly when working with complex quantities.

A user wanted to form  $A*x$  but when he tried, got a conformability error. He thought  $x$  was a column vector, but it turned out to be a row vector, or perhaps it was the other way around. Anyway, he then coded  $A*x'$ , and the program worked and, even better, produced the correct answers. In his test,  $x$  was real.

Later, the user ran the program with complex  $x$ , and the program generated incorrect results, although it took him a while to notice. Study and study his code he did, before he thought about the innocuous  $A*x'$ . The transpose operator had not only changed  $x$  from being a row into being a column but had taken the conjugate of each element of  $x$ ! He changed the code to read  $A*\text{transposeonly}(x)$ .



The user no doubt wondered why the `'` transpose operator was not defined at the outset to be equivalent to `transposeonly()`. If it had been, then rather than telling the story of the man who was bitten by conjugate transpose when he only wanted the transpose, we would have told the story of the woman who was bitten by the transpose when she needed the conjugate transpose. There are, in fact, more of the latter stories than there are of the former.

## Also see

[M-5] `conj()` — Complex conjugate

[M-5] `_transpose()` — Transposition in place

[M-5] `transposeonly()` — Transposition without conjugation

[M-2] `exp` — Expressions

[M-2] `intro` — Language definition

## Syntax

```
function functionname(|arg [, arg [, ... ]]) { ... }
```

```
function functionname(arg, |arg [, ... ] ) { ... }
```

```
function functionname(arg, arg , |...) { ... }
```

The vertical (or) bar separates required arguments from optional arguments in function declarations. The bar may appear at most once.

## Description

Mata functions may have various numbers of arguments. How you write programs that allow these optional arguments is described below.

## Remarks

Remarks are presented under the following headings:

[What are optional arguments?](#)

[How to code optional arguments](#)

[Examples revisited](#)

## What are optional arguments?

### ► Example 1

You write a function named `ditty()`. Function `ditty()` allows the caller to specify two or three arguments:

```
real matrix ditty(real matrix A, real matrix B, real scalar scale)
```

```
real matrix ditty(real matrix A, real matrix B)
```

If the caller specifies only two arguments, results are as if the caller had specified the third argument equal to missing; that is, `ditty(A, B)` is equivalent to `ditty(A, B, .)`



### ► Example 2

You write function `gash()`. Function `gash()` allows the caller to specify one or two arguments:

```
real matrix gash(real matrix A, real matrix B)
```

```
real matrix gash(real matrix A)
```

If the caller specifies only one argument, results are as if  $J(0,0,.)$  were specified for the second. ◀

### ► Example 3

You write function `easygoing()`. Function `easygoing()` takes three arguments but allows the caller to specify three, two, one, or even no arguments:

```
real scalar easygoing(real matrix A, real matrix B, real scalar scale)
real scalar easygoing(real matrix A, real matrix B)
real scalar easygoing(real matrix A)
real scalar easygoing()
```

If *scale* is not specified, results are as if *scale* = 1 were specified. If *B* is not specified, results are as if *B* = *A* were specified. If *A* is not specified, results are as if *A* =  $I(2)$  were specified. ◀

### ► Example 4

You write function `midsection()`. `midsection()` takes three arguments, but users may specify only two—the first and last—if they wish.

```
real matrix midsection(real matrix A, real vector w, real matrix B)
real matrix midsection(real matrix A, real matrix B)
```

If *w* is not specified, results are as if  $w = J(1, \text{cols}(A), 1)$  was specified. ◀

## How to code optional arguments

When you code

```
function nebulous(a, b, c)
{
    ...
}
```

you are stating that function `nebulous()` requires three arguments. If the caller specifies fewer or more, execution will abort.

If you code

```
function nebulous(a, b, |c)
{
    ...
}
```

you are stating that the last argument is optional. Note the vertical or bar in front of *c*.

If you code

```
function nebulous(a, |b, c)
{
    ...
}
```

you are stating that the last two arguments are optional; the user may specify one, two, or three arguments.

If you code

```
function nebulous(|a, b, c)
{
    ...
}
```

you are stating that all arguments are optional; the user may specify zero, one, two, or three arguments.

The arguments that the user does not specify will be filled in according to the arguments' type,

If the argument type is ...	The default value will be ...
undeclared	J(0, 0, .)
transmorphic matrix	J(0, 0, .)
real matrix	J(0, 0, .)
complex matrix	J(0, 0, 1i)
string matrix	J(0, 0, "")
pointer matrix	J(0, 0, NULL)
transmorphic rowvector	J(1, 0, .)
real rowvector	J(1, 0, .)
complex rowvector	J(1, 0, 1i)
string rowvector	J(1, 0, "")
pointer rowvector	J(1, 0, NULL)
transmorphic colvector	J(0, 1, .)
real colvector	J(0, 1, .)
complex colvector	J(0, 1, 1i)
string colvector	J(0, 1, "")
pointer colvector	J(0, 1, NULL)
transmorphic vector	J(1, 0, .)
real vector	J(1, 0, .)
complex vector	J(1, 0, 1i)
string vector	J(1, 0, "")
pointer vector	J(1, 0, NULL)
transmorphic scalar	J(1, 1, .)
real scalar	J(1, 1, .)
complex scalar	J(1, 1, C(.))
string scalar	J(1, 1, "")
pointer scalar	J(1, 1, NULL)

Also, the function `args()` (see [M-5] `args()`) will return the number of arguments that the user specified.

The vertical bar can be specified only once. That is sufficient, as we will show.

## Examples revisited

### ► Example 1

In this example, real matrix function `ditty(A, B, scale)` allowed real scalar *scale* to be optional. If *scale* was not specified, results were as if *scale*=. had been specified. This can be coded

```
real matrix ditty(real matrix A, real matrix B, |real scalar scale)
{
    ...
}
```

The body of the code is written just as if *scale* were not optional because, if the caller does not specify the argument, the missing argument is automatically filled in with missing, per the table above. ◀

### ► Example 2

Real matrix function `gash(A, B)` allowed real matrix *B* to be optional, and if not specified, *B* = `J(0,0,.)` was assumed. Hence, this is coded just as example 1 was coded:

```
real matrix gash(real matrix A, |real matrix B)
{
    ...
}
```

◀

### ► Example 3

Real scalar function `easygoing(A, B, scale)` allowed all arguments to be optional. *scale* = 1 was assumed, *B* = *A*, and if necessary, *A* = `I(2)`.

```
real scalar easygoing(|real matrix A, real matrix B,
                      real scalar scale)
{
    ...
    if (args()==2) scale = 1
    else if (args()==1) {
        B = A
        scale = 1
    }
    else if (args()==0) {
        A = B = I(2)
        scale = 1 ;
    }
    ...
}
```

◀

### ► Example 4

Real matrix function `midsection(A, w, B)` allowed  $w$ —its middle argument—to be omitted. If  $w$  was not specified, `J(1, cols(A), 1)` was assumed. Here is one solution:

```
real matrix midsection(a1, a2, |a3)
{
    if (args()==3) return(midsection_u(a1,      a2,      a3))
    else          return(midsection_u(a1, J(1,cols(a1),1), a2))
}
real matrix midsection_u(real matrix A, real vector w, real matrix B)
{
    ...
}
```

We will never tell callers about the existence of `midsection_u()` even though `midsection_u()` is our real program.

What we did above was write `midsection()` to take two or three arguments, and then we called `midsection_u()` with the arguments in the correct position.



## Also see

[\[M-2\] intro](#) — Language definition

## Syntax

`pointer` $[ (totype) ] [ orgtype ] [ function ] \dots$

where *totype* is

$[ eltype ] [ orgtype ] [ function ]$

and where *eltype* and *orgtype* are

<i>eltype</i>	<i>orgtype</i>
transmorphic	matrix
numeric	vector
real	rowvector
complex	colvector
string	scalar
<code>pointer</code> $[ (towhat) ]$	

`pointer` $[ (totype) ] [ orgtype ]$  can be used in front of declarations, be they function declarations, argument declarations, or variable definitions.

## Description

Pointers are objects that contain the addresses of other objects. The `*` prefix operator obtains the contents of an address. Thus if *p* is a pointer, `*p` refers to the contents of the object to which *p* points. Pointers are an advanced programming concept. Most programs, including involved and complicated ones, can be written without them.

In Mata, pointers are commonly used to

1. put a collection of objects under one name and
2. pass functions to functions.

One need not understand everything about pointers merely to pass functions to functions; see [\[M-2\] `ftof`](#).

## Remarks

Remarks are presented under the following headings:

- What is a pointer?*
- Pointers to variables*
- Pointers to expressions*
- Pointers to functions*
- Pointers to pointers*
- Pointer arrays*
- Mixed pointer arrays*
- Definition of NULL*
- Use of parentheses*
- Pointer arithmetic*
- Listing pointers*
- Declaration of pointers*
- Use of pointers to collect objects*
- Efficiency*

## What is a pointer?

A pointer is the address of a variable or a function. Say that variable  $X$  contains a matrix. Another variable  $p$  might contain 137,799,016, and if 137,799,016 were the address at which  $X$  were stored, then  $p$  would be said to point to  $X$ . Addresses are seldom written in base 10, so rather than saying  $p$  contains 137,799,016, we would be more likely to say that  $p$  contains 0x836a568, which is the way we write numbers in base 16. Regardless of how we write addresses, however,  $p$  contains a number and that number corresponds to the address of another variable.

In our program, if we refer to  $p$ , we are referring to  $p$ 's contents, the number 0x836a568. The monadic operator  $*$  is defined as “refer to the contents of the address” or “dereference”:  $*p$  means  $X$ . We could code  $Y = *p$  or  $Y = X$ , and either way, we would obtain the same result. In our program, we could refer to  $X[i, j]$  or  $(*p)[i, j]$ , and either way, we would obtain the  $i, j$  element of  $X$ .

The monadic operator  $\&$  is how we put addresses into  $p$ . To load  $p$  with the address of  $X$ , we code  $p = \&X$ .

The special address 0 (zero, written in hexadecimal as 0x0), also known as NULL, is how we record that a pointer variable points to nothing. A pointer variable contains NULL or it contains the address of another variable.

Or it contains the address of a function. Say that  $p$  contains 0x836a568 and that 0x836a568, rather than being the address of matrix  $X$ , is the address of function  $f()$ . To get the address of  $f()$  into  $p$ , just as we coded  $p = \&X$  previously, we code  $p = \&f()$ . The  $()$  at the end tells  $\&$  that we want the address of a function. We code  $()$  on the end regardless of the number of arguments  $f()$  requires because we are not executing  $f()$ , we are just obtaining its address.

To execute the function at 0x836a568—now we will assume that  $f()$  takes two arguments and call them  $i$  and  $j$ —we code  $(*p)(i, j)$  just as we coded  $(*p)[i, j]$  when  $p$  contained the address of matrix  $X$ .



## Pointers to variables

To create a pointer  $p$  to a variable, you code

```
 $p = \&vname$ 
```

For instance, if  $X$  is a matrix,

```
 $p = \&X$ 
```

stores in  $p$  the address of  $X$ . Subsequently, referring to  $*p$  and referring to  $X$  amount to the same thing. That is, if  $X$  contained a  $3 \times 3$  identity matrix and you coded

```
 $*p = \text{Hilbert}(4)$ 
```

then after that you would find that  $X$  contained the  $4 \times 4$  Hilbert matrix.  $X$  and  $*p$  are the same matrix.

If  $X$  contained a  $3 \times 3$  identity matrix and you coded

```
 $(*p)[2,3] = 4$ 
```

you would then find  $X[2,3]$  equal to 4.

You cannot, however, point to the interior of objects. That is, you cannot code

```
 $p = \&X[2,3]$ 
```

and get a pointer that is equivalent to  $X[2,3]$  in the sense that if you later coded  $*p=2$ , you would see the change reflected in  $X[2,3]$ . The statement  $p = \&X[2,3]$  is valid, but what it does, we will explain in [Pointers to expressions](#) below.

By the way, variables can be sustained by being pointed to. Consider the program

```
pointer(real matrix) scalar example(real scalar n)
{
    real matrix    tmp
    tmp = I(3)
    return(&tmp)
}
```

Ordinarily, variable `tmp` would be destroyed when `example()` concluded execution. Here, however, `tmp`'s existence will be sustained because of the pointer to it. We might code

```
 $p = \text{example}(3)$ 
```

and the result will be to create  $*p$  containing the  $3 \times 3$  identity matrix. The memory consumed by that matrix will be freed when it is no longer being pointed to, which will occur when  $p$  itself is freed, or, before that, when the value of  $p$  is changed, perhaps by

```
 $p = \text{NULL}$ 
```

For a discussion of pointers to structures, see [\[M-2\] struct](#). For a discussion of pointers to classes, see [\[M-2\] class](#).

## Pointers to expressions

You can code

```
p = &(2+3)
```

and the result will be to create *\*p* containing 5. Mata creates a temporary variable to contain the evaluation of the expression and sets *p* to the address of the temporary variable. That temporary variable will be freed when *p* is freed or, before that, when the value of *p* is changed, just as *tmp* was freed in the example in the previous section.

When you code

```
p = &X[2,3]
```

the result is the same. The expression is evaluated and the result of the expression stored in a temporary variable. That is why subsequently coding *\*p=2* does not change *X[2,3]*. All *\*p=2* does is change the value of the temporary variable.

Setting pointers equal to the value of expressions can be useful. In the following code fragment, we create  $n \times 5 \times 5$  matrices for later use:

```
pvec = J(1, n, NULL)
for (i=1; i<=n; i++) pvec[i] = &(J(5, 5, .))
```

## Pointers to functions

When you code

```
p = &functionname()
```

the address of the function is stored in *p*. You can later execute the function by coding

```
... (*p)(...)
```

Distinguish carefully between

```
p = &functionname()
```

and

```
p = &(functionname())
```

The latter would execute *functionname()* with no arguments and then assign the returned result to a temporary variable.

For instance, assume that you wish to write a function *neat()* that will calculate the derivative of another function, which function you will pass to *neat()*. Your function, we will pretend, returns a real scalar.

You could do that as follows

```
real scalar neat(pointer(function) p, other args...)
{
    ...
}
```

although you could be more explicit as to the characteristics of the function you are passed:

```
real scalar neat(pointer(real scalar function) p, other args...)
{
    ...
}
```

In any case, inside the body of your function, where you want to call the passed function, you code

```
(*p)(arguments)
```

For instance, you might code

```
approx = ( (*p)(x+delta)-(*p)(x) ) / delta
```

The caller of your `neat()` function, wanting to use it with, say, function `zeta_i_just_wrote()`, would code

```
result = neat(&zeta_i_just_wrote(), other args...)
```

For an example of the use of pointers in sophisticated numerical calculations, see [Størving \(2007\)](#).

## Pointers to pointers

Pointers to pointers (to pointers ...) are allowed, for instance, if  $X$  is a matrix,

```
p1 = &X
p2 = &p1
```

Here  $*p2$  is equivalent to  $p1$ , and  $**p2$  is equivalent to  $X$ .

Similarly, we can construct a pointer to a pointer to a function:

```
q1 = &f()
q2 = &p1
```

Here  $*q2$  is equivalent to  $q1$ , and  $**q2$  is equivalent to  $f()$ .

When constructing pointers to pointers, never type  $\&\&$ —such as  $\&\&x$ —to obtain the address of the address of  $x$ . Type  $\&(\&x)$  or  $\&\&x$ .  $\&\&$  is a synonym for  $\&$ , included for those used to coding in C.

## Pointer arrays

You may create an array of pointers, such as

```
P = (&X1, &X2, &X3)
```

or

```
Q = (&f1(), &f2(), &f3())
```

Here  $*P[2]$  is equivalent to  $X2$  and  $*Q[2]$  is equivalent to  $f2()$ .

## Mixed pointer arrays

You may create mixed pointer arrays, such as

$$R = (\&X, \&f())$$

Here  $\ast R[2]$  is equivalent to  $f()$ .

You may not, however, create arrays of pointers mixed with real, complex, or string elements. Mata will abort with a type-mismatch error.

## Definition of NULL

NULL is the special pointer value that means “points to nothing” or undefined. NULL is like 0 in many ways—for instance, coding `if (X)` is equivalent to coding `if (X!=NULL)`, but NULL is distinct from zero in other ways. 0 is a numeric value; NULL is a pointer value.

## Use of parentheses

Use parentheses to make your meaning clear.

In the table below, we assume

$$\begin{aligned} p &= \&X \\ P &= (\&X11, \&X12 \setminus \&X21, \&X22) \\ q &= \&f() \\ Q &= (\&f11(), \&f12() \setminus \&f21(), \&f22()) \end{aligned}$$

where  $X, X11, X12, X21$ , and  $X22$  are matrices and  $f(), f11(), f12(), f21()$ , and  $f22()$  are functions.

Expression	Meaning
$\ast p$	$X$
$\ast p[1,1]$	$X$
$(\ast p)[1,1]$	$X[1,1]$
$\ast P[1,2]$	$X12$
$(\ast P[1,2])[3,4]$	$X12[3,4]$
$\ast q(a,b)$	execute function $q()$ of $a, b$ ; dereference that
$(\ast q)(a,b)$	$f(a,b)$
$(\ast q[1,1])(a,b)$	$f(a,b)$
$\ast Q[1,2](a,b)$	nonsense
$(\ast Q[1,2])(a,b)$	$f12(a,b)$

## Pointer arithmetic

Arithmetic with pointers (which is to say, with addresses) is not allowed:

```
: y = 2
: x = &y
: x+2
<stmt>: 3205 undefined operation on pointer
```

Do not confuse the expression `x+2` with the expression `*x+2`, which is allowed and in fact evaluates to 4.

You may use the equality and inequality operators `==` and `!=` with pairs of pointer values:

```
if (p1 == p2) {
    ...
}
if (p1 != p2) {
    ...
}
```

Also pointer values may be assigned and compared with the value `NULL`, which is much like, but still different from, zero: `NULL` is a  $1 \times 1$  scalar containing an address value of 0. An unassigned pointer has the value `NULL`, and you may assign the value `NULL` to pointers:

```
p = NULL
```

Pointer values may be compared with `NULL`,

```
if (p1 == NULL) {
    ...
}
if (p1 != NULL) {
    ...
}
```

but if you attempt to dereference a `NULL` pointer, you will get an error:

```
: x = NULL
: *x + 2
<stmt>: 3120 attempt to dereference NULL pointer
```

Concerning logical expressions, you may directly examine pointer values:

```
if (p1) {
    ...
}
```

The above is interpreted as if `if (p1!=NULL)` were coded.

## Listing pointers

You may list pointers:

```
: y = 2
: x = &y
: x
0x8359e80
```

What is shown, 0x8359e80, is the memory address of `y` during our Stata session. If you typed the above lines, the address you would see could differ, but that does not matter.

Listing the value of pointers often helps in debugging because, by comparing addresses, you can determine where pointers are pointing and whether some are pointing to the same thing.

In listings, NULL is presented as 0x0.

## Declaration of pointers

Declaration of pointers, as with all declarations (see [M-2] [declarations](#)) is optional. That basic syntax is

```
pointer[ (totype) ] orgtype [ function ] ...
```

For instance,

```
pointer(real matrix) scalar p1
```

declares that *p1* is a pointer scalar and that it points to a real matrix, and

```
pointer(complex colvector) rowvector p2
```

declares that *p2* is a rowvector of pointers and that each pointer points to a complex colvector, and

```
pointer(real scalar function) scalar p3
```

declares that *p3* is a pointer scalar, and that it points to a function that returns a real scalar, and

```
pointer(pointer(real matrix function) rowvector) colvector p4
```

declares that *p4* is a column vector of pointers to pointers, the pointers to which each element points are rowvectors, and each of those elements points to a function returning a real matrix.

You can omit the pieces you wish.

```
pointer() scalar p5
```

declares that *p5* is a pointer scalar—to what being uncertain.

```
pointer scalar p5
```

means the same thing.

```
pointer p6
```

declares that *p6* is a pointer, but whether it is a matrix, vector, or scalar, is unsaid.

## Use of pointers to collect objects

Assume that you wish to write a function in two parts: `result_setup()` and `repeated_result()`.

In the first part, `result_setup()`, you will be passed a matrix and a function by the user, and you will make a private calculation that you will use later, each time `repeated_result()` is called. When `repeated_result()` is called, you will need to know the matrix, the function, and the value of the private calculation that you previously made.

One solution is to adopt the following design. You request the user code

```
resultinfo = result_setup(setup args...)
```

on the first call, and

```
value = repeated_result(resultinfo, other args...)
```

on subsequent calls. The design is that you will pass the information between the two functions in `resultinfo`. Here `resultinfo` will need to contain three things: the original matrix, the original function, and the private result you calculated. The user, however, never need know the details, and you will simply request that the user declare `resultinfo` as a pointer vector.

Filling in the details, your code

```
pointer vector result_setup(real matrix X, pointer(function) f)
{
    real matrix      privmat
    pointer vector   info

    ...
    privmat = ...
    ...
    info = (&X, f, &privmat)
    return(info)
}

real matrix repeated_result(pointer vector info, ...)
{
    pointer(function) scalar  f
    pointer(matrix)   scalar  X
    pointer(matrix)   scalar  privmat

    f = info[2]
    X = info[1]
    privmat = info[3]

    ...
    ... (*f)(...) ...
    ... (*X) ...
    ... (*privmat) ...
    ...
}
```

It was not necessary to unload `info[]` into the individual scalars. The lines using the passed values could just as well have read

```
... (*info[2])(...) ...
... (*info[1]) ...
... (*info[3]) ...
```

## Efficiency

When calling subroutines, it is better to pass the evaluation of pointer scalar arguments rather than the pointer scalar itself, because then the subroutine can run a little faster. Say that `p` points to a real matrix. It is better to code

```
... mysub(*p) ...
```

rather than

```
... mysub(p) ...
```

and then to write `mysub()` as

```
function mysub(real matrix X)
{
    ... X ...
}
```

rather than

```
function mysub(pointer(real matrix) scalar p)
{
    ... (*p) ...
}
```

Dereferencing a pointer (obtaining `*p` from `p`) does not take long, but it does take time. Passing `*p` rather than `p` can be important if `mysub()` loops and performs the evaluation of `*p` hundreds of thousands or millions of times.

## Diagnostics

The prefix operator `*` (called the dereferencing operator) aborts with error if it is applied to a nonpointer object.

Arithmetic may not be performed on undereferenced pointers. Arithmetic operators abort with error.

The prefix operator `&` aborts with error if it is applied to a built-in function.

## Reference

Støvring, H. 2007. [A generic function evaluator implemented in Mata](#). *Stata Journal* 7: 542–555.



## Also see

[M-2] [intro](#) — Language definition

# Title

[M-2] **pragma** — Suppressing warning messages

## Syntax

```
pragma unset varname
pragma unused varname
```

## Description

`pragma` informs the compiler of your intentions so that the compiler can avoid presenting misleading warning messages and so that the compiler can better optimize the code.

## Remarks

Remarks are presented under the following headings:

*pragma unset*  
*pragma unused*

### **pragma unset**

The `pragma`

```
pragma unset X
```

suppresses the warning message

**note:** variable *X* may be used before set

The `pragma` has no effect on the resulting compiled code.

In general, the warning message flags logical errors in your program, such as

```
: function problem(real matrix a, real scalar j)
> {
>     real scalar i
>
>     j = i
>     ...
> }
note: variable i may be used before set
```

Sometimes, however, the message is misleading:

```
: function notaproblem(real matrix a, real scalar j)
> {
>     real matrix V
>
>     st_view(V, ...)
>     ...
> }
note: variable V may be used before set
```

In the above, function `st_view()` (see [M-5] `st_view()`) defines `V`, but the compiler does not know that.

The warning message causes no problem but, if you wish to suppress it, change the code to read

```
: function notaproblem(real matrix a, real scalar j)
> {
>   real matrix V
>
>   pragma unset V
>   st_view(V, ...)
>   ...
> }
```

`pragma unset V` states that you know `V` is unset and that, for warning messages, the compiler should act as if `V` were set at this point in your code.

## pragma unused

The pragma

```
pragma unused X
```

suppresses the warning messages

```
note: argument X unused
note: variable X unused
note: variable X set but not used
```

The pragma has no effect on the resulting compiled code.

Intentionally unused variables most often arise with respect to function arguments. Your code

```
: function resolve(A, B, C)
> {
>   ...
> }
note: argument C unused
```

and you know well that you are not using `C`. You include the unnecessary argument because you are attempting to fit into a standard or you know that, later, you may wish to change the function to include `C`. To suppress the warning message, change the code to read

```
: function resolve(A, B, C)
> {
>   ...
>   pragma unused C
>   ...
> }
```

The pragma states that you know `C` is unused and, for the purposes of warning messages, the compiler should act as if `C` were used at this point in your code.

Unused variables can also arise, and in general, they should simply be removed,

```
: function resin(X, Y)
> {
>   real scalar i
>   ...
>   ... code in which i never appears
>   ...
> }
note: variable i unused
```

Rather than using the pragma to suppress the message, you should remove the line `real scalar i`.

Warnings are also given for variables that are set and not used:

```
: function thwart(X, Y)
> {
>   real scalar i
>   ...
>   i = 1
>   ...
>   ... code in which i never appears
>   ...
> }
note: variable i set but unused
```

Here you should remove both the `real scalar i` and `i = 1` lines.

It is possible, however, that the set-but-unused variable was intentional:

```
: function thwart(X, Y)
> {
>   real scalar i
>   ...
>   i = somefunction(...)
>   ...
>   ... code in which i never appears
>   ...
> }
note: variable i set but not used
```

You assigned the value of `somefunction()` to `i` to prevent the result from being displayed. Here you could use `pragma unused i` to suppress the warning message, but a better alternative would be

```
: function thwart(X, Y)
> {
>   ...
>   (void) somefunction(...)
>   ...
> }
```

See [Assignment suppresses display, as does \(void\) in \[M-2\] exp](#).

## Also see

[\[M-2\] intro](#) — Language definition

## Syntax

Reserved words are

aggregate	float	pointer	union
array	for	polymorphic	unsigned
	friend	pragma	using
boolean	function	private	
break		protected	vector
byte	global	public	version
	goto		virtual
case		quad	void
catch	if		volatile
class	inline	real	
colvector	int	return	while
complex		rowvector	
const	local		
continue	long	scalar	
		short	
default	mata	signed	
delegate	matrix	static	
delete		string	
do	namespace	struct	
double	new	super	
	NULL	switch	
else	numeric		
eltypedef		template	
end	operator	this	
enum	orgtypedef	throw	
explicit		transmorphic	
export		try	
external		typedef	
		typename	

## Description

Reserved words are words reserved by the Mata compiler; they may not be used to name either variables or functions.

## Remarks

Remarks are presented under the following headings:

*Future developments*  
*Version control*

## Future developments

Many of the words above are reserved for the future implementation of new features. For instance, the words `aggregate`, `array`, `boolean`, `byte`, etc., currently play no role in Mata, but they are reserved.

You cannot infer much about short-run development plans from the presence or absence of a word from the list. The list was constructed by including words that would be needed to add certain features, but we have erred on the side of reserving too many words because it is better to give back a word than to take it away later. Taking away a word can cause previously written code to break.

Also, features can be added without reserving words if the word will be used only within a specific context. Our original list was much longer, but then we struck from it such context-specific words.

## Version control

Even if we should need to reserve new words in the future, you can ensure that you need not modify your programs by engaging in version control; see [\[M-2\] \*\*version\*\*](#).

## Also see

[\[M-1\] \*\*naming\*\*](#) — Advice on naming functions and variables

[\[M-2\] \*\*version\*\*](#) — Version control

[\[M-2\] \*\*intro\*\*](#) — Language definition

# Title

[M-2] **return** — return and return(*exp*)

## Syntax

```
return
```

```
return(exp)
```

## Description

`return` causes the function to stop execution and return to the caller, returning nothing.

`return(exp)` causes the function to stop execution and return to the caller, returning the evaluation of *exp*.

## Remarks

Remarks are presented under the following headings:

*Functions that return results*

*Functions that return nothing (void functions)*

### Functions that return results

`return(exp)` specifies the value to be returned. For instance, you have written a program to return the sum of two numbers:

```
function mysum(a, b)
{
    return(a+b)
}
```

`return(exp)` may appear multiple times in the program. The following program calculates *x* factorial; it assumes *x* is an integer greater than 0:

```
real scalar myfactorial(real scalar x)
{
    if (x<=0) return(1)
    return(x*factorial(x-1))
}
```

If  $x \leq 0$ , the function returns 1; execution does not continue to the next line.

Functions that return a result always include one or more `return(exp)` statements.

## Functions that return nothing (void functions)

A function is said to be void if it returns nothing. The following program changes the diagonal of a matrix to be 1:

```
function fixdiag(matrix A)
{
    real scalar    i

    for (i=1; i<=rows(A); i++) A[i,i] = 1
}
```

This function does not even include a `return` statement; execution just ends. That is fine, although the function could just as well read

```
function fixdiag(matrix A)
{
    real scalar    i

    for (i=1; i<=rows(A); i++) A[i,i] = 1
    return
}
```

The use of `return` is when the function has reason to end early:

```
void fixmatrix(matrix A, scalar how)
{
    real scalar    i, j

    for (i=1; i<=rows(A); i++) A[i,i] = 1
    if (how==0) return
    for (i=1; i<=rows(A); i++) {
        for (j=1; j<i; j++) A[i,j] = 0
    }
}
```

## Also see

[M-5] [exit\(\)](#) — Terminate execution

[M-2] [intro](#) — Language definition



# Title

## [M-2] semicolons — Use of semicolons

## Syntax

*stmt*

*stmt ;*

## Description

Mata allows, but does not require, semicolons.

Use of semicolons is discussed below, along with advice on the possible interactions of Stata's `#delimit` instruction; see [P] [#delimit](#).

## Remarks

Remarks are presented under the following headings:

*Optional use of semicolons*

*You cannot break a statement anywhere even if you use semicolons*

*Use of semicolons to create multistatement lines*

*Significant semicolons*

*Do not use #delimit ;*

## Optional use of semicolons

You can code your program to look like this

```
real scalar foo(real matrix A)
{
    real scalar    i, sum

    sum = 0
    for (i=1; i<=rows(A); i++) {
        sum = sum + A[i,i]
    }
    return(sum)
}
```

or you can code your program to look like this:

```
real scalar foo(real matrix A)
{
    real scalar    i, sum ;

    sum = 0 ;
    for (i=1; i<=rows(A); i++) {
        sum = sum + A[i,i] ;
    }
    return(sum) ;
}
```

That is, you may omit or include semicolons at the end of statements. It makes no difference. You can even mix the two styles:

```
real scalar foo(real matrix A)
{
    real scalar    i, sum ;

    sum = 0 ;
    for (i=1; i<=rows(A); i++) {
        sum = sum + A[i,i]
    }
    return(sum)
}
```

## You cannot break a statement anywhere even if you use semicolons

Most languages that use semicolons follow the rule that a statement continues up to the semicolon.

Mata follows a different rule: a statement continues across lines until it looks to be complete, and semicolons force the end of statements.

For instance, consider the statement  $x=b-c$  appearing in some program. In the code, might appear

```
x = b -
c
```

or

```
x = b -
c ;
```

and, either way, Mata will understand the statement to be  $x=b-c$ , because the statement could not possibly end at the minus:  $x=b-$  makes no sense.

On the other hand,

```
x = b
- c
```

would be interpreted by Mata as two statements:  $x=b$  and  $-c$  because  $x = b$  looks like a completed statement to Mata. The first statement will assign  $b$  to  $x$ , and the second statement will display the negative value of  $c$ .

Adding a semicolon will not help:

```
x = b
- c ;
```

$x = b$  is still, by itself, a complete statement. All that has changed is that the second statement ends in a semicolon, and that does not matter.

Thus remember always to break multiline statements at places where the statement could not possibly be interpreted as being complete, such as

```
x = b -
    c + (d
        + e)
myfunction(A,
           B, C,
           )
```

Do this whether or not you use semicolons.

## Use of semicolons to create multistatement lines

Semicolons allow you to put more than one statement on a line. Rather than coding

```
a = 2
b = 3
```

you can code

```
a = 2 ; b = 3 ;
```

and you can even omit the trailing semicolon:

```
a = 2 ; b = 3
```

Whether you code separate statements on separate lines or the same line is just a matter of style; it does not change the meaning. Coding

```
for (i=1; i<n; i++) a[i] = -a[i] ; sum = sum + a[i] ;
```

still means

```
for (i=1; i<n; i++) a[i] = -a[i] ;
sum = sum + a[i] ;
```

and, without doubt, the programmer intended to code

```
for (i=1; i<n; i++) {
    a[i] = -a[i] ;
    sum = sum + a[i] ;
}
```

which has a different meaning.

## Significant semicolons

Semicolons are not all style. The syntax for the `for` statement is (see [\[M-2\] for](#))

```
for (exp1; exp2; exp3) stmt
```

Say that the complete `for` loop that we want to code is

```
for (x=init(); !converged(x); iterate(x))
```

and that there is no *stmt* following it. Then we must code

```
for (x=init(); !converged(x); iterate(x)) ;
```

Here we use the semicolon to force the end of the statement. Say we omitted it, and the code read

```
...
for (x=init(); !converged(x); iterate(x))
x = -x
...
```

The `for` statement would look incomplete to Mata, so it would interpret our code as if we had coded

```
for (x=init(); !converged(x); iterate(x)) {
    x = -x
}
```

Here the semicolon is significant.

Significant semicolons only happen following `for` and `while`.

## Do not use `#delimit ;`

What follows has to do with Stata's `#delimit ;` mode. If you do not know what it is or if you never use it, you can skip what follows.

Stata has an optional ability to allow its lines to continue up to semicolons. In Stata, you code

```
. #delimit ;
```

and the delimiter is changed to semicolon until your do-file or ado-file ends, or until you code

```
. #delimit cr
```

We recommend that you do not use Mata when Stata is in `#delimit ;` mode. Mata will not mind, but you will confuse yourself.

When Mata gets control, if `#delimit ;` is on, Mata turns it off temporarily, and then Mata applies its own rules, which we have summarized above.

## Also see

[\[M-2\] intro](#) — Language definition

[\[P\] #delimit](#) — Change delimiter

## Syntax

```
struct structname {  
    declaration(s)  
}
```

such as

```
struct mystruct {  
    real scalar    n1, n2  
    real matrix    X  
}
```

## Description

A structure contains a set of variables tied together under one name.

Structures may not be used interactively. They are used in programming.

## Remarks

Remarks are presented under the following headings:

*Introduction*

*Structures and functions must have different names*

*Structure variables must be explicitly declared*

*Declare structure variables to be scalars whenever possible*

*Vectors and matrices of structures*

*Structures of structures*

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*Operators and functions for use with entire structures*

*Listing structures*

*Use of transmorphics as passthru*

*Saving compiled structure definitions*

*Saving structure variables*

## Introduction

Here is an overview of the use of structures:

```
struct mystruct {
    real scalar    n1, n2
    real matrix    X
}

function myfunc()
{
    struct mystruct scalar e

    ...
    e.n1 = ...
    e.n2 = ...
    e.X  = ...
    ...
    ... mysubroutine(e, ...)
    ...
}

function mysubroutine(struct mystruct scalar x, ...)
{
    struct mystruct scalar    y
    ...
    ... x.n1 ... x.n2 ... x.X ...
    ...
    y = mysubfcn(x)
    ...
    ... y.n1 ... y.n2 ... y.X ...
    ... x.n1 ... x.n2 ... x.X ...
    ...
}

struct mystruct scalar mysubfcn(struct mystruct scalar x)
{
    struct mystruct scalar    result

    result = x
    ... result.n1 ... result.n2 ... result.X ...
    return(result)
}
```

Note the following:

1. We first defined the structure. Definition does not create variables; definition defines what we mean when we refer to a `struct mystruct` in the future. This definition is done outside and separately from the definition of the functions that will use it. The structure is defined before the functions.
2. In `myfunc()`, we declared that variable `e` is a `struct mystruct scalar`. We then used variables `e.n1`, `e.n2`, and `e.X` just as we would use any other variable. In the call to `mysubroutine()`, however, we passed the entire `e` structure.

3. In `mysubroutine()`, we declared that the first argument received is a `struct mystruct` scalar. We chose to call it `x` rather than `e` to emphasize that names are not important. `y` is also a `struct mystruct` scalar.
4. `mysubfcn()` not only accepts a `struct mystruct` as an argument but also returns a `struct mystruct`. One of the best uses of structures is as a way to return multiple, related values.

The line `result=x` copied all the values in the structure; we did not need to code `result.n1=x.n1`, `result.n2=x.n2`, and `result.X=x.X`.

## Structures and functions must have different names

You define structures much as you define functions, at the colon prompt, with the definition enclosed in braces:

```
: struct twopart {
>     real scalar    n1, n2
> }
: function setuphistory()
> {
>     ...
> }
```

Structures and functions may not have the same names. If you call a structure `twopart`, then you cannot have a function named `twopart()`, and vice versa.

## Structure variables must be explicitly declared

Declarations are usually optional in Mata. You can code

```
real matrix swaprows(real matrix A, real scalar i1, real scalar i2)
{
    real matrix    B
    real rowvector  v

    B = A
    v = B[i1, .]
    B[i1, .] = B[i2, .]
    B[i2, .] = v
    return(B)
}
```

or you can code

```
function swaprows(A, i1, i2)
{
    B = A
    v = B[i1, .]
    B[i1, .] = B[i2, .]
    B[i2, .] = v
    return(B)
}
```

When a variable, argument, or returned value is a structure, however, you must explicitly declare it:

```
function makecalc()
{
    struct twopart scalar    t
    t.n1 = t.n2 = 0
    ...
}

function clear_twopart(struct twopart scalar t)
{
    t.n1 = t.n2 = 0
}

struct twopart scalar new_twopart()
{
    struct twopart scalar    t
    t.n1 = t.n2 = 0
    return(t)
}
```

In the functions above, we refer to variables `t.n1` and `t.n2`. The Mata compiler cannot interpret those names unless it knows the definition of the structure.

Aside: all structure references are resolved at compile time, not run time. That is, terms like `t.n1` are not stored in the compiled code and resolved during execution. Instead, the Mata compiler accesses the structure definition when it compiles your code. The compiler knows that `t.n1` refers to the first element of the structure and generates efficient code to access it.

## Declare structure variables to be scalars whenever possible

In our declarations, we code things like

```
struct twopart scalar    t
```

and do not simply code

```
struct twopart           t
```

although the simpler statement would be valid.

Structure variables can be scalars, vectors, or matrices; when you do not say which, matrix is assumed.

Most uses of structures are as scalars, and the compiler will generate more efficient code if you tell it that the structures are scalars. Also, when you use structure vectors or matrices, there is an extra step you need to fill in, as described in the next section.

## Vectors and matrices of structures

Just as you can have real scalars, vectors, or matrices, you can have structure scalars, vectors, or matrices. The following are all valid:



```

struct twopart scalar    t
struct twopart vector    t
struct twopart rowvector t
struct twopart colvector t
struct twopart matrix    t

```

In a `struct twopart matrix`, every element of the matrix is a separate structure. Say that the matrix were  $2 \times 3$ . Then you could refer to any of the following variables,

```

t[1,1].n1
t[1,2].n1
t[1,3].n1
t[2,1].n1
t[2,2].n1
t[2,3].n1

```

and similarly for `t[i,j].n2`.

If `struct twopart` also contained a matrix `X`, then

```
t[i,j].X
```

would refer to the  $(i,j)$ th matrix.

```
t[i,j].X[k,l]
```

would refer to the  $(k,l)$ th element of the  $(i,j)$ th matrix.

If `t` is to be a  $2 \times 3$  matrix, you must arrange to make it  $2 \times 3$ . After the declaration

```
struct twopart matrix t
```

`t` is  $0 \times 0$ . This result is no different from the situation where `t` is a `real matrix` and after declaration, `t` is  $0 \times 0$ .

Whether `t` is a `real matrix` or a `struct twopart matrix`, you allocate `t` by assignment. Let's pretend that `t` is a `real matrix`. There are three solutions to the allocation problem:

- (1) `t = x`
- (2) `t = somefunction(...)`
- (3) `t = J(r, c, v)`

All three are so natural that you do not even think of them as allocation; you think of them as definition.

The situation with structures is the same.

Let's take each in turn.

1. `x` contains a  $2 \times 3$  `struct twopart`. You code

```
t = x
```

and now `t` contains a copy of `x`. `t` is  $2 \times 3$ .

2. `somefunction(...)` returns a  $2 \times 3$  struct `twopart`. You code

```
t = somefunction(...)
```

and now `t` contains the  $2 \times 3$  result.

3. Mata function `J(r, c, v)` returns an  $r \times c$  matrix, every element of which is set to *v*. So pretend that variable `tpc` contains a struct `twopart` scalar. You code

```
t = J(2, 3, tpc)
```

and now `t` is  $2 \times 3$ , every element of which is a copy of `tpc`. Here is how you might do that:

```
function ...(...)
{
    struct twopart scalar    tpc
    struct twopart matrix    t

    ...
    t = J(2, 3, tpc)
    ...
}
```

Finally, there is a fourth way to create structure vectors and matrices. When you defined

```
struct twopart {
    real scalar    n1, n2
}
```

Mata not only recorded the structure definition but also created a function named `twopart()` that returns a struct `twopart`. Thus, rather than enduring all the rigmarole of creating a matrix from a preallocated scalar, you could simply code

```
t = J(2, 3, twopart())
```

In fact, the function `twopart()` that Mata creates for you allows 0, 1, or 2 arguments:

<code>twopart()</code>	returns a $1 \times 1$ struct <code>twopart</code>
<code>twopart(<i>r</i>)</code>	returns an $r \times 1$ struct <code>twopart</code>
<code>twopart(<i>r</i>, <i>c</i>)</code>	returns an $r \times c$ struct <code>twopart</code>

so you could code

```
t = twopart(2, 3)
```

or you could code

```
t = J(2, 3, twopart())
```

and whichever you code makes no difference.

Either way, what is in `t`? Each element contains a separate struct `twopart`. In each struct `twopart`, the scalars have been set to missing (`.`, `""`, or `NULL`, as appropriate), the vectors and row vectors have been made  $1 \times 0$ , the column vectors  $0 \times 1$ , and the matrices  $0 \times 0$ .

## Structures of structures

Structures may contain other structures:

```
struct twopart {
    real scalar    n1, n2
}

struct pair_of_twoparts {
    struct twopart scalar  t1, t2
}
```

If `t` were a `struct pair_of_twoparts` scalar, then the members of `t` would be

```
t.t1      a struct twopart scalar
t.t2      a struct twopart scalar
t.t1.n1   a real scalar
t.t1.n2   a real scalar
t.t2.n1   a real scalar
t.t2.n2   a real scalar
```

You may also create structures of structures of structures, structures of structures of structures of structures, and so on. You may not, however, include a structure in itself:

```
struct recursive {
    ...
    struct recursive scalar  r
    ...
}
```

Do you see how, even in the scalar case, `struct recursive` would require an infinite amount of memory to store?

## Pointers to structures

What you can do is this:

```
struct recursive {
    ...
    pointer(struct recursive scalar) scalar  r
    ...
}
```

Thus, if `r` were a `struct recursive` scalar, then `*r.r` would be the next structure, or `r.r` would be `NULL` if there were no next structure. Immediately after allocation, `r.r` would equal `NULL`.

In this way, you may create linked lists.

Mata provides operator `->` for accessing members of pointers to structures.

Let `rec` be a `struct recursive`, and assume that `struct recursive` also had member `real scalar n`, so that `rec.n` would be `rec`'s `n` value. The value of the next structure's `n` would be `rec.r->n` (assuming `rec.r!=NULL`).

The syntax of `->` is

*exp1* `->` *exp2*

where *exp1* evaluates to a structure pointer and *exp2* indexes the structure.

## Operators and functions for use with structure members

All operators, all functions, and all features of Mata work with members of structures. That is, given

```
struct example {
    real scalar n
    real matrix X
}

function ...(...)
{
    real scalar rs
    real matrix rm
    struct example scalar ex
    ...
}
```

then `ex.n` and `ex.X` may be used anyplace `rs` and `rm` would be valid.

## Operators and functions for use with entire structures

Some operators and functions can be used with entire structures, not just the structure's elements. Given

```
struct mystruct scalar    ex1, ex2, ex3, ex4
struct mystruct matrix    E, F, G
```

1. You may use `==` and `!=` to test for equality:

```
if (ex1==ex2) ...
if (ex1!=ex2) ...
```

Two structures are equal if their members are equal.

In the example, `struct mystruct` itself contains no substructures. If it did, the definition of equality would include checking the equality of substructures, sub-substructures, etc.

In the example, `ex1` and `ex2` are scalars. If they were matrices, each element would be compared, and the matrices would be equal if the corresponding elements were equal.

2. You may use `:=` and `:=` to form pattern matrices of equality and inequality.
3. You may use the [comma](#) and [backslash](#) operators to form vectors and matrices of structures:

```
ex = ex1, ex2 \ ex3, ex4
```

4. You may use `&` to obtain pointers to structures:

```
ptr_to_ex1 = &ex1
```

5. You may use `subscripting` to access and copy structure members:

```
ex1 = E[1,2]
E[1,2] = ex1
F = E[2,.]
E[2,.] = F
G = E[|1,1\2,2|]
E[|1,1\2,2|] = G
```

6. You may use the `rows()` and `cols()` functions to obtain the number of rows and columns of a matrix of structures.
7. You may use `eltype()` and `orgtype()` with structures. `eltype()` returns `struct`; `orgtype()` returns the usual results.
8. You may use most functions that start with the letters *is*, as in `isreal()`, `iscomplex()`, `isstring()`, etc. These functions return 1 if true and 0 if false and with structures, usually return 0.
9. You may use `swap()` with structures.

## Listing structures

To list the contents of a structure variable, as for debugging purposes, use function `liststruct()`; see [M-5] `liststruct()`.

Using the default, unassigned-expression method to list structures is not recommended, because all that is shown is a pointer value instead of the structure itself.

## Use of transmorphics as `passthru`

A transmorphic matrix can theoretically hold anything, so when we told you that structures had to be explicitly declared, that was not exactly right. Say that function `twopart()` returns a `struct twopart` scalar. You could code

```
x = twopart()
```

without declaring `x` (or declaring it `transmorphic`), and that would not be an error. What you could not do would be to then refer to `x.n1` or `x.n2`, because the compiler would not know that `x` contains a `struct twopart` and so would have no way of interpreting the variable references.

This property can be put to good use in implementing handles and `passthru`.

Say that you are implementing a complicated system. Just to fix ideas, we'll pretend that the system finds the maximum of user-specified functions and that the system has many bells and whistles. To track a given problem, let's assume that your system needs many variables. One variable records the method to be used. Another records whether numerical derivatives are to be used. Another records the current gradient vector. Another records the iteration count, and so on. There might be hundreds of these variables.

You bind all these variables together in one structure:

```
struct maxvariables {
    real scalar  method
    real scalar  use_numeric_d
    real vector  gradient
    real scalar  iteration
    ...
}
```

You design a system with many functions, and some functions call others, but because all the status variables are bound together in one structure, it is easy to pass the values from one function to another.

You also design a system that is easy to use. It starts by the user “opening” a problem,

```
handle = maximize_open()
```

and from that point on the user passes the handle around to the other maximize routines:

```
maximize_set_use_numeric_d(handle, 1)
maximize_set_function_to_max(handle, &myfunc())
...
maximize_maximize_my_function(handle)
```

In this way, you, the programmer of this system, can hold on to values from one call to the next, and you can change the values, too.

What you never do, however, is tell the user that the handle is a `struct maxvariables`. You just tell the user to open a problem by typing

```
handle = maximize_open()
```

and then to pass the handle returned to the other `maximize_*`() routines. If the user feels that he must explicitly declare the handle, you tell him to declare it:

```
transmorphic scalar handle
```

What is the advantage of this secrecy? You can be certain that the user never changes any of the values in your `struct maxvariables` because the compiler does not even know what they are.

Thus you have made your system more robust to user errors.

## Saving compiled structure definitions

You save compiled structure definitions just as you save compiled function definitions; see [\[M-3\] mata mosave](#) and [\[M-3\] mata mlib](#).

When you define a structure, such as `twopart`,

```
struct twopart {
    real scalar  n1, n2
}
```

that also creates a function, `twopart()`, that creates instances of the structure.

Saving `twopart()` in a `.mo` file, or in a `.mlib` library, saves the compiled definition as well. Once `twopart()` has been saved, you may write future programs without bothering to define `struct twopart`. The definition will be automatically found.

## Saving structure variables

Variables containing structures may be saved on disk just as you would save any other variable. No special action is required. See [M-3] [mata matsave](#) and see the function `fputmatrix()` in [M-5] [fopen\(\)](#). `mata matsave` and `fputmatrix()` both work with structure variables, although their entries do not mention them.

## Reference

Gould, W. W. 2007. [Mata Matters: Structures](#). *Stata Journal* 7: 556–570.

## Also see

[M-2] [declarations](#) — Declarations and types

[M-2] [intro](#) — Language definition

# Title

[M-2] **subscripts** — Use of subscripts

## Syntax

$x[\textit{real vector } r, \textit{ real vector } c]$

$x[|\textit{real matrix sub}|]$

Subscripts may be used on the left or right of the [equal-assignment operator](#).

## Description

Subscripts come in two styles.

In `[subscript]` syntax—called list subscripts—an element or a matrix is specified:

$x[1,2]$	the 1,2 element of $x$ ; a scalar
$x[(1\backslash 3\backslash 2), (4,5)]$	the $3 \times 2$ matrix composed of rows 1, 3, and 2 and columns 4 and 5 of $x$ :

$$\begin{bmatrix} x_{14} & x_{15} \\ x_{34} & x_{35} \\ x_{24} & x_{25} \end{bmatrix}$$

In `[|subscript|]` syntax—called range subscripts—an element or a contiguous submatrix is specified:

$x[ 1,2 ]$	same as $x[1,2]$ ; a scalar
$x[ 2,3 \backslash 4,7 ]$	$3 \times 4$ submatrix of $x$ :

$$\begin{bmatrix} x_{23} & x_{24} & x_{25} & x_{26} & x_{27} \\ x_{33} & x_{34} & x_{35} & x_{36} & x_{37} \\ x_{43} & x_{44} & x_{45} & x_{46} & x_{47} \end{bmatrix}$$

Both style subscripts may be used in expressions and may be used on the left-hand side of the equal-assignment operator.

## Remarks

Remarks are presented under the following headings:

[List subscripts](#)

[Range subscripts](#)

[When to use list subscripts and when to use range subscripts](#)

[A fine distinction](#)



## List subscripts

List subscripts—also known simply as subscripts—are obtained when you enclose the subscripts in square brackets, [ and ]. List subscripts come in two basic forms:

$x[ivec, jvec]$	matrix composed of rows <i>ivec</i> and columns <i>jvec</i> of matrix <i>x</i>
$v[kvec]$	vector composed of elements <i>kvec</i> of vector <i>v</i>

where *ivec*, *jvec*, *kvec* may be a vector or a scalar, so the two basic forms include

$x[i, j]$	scalar <i>i, j</i> element
$x[i, jvec]$	row vector of row <i>i</i> , elements <i>jvec</i>
$x[ivec, j]$	column vector of column <i>j</i> , elements <i>ivec</i>
$v[k]$	scalar <i>k</i> th element of vector <i>v</i>

Also missing value may be specified to mean all the rows or all the columns:

$x[i, .]$	row vector of row <i>i</i> of <i>x</i>
$x[. , j]$	column vector of column <i>j</i> of <i>x</i>
$x[ivec, .]$	matrix of rows <i>ivec</i> , all columns
$x[. , jvec]$	matrix of columns <i>jvec</i> , all rows
$x[. , .]$	the entire matrix

Finally, Mata assumes missing value when you omit the argument entirely:

$x[i, ]$	same as $x[i, .]$
$x[ivec, ]$	same as $x[ivec, .]$
$x[, j]$	same as $x[. , j]$
$x[, jvec]$	same as $x[. , jvec]$
$x[, ]$	same as $x[. , .]$

Good style is to specify *ivec* as a column vector and *jvec* as a row vector, but that is not required:

$x[(1\backslash 2\backslash 3), (1, 2, 3)]$	good style
$x[(1, 2, 3), (1\backslash 2\backslash 3)]$	same as $x[(1\backslash 2\backslash 3), (1, 2, 3)]$
$x[(1\backslash 2\backslash 3), (1\backslash 2\backslash 3)]$	same as $x[(1\backslash 2\backslash 3), (1, 2, 3)]$
$x[(1, 2, 3), (1\backslash 2\backslash 3)]$	same as $x[(1\backslash 2\backslash 3), (1, 2, 3)]$

Similarly, good style is to specify *kvec* as a column when *v* is a column vector and to specify *kvec* as a row when *v* is a row vector, but that is not required and what is returned is a column vector if *v* is a column and a row vector if *v* is a row:

$rowv[(1, 2, 3)]$	good style for specifying row vector
$rowv[(1\backslash 2\backslash 3)]$	same as $rowv[(1, 2, 3)]$
$colv[(1\backslash 2\backslash 3)]$	good style for specifying column vector
$colv[(1, 2, 3)]$	same as $colv[(1\backslash 2\backslash 3)]$

Subscripts may be used in expressions following a variable name:

```
first = list[1]
multiplier = x[3,4]
result = colsum(x[,j])
```

Subscripts may be used following an expression to extract a submatrix from a result:

```
allneeded = invsym(x)[(1::4), .] * multiplier
```

Subscripts may be used on the left-hand side of the equal-assignment operator:

```
x[1,1] = 1
x[1,.] = y[3,.]
x[(1::4), (1..4)] = I(4)
```

## Range subscripts

Range subscripts appear inside the difficult to type `[|` and `|]` brackets. Range subscripts come in four basic forms:

<code>x[ i,j ]</code>	<i>i,j</i> element; same result as <code>x[i,j]</code>
<code>v[ k ]</code>	<i>k</i> th element of vector; same result as <code>v[k]</code>
<code>x[ i,j \ k,l ]</code>	submatrix, vector, or scalar formed using ( <i>i,j</i> ) as top-left corner and ( <i>k,l</i> ) as bottom-right corner
<code>v[ i \ k ]</code>	subvector or scalar of elements <i>i</i> through <i>k</i> ; result is row vector if <i>v</i> is row vector, column vector if <i>v</i> is column vector

Missing value may be specified for a row or column to mean all rows or all columns when a  $1 \times 2$  or  $1 \times 1$  subscript is specified:

<code>x[ i, . ]</code>	row <i>i</i> of <i>x</i> ; same as <code>x[i, .]</code>
<code>x[  . , j ]</code>	column <i>j</i> of <i>x</i> ; same as <code>x[ . , j]</code>
<code>x[  . , . ]</code>	entire matrix; same as <code>x[ . , .]</code>
<code>v[  .  ]</code>	entire vector; same as <code>v[.]</code>

Also missing may be specified to mean the number of rows or the number of columns of the matrix being subscripted when a  $2 \times 2$  subscript is specified:

<code>x[ 1,2 \ 4, . ]</code>	equivalent to <code>x[ 1,2 \ 4, cols(x) ]</code>
<code>x[ 1,2 \ . , 3 ]</code>	equivalent to <code>x[ 1,2 \ rows(x), 3 ]</code>
<code>x[ 1,2 \ . , . ]</code>	equivalent to <code>x[ 1,2 \ rows(x), cols(x) ]</code>

With range subscripts, what appears inside the square brackets is in all cases interpreted as a matrix expression, so in

```
sub = (1,2)
... x[|sub|] ...
```

`x[sub]` refers to `x[1,2]`.

Range subscripts may be used in all the same contexts as list subscripts; they may be used in expressions following a variable name

```
submat = result[|1,1 \ 3,3|]
```

they may be used to extract a submatrix from a calculated result

```
allneeded = invsym(x)[|1,1 \ 4,4|]
```

and they may be used on the left-hand side of the equal-assignment operator:

```
x[|1,1 \ 4,4|] = I(4)
```

## When to use list subscripts and when to use range subscripts

Everything a range subscript can do, a list subscript can also do. The one seemingly unique feature of a range subscript,

```
x[|i1,j1 \ i2,j2|]
```

is perfectly mimicked by

```
x[(i1:i2), (j1..j2)]
```

The range-subscript construction, however, executes more quickly, and so that is the purpose of range subscripts: to provide a fast way to extract contiguous submatrices. In all other cases, use list subscripts because they are faster.

Use list subscripts to refer to scalar values:

```
result = x[1,3]
x[1,3] = 2
```

Use list subscripts to extract entire rows or columns:

```
obs = x[., 3]
var = x[4, .]
```

Use list subscripts to permute the rows and columns of matrices:

```
: x = (1,2,3,4 \ 5,6,7,8 \ 9,10,11,12)
: y = x[(1\3\2), .]
: y
```

	1	2	3	4
1	1	2	3	4
2	9	10	11	12
3	5	6	7	8

```
: y = x[., (1,3,2,4)]
: y
```

	1	2	3	4
1	1	3	2	4
2	5	7	6	8
3	9	11	10	12

```
: y=x[(1\3\2), (1,3,2,4)]
```

```

: y

```

	1	2	3	4
1	1	3	2	4
2	9	11	10	12
3	5	7	6	8

Use list subscripts to duplicate rows or columns:

```

: x = (1,2,3,4 \ 5,6,7,8 \ 9,10,11,12)
: y = x[(1\2\3\1), .]
: y

```

	1	2	3	4
1	1	2	3	4
2	5	6	7	8
3	9	10	11	12
4	1	2	3	4

```

: y = x[., (1,2,3,4,2)]
: y

```

	1	2	3	4	5
1	1	2	3	4	2
2	5	6	7	8	6
3	9	10	11	12	10

```

: y = x[(1\2\3\1), (1,2,3,4,2)]
: y

```

	1	2	3	4	5
1	1	2	3	4	2
2	5	6	7	8	6
3	9	10	11	12	10
4	1	2	3	4	2

### A fine distinction

There is a fine distinction between  $x[i,j]$  and  $x[|i,j|]$ . In  $x[i,j]$ , there are two arguments,  $i$  and  $j$ . The comma separates the arguments. In  $x[|i,j|]$ , there is one argument:  $i,j$ . The comma is the [column-join operator](#).

In Mata, comma means mostly the column-join operator:

```
newvec = oldvec, addedvalues
qsum = (x,1)'(x,1)
```

There are, in fact, only two exceptions. When you type the arguments for a function, the comma separates one argument from the next:

```
result = f(a,b,c)
```

In the above example,  $f()$  receives three arguments:  $a$ ,  $b$ , and  $c$ . If we wanted  $f()$  to receive one argument,  $(a,b,c)$ , we would have to enclose the calculation in parentheses:

```
result = f((a,b,c))
```

That is the first exception. When you type the arguments inside a function, comma means argument separation. You get back to the usual meaning of comma—the column-join operator—by opening another set of parentheses.

The second exception is in [list subscripting](#):

$$x[i,j]$$

Inside the list-subscript brackets, comma means argument separation. That is why you have seen us type vectors inside parentheses:

$$x[(1\backslash 2\backslash 3), (1,2,3)]$$

These are the two exceptions. Range subscripting is not an exception. Thus in

$$x[|i, j|]$$

there is one argument,  $i,j$ . With range subscripts, you may program constructs such as

```
IJ      = (i,j)
RANGE  = (1,2 \ 4,4)
...
... x[|IJ|] ... x[|RANGE|] ...
```

You may not code in this way with list subscripts. In particular,  $x[IJ]$  would be interpreted as a request to extract elements  $i$  and  $j$  from vector  $x$ , and would be an error otherwise.  $x[RANGE]$  would always be an error.

We said earlier that list subscripts  $x[i,j]$  are a little faster than range subscripts  $x[|i,j|]$ . That is true, but if  $IJ=(i,j)$  already,  $x[|IJ|]$  is faster than  $x[i,j]$ . You would, however, have to execute many millions of references to  $x[|IJ|]$  before you could measure the difference.

## Conformability

$x[i, j]$ :

$x$ :	$r \times c$			
$i$ :	$m \times 1$	or	$1 \times m$	(does not matter which)
$j$ :	$1 \times n$	or	$n \times 1$	(does not matter which)
result:	$m \times n$			

$x[i, .]$ :

$x$ :	$r \times c$			
$i$ :	$m \times 1$	or	$1 \times m$	(does not matter which)
result:	$m \times c$			

$x[. , j]$ :

$x$ :	$r \times c$			
$j$ :	$1 \times n$	or	$n \times 1$	(does not matter which)
result:	$r \times n$			

$x[. , .]$ :

$x$ :	$r \times c$
result:	$r \times c$

$x[i]:$	$x:$	$n \times 1$		$1 \times n$
	$i:$	$m \times 1$	or	$1 \times m$
	$result:$	$m \times 1$		$1 \times m$ or $m \times 1$
$x[.]:$	$x:$	$n \times 1$		$1 \times n$
	$result:$	$n \times 1$		$1 \times n$
$x[ k ]:$	$x:$	$r \times c$		
	$k:$	$1 \times 2$		
	$result:$	$1 \times 1$	if $k[1]<.$ and $k[2]<.$	
		$r \times 1$	if $k[1]>=.$ and $k[2]<.$	
		$1 \times c$	if $k[1]<.$ and $k[2]>=.$	
		$r \times c$	if $k[1]>=.$ and $k[2]>=.$	
$x[ k ]:$	$x:$	$r \times c$		
	$k:$	$2 \times 2$		
	$result:$	$k[2,1]-k[1,1]+1 \times k[2,2]-k[1,2]+1$		
		(in the above formula, if $k[2,1]>=.$ , treat as if $k[2,1]=r$ , and similarly, if $k[2,2]>=.$ , treat as if $k[2,2]=c$ )		
$x[ k ]:$	$x:$	$r \times 1$		$1 \times c$
	$k:$	$2 \times 1$		$2 \times 1$
	$result:$	$k[2]-k[1]+1 \times 1$		$1 \times k[2]-k[1]+1$
		(if $k[2]>=.$ , treat as if $k[2]=r$ )		(if $k[2]>=.$ , treat as if $k[2]=c$ )

## Diagnostics

Both styles of subscripts abort with error if the subscript is out of range, if a reference is made to a nonexisting row or column.

## Reference

Gould, W. W. 2007. [Mata Matters: Subscripting](#). *Stata Journal* 7: 106–116.

## Also see

[\[M-2\] intro](#) — Language definition

## Syntax

The basic language syntax is

*istmt*

where

```
istmt :=      stmt
               function name(farglist) fstmt
               ftype name(farglist) fstmt
               ftype function name(farglist) fstmt

stmt :=       nothing
               ;                                     (meaning nothing)
               version number
               { stmt ... }
               exp
               pragma pstmt
               if (exp) stmt
               if (exp) stmt else stmt
               for (exp;exp;exp) stmt
               while (exp) stmt
               do stmt while (exp)
               break
               continue
               label:
               goto label
               return
               return(exp)

fstmt :=      stmt
               type arglist
               external type arglist

arglist :=     name
               name()
               name, arglist
               name() , arglist

farglist :=    nothing
               efarglist

efarglist :=   felement
               felement, efarglist
               | felement
               | felement, efarglist
```

<i>felement</i> :=	<i>name</i> <i>type name</i> <i>name()</i> <i>type name()</i>
<i>ftype</i> :=	<i>type</i> void
<i>type</i> :=	<i>eltype</i> <i>orgtype</i> <i>eltype orgtype</i>
<i>eltype</i> :=	transmorphic string numeric real complex pointer pointer( <i>ptrtype</i> )
<i>orgtype</i> :=	matrix vector rowvector colvector scalar
<i>ptrtype</i> :=	<i>nothing</i> <i>type</i> <i>type function</i> function
<i>pstmt</i> :=	unset <i>name</i> unused <i>name</i>
<i>name</i> :=	identifier up to 32 characters long
<i>label</i> :=	identifier up to 8 characters long
<i>exp</i> :=	expression as defined in <a href="#">[M-2]</a> <b>exp</b>

## Description

Mata is a C-like compiled-into-pseudocode language with matrix extensions and run-time linking.

## Remarks

Remarks are presented under the following headings:

- [Treatment of semicolons](#)
- [Types and declarations](#)
- [Void matrices](#)
- [Void functions](#)
- [Operators](#)
- [Subscripts](#)
- [Implied input tokens](#)
- [Function argument-passing convention](#)
- [Passing functions to functions](#)
- [Optional arguments](#)



After reading [M-2] [syntax](#), see [M-2] [intro](#) for a list of entries that give more explanation of what is discussed here.

## Treatment of semicolons

Semicolon (;) is treated as a line separator. It is not required, but it may be used to place two statements on the same physical line:

```
x = 1 ; y = 2 ;
```

The last semicolon in the above example is unnecessary but allowed.

Single statements may continue onto more than one line if the continuation is obvious. Take “obvious” to mean that there is a hanging open parenthesis or a hanging dyadic operator; for example,

```
x = (
      3)
x = x +
      2
```

See [M-2] [semicolons](#) for more information.

## Types and declarations

The *type* of a variable or function is described by

*eltype orgtype*

where *eltype* and *orgtype* are each one of

<i>eltype</i>	<i>orgtype</i>
transmorphic	matrix
numeric	vector
real	rowvector
complex	colvector
string	scalar
pointer	

For example, a variable might be real scalar, or complex matrix, or string vector.

Mata also has structures—the *eltype* is `struct name`—but these are not discussed here. For a discussion of structures, see [M-2] [struct](#).

Mata also has classes—the *eltype* is `class name`—but these are not discussed here. For a discussion of classes, see [M-2] [class](#).

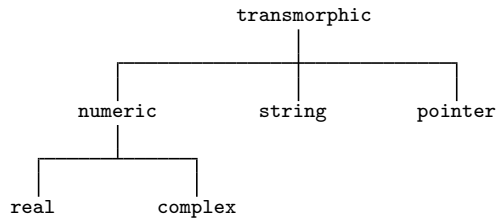
Declarations are optional. When the *type* of a variable or function is not declared, it is assumed to be a transmorphic matrix. In particular:

1. *eltype* specifies the type of the elements. When *eltype* is not specified, transmorphic is assumed.

2. *orgtype* specifies the organization of the elements. When *orgtype* is not specified, *matrix* is assumed.

All *types* are special cases of *transmorphic matrix*.

The nesting of *eltypes* is



*orgtypes* amount to nothing more than a constraint on the number of rows and columns of a matrix:

<i>orgtype</i>	Constraint
matrix	$r \geq 0$ and $c \geq 0$
vector	$r = 1$ and $c \geq 0$ or $r \geq 0$ and $c = 1$
rowvector	$r = 1$ and $c \geq 0$
colvector	$r \geq 0$ and $c = 1$
scalar	$r = 1$ and $c = 1$

See [M-2] [declarations](#).

## Void matrices

A matrix (vector, row vector, or column vector) that is  $0 \times 0$ ,  $r \times 0$ , or  $0 \times c$  is said to be *void*; see [M-2] [void](#).

The function `J(r, c, val)` returns an  $r \times c$  matrix with each element containing *val*; see [M-5] [J\(\)](#).

`J()` can be used to create void matrices.

See [M-2] [void](#).

## Void functions

Rather than *eltype orgtype*, a function can be declared to return nothing by being declared to return *void*:

```
void function example(matrix A)
{
    real scalar i
    for (i=1; i<=rows(A); i++) A[i,i] = 1
}
```

A function that returns nothing (does not include a `return(exp)` statement), in fact returns `J(0, 0, .)`, and the above function could equally well be coded as

```
void function example(matrix A)
{
    real scalar    i
    for (i=1; i<=rows(A); i++) A[i,i] = 1
    return(J(0, 0, .))
}
```

or

```
void function example(matrix A)
{
    real scalar    i
    for (i=1; i<=rows(A); i++) A[i,i] = 1
    return(J(0,0,.))
}
```

Therefore, void also is a special case of transmorphic matrix (it is in fact a  $0 \times 0$  real matrix). Since declarations are optional (but recommended both for reasons of style and for reasons of efficiency), the above function could also be coded as

```
function example(A)
{
    for (i=1; i<=rows(A); i++) A[i,i] = 1
}
```

See [\[M-2\] declarations](#).

## Operators

Mata provides the usual assortment of operators; see [\[M-2\] exp](#).

The monadic prefix operators are

```
-    !    ++    --    &    *
```

Prefix operators `&` and `*` have to do with pointers; see [\[M-2\] pointers](#).

The monadic postfix operators are

```
'    ++    --
```

Note the inclusion of postfix operator `'` for transposition. Also, for  $Z$  complex,  $Z'$  returns the conjugate transpose. If you want the transposition without conjugation, see [\[M-5\] transposeonly\(\)](#).

The dyadic operators are

```
=    ?    \    ::    ,    ..    |    &    ==    >=    <=    <    >
!=    +    -    *    #    ^
```

Also, `&&` and `||` are included as synonyms for `&` and `|`.

The operators `==` and `!=` do not require conformability, nor do they require that the matrices be of the same type. In such cases, the matrices are unequal (`==` is false and `!=` is true). For complex arguments, `<`, `<=`, `>`, and `>=` refer to length of the complex vector. `==` and `!=`, however, refer not to length but to actual components. See [\[M-2\] op\\_logical](#).

The operators `,` and `\` are the row-join and column-join operators.  $(1,2,3)$  constructs the row vector  $(1,2,3)$ .  $(1\backslash 2\backslash 3)$  constructs the column vector  $(1,2,3)'$ .  $(1,2\backslash 3,4)$  constructs the matrix with first row  $(1,2)$  and second row  $(3,4)$ .  $a,b$  joins two scalars, vectors, or matrices rowwise.  $a\backslash b$  joins two scalars, vectors, or matrices columnwise. See [M-2] [op\\_join](#).

`..` and `::` refer to the row-to and column-to operators.  $1..5$  is  $(1,2,3,4,5)$ .  $1::5$  is  $(1\backslash 2\backslash 3\backslash 4\backslash 5)$ .  $5..1$  is  $(5,4,3,2,1)$ .  $5::1$  is  $(5\backslash 4\backslash 3\backslash 2\backslash 1)$ . See [M-2] [op\\_range](#).

For `|`, `&`, `==`, `>=`, `<=`, `<`, `>`, `!=`, `+`, `-`, `*`, `/`, and `^`, there is `:op` at precedence just below `op`. These operators perform the elementwise operation. For instance,  $A*B$  refers to matrix multiplication;  $A:*B$  refers to elementwise multiplication. Moreover, elementwise is generalized to cases where  $A$  and  $B$  do not have the same number of rows and the same number of columns. For instance, if  $A$  is a  $1 \times c$  row vector and  $B$  is a  $r \times c$  matrix, then  $||C_{ij}|| = ||A_j|| * ||B_{ij}||$  is returned. See [M-2] [op\\_colon](#).

Subscripts

$A[i,j]$  returns the  $i,j$  element of  $A$ .

$A[k]$  returns  $A[1,k]$  if  $A$  is  $1 \times c$  and  $A[k,1]$  if  $A$  is  $r \times 1$ . That is, in addition to declared vectors, any  $1 \times c$  matrix or  $r \times 1$  matrix may be subscripted by one index. Similarly, any vector can be subscripted by two indices.

$i, j$ , and  $k$  may be vectors as well as scalars. For instance,  $A[(3\backslash 4\backslash 5), 4]$  returns a  $3 \times 1$  column vector containing rows 3 to 5 of the 4th column.

$i, j$ , and  $k$  may be missing value.  $A[., 4]$  returns a column vector of the 4th column of  $A$ .

The above subscripts are called list-style subscripts. Mata provides a second format called range-style subscripts that is especially useful for selecting submatrices.  $A[|3,3\backslash 5,5|]$  returns the  $3 \times 3$  submatrix of  $A$  starting at  $A[3,3]$ .

See [M-2] [subscripts](#).

Implied input tokens

Before interpreting and compiling a line, Mata makes the following substitutions to what it sees:

Input sequence	Interpretation
<i>'name</i>	<i>'*name</i>
[ ,	[ . ,
, ]	, . ]

Hence, coding  $X'Z$  is equivalent to coding  $X'*Z$ , and coding  $x = z[1,]$  is equivalent to coding  $x = z[1, .]$ .

## Function argument-passing convention

Arguments are passed to functions by address, also known as by name or by reference. They are not passed by value. When you code

```
... f(A) ...
```

it is the address of `A` that is passed to `f()`, not a copy of the values in `A`. `f()` can modify `A`.

Most functions do not modify their arguments, but some do. `lud(A, L, U, p)`, for instance, calculates the LU decomposition of `A`. The function replaces the contents of `L`, `U`, and `p` with matrices such that  $L[p,]*U = A$ .

Oldtimers will have heard of the FORTRAN programmer who called a subroutine and passed to it a second argument of 1. Unbeknownst to him, the subroutine changed its second argument, with the result that the constant 1 was changed throughout the rest of his code. That cannot happen in Mata. When an expression is passed as an argument (and constants are expressions), a temporary variable containing the evaluation is passed to the function. Modifications to the temporary variable are irrelevant because the temporary variable is discarded once the function returns. Thus if `f()` modifies its second argument and you call it by coding `f(A, 2)`, because 2 is copied to a temporary variable, the value of the literal 2 will remain unchanged on the next call.

If you call a function with an expression that includes the assignment operator, it is the left-hand side of the expression that is passed. That is, coding

```
f(a, b=c)
```

has the same result as coding

```
b = c
f(a, b)
```

If function `f()` changes its second argument, it will be `b` and not `c` that is modified.

Also, Mata attempts not to create unnecessary copies of matrices. For instance, consider

```
function changearg(x) x[1,1] = 1
```

`changearg(mymat)` changes the 1,1 element of `mymat` to 1. Now let us define

```
function cp(x) return(x)
```

Coding `changearg(cp(mymat))` would still change `mymat` because `cp()` returned `x` itself. On the other hand, if we defined `cp()` as

```
function cp(x)
{
    matrix t
    t = x
    return(t)
}
```

then coding `changearg(cp(mymat))` would not change `mymat`. It would change a temporary matrix which would be discarded once `changearg()` returned.

## Passing functions to functions

One function may receive another function as an argument using pointers. One codes

```
function myfunc(pointer(function) f, a, b)
{
    ... (*f)(a) ... (*f)(b) ...
}
```

although the `pointer(function)` declaration, like all declarations, is optional. To call `myfunc()` and tell it to use function `prima()` for `f()`, and 2 and 3 for `a` and `b`, one codes

```
myfunc(&prima(), 2, 3)
```

See [M-2] [ftof](#) and [M-2] [pointers](#).

## Optional arguments

Functions may be coded to allow receiving a variable number of arguments. This is done by placing a vertical or bar (`|`) in front of the first argument that is optional. For instance,

```
function mynorm(matrix A, |scalar power)
{
    ...
}
```

The above function may be called with one matrix or with a matrix followed by a scalar.

The function `args()` (see [M-5] [args\(\)](#)) can be used to determine the number of arguments received and to set defaults:

```
function mynorm(matrix A, |scalar power)
{
    ...
    (args()==1) power = 2
    ...
}
```

See [M-2] [optargs](#).

## Reference

Gould, W. W. 2005. *Mata Matters: Translating Fortran*. *Stata Journal* 5: 421–441.

## Also see

[M-2] [intro](#) — Language definition

## Syntax

### *Syntax 1*

```
. version #[. #]
. mata:
: ...
: function name(...)
: {
:     ...
: }
: ...
: end
```

### *Syntax 2*

```
: function name(...)
: {
:     version #[. #]
:     ...
: }
```

## Description

In syntax 1, Stata's `version` command (see [\[P\] version](#)) sets the version before entering Mata. This specifies both the compiler and library versions to be used. Syntax 1 is recommended.

In syntax 2, Mata's `version` command sets the version of the library functions that are to be used. Syntax 2 is rarely used.

## Remarks

Remarks are presented under the following headings:

[Purpose of version control](#)  
[Recommendations for do-files](#)  
[Recommendations for ado-files](#)  
[Compile-time and run-time versioning](#)

## Purpose of version control

Mata is under continual development, which means not only that new features are being added but also that old features sometimes change how they work. Old features changing how they work is supposedly an improvement—it generally is—but that also means old programs might stop working or, worse, work differently.

version provides the solution.

If you are working interactively, nothing said here matters.

If you use Mata in do-files or ado-files, we recommend that you set version before entering Mata.

## Recommendations for do-files

The recommendation for do-files that use Mata is the same as for do-files that do not use Mata: specify the version number of the Stata you are using on the top line:

```
----- begin myfile.do -----  
version 12  
...  
----- end myfile.do -----
```

To determine the number that should appear after version, type about at the Stata prompt:

```
. about  
Stata/SE 12.0  
(output omitted)
```

We are using Stata 12.0.

Coding version 12 will not benefit us today but, in the future, we will be able to rerun our do-file and obtain the same results.

By the way, a do-file is any file that you intend to execute using Stata's do or run commands (see [\[R\] do](#)), regardless of the file suffix. Many users (us included) save Mata source code in .mata files and then type do *myfile.mata* to compile. .mata files are do-files; we include the version line:

```
----- begin myfile.mata -----  
version 12  
mata:  
...  
end  
----- end myfile.mata -----
```

## Recommendations for ado-files

Mata functions may be included in ado-files; see [\[M-1\] ado](#). In such files, set version before entering Mata along with, as usual, setting the version at the top of your program:

```
----- begin myfile.ado -----  
  
program myfile  
    version 12    ← as usual  
    ...  
end  
  
version 12        ← new  
mata:  
...  
end  
----- end myfile.ado -----
```



## Compile-time and run-time versioning

What follows is detail. We recommend always following the recommendations above.

There are actually two version numbers that matter—the version number set at the time of compilation, which affects how the source code is interpreted, and the version of the libraries used to supply subroutines at the time of execution.

The `version` command that we used in the previous sections is in fact Stata's `version` command (see [\[P\] version](#)), and it sets both versions:

```
. version 12
. mata:
: function example()
: {
:     ...
: }
: end
```

In the above, we compile `example()` by using the version 12 syntax of the Mata language, and any functions `example()` calls will be the 12 version of those functions. Setting `version 12` before entering Mata ensured all of that.

In the following example, we compile using version 12 syntax and use version 12.2 functions:

```
. version 12
. mata:
: function example()
: {
:     version 12.2
:     ...
: }
: end
```

In the following example, we compile using version 12.2 syntax and use version 12 functions:

```
. version 12.2
. mata:
: function example()
: {
:     version 12
:     ...
: }
: end
```

It is, however, very rare that you will want to compile and execute at different version levels.

## Also see

[\[M-5\] callersversion\(\)](#) — Obtain version number of caller

[\[M-2\] intro](#) — Language definition

## Syntax

<code>J(0, 0, .)</code>	$0 \times 0$ real matrix
<code>J(<i>r</i>, 0, .)</code>	$r \times 0$ real matrix
<code>J(0, <i>c</i>, .)</code>	$0 \times c$ real matrix
<code>J(0, 0, 1i)</code>	$0 \times 0$ complex matrix
<code>J(<i>r</i>, 0, 1i)</code>	$r \times 0$ complex matrix
<code>J(0, <i>c</i>, 1i)</code>	$0 \times c$ complex matrix
<code>J(0, 0, "")</code>	$0 \times 0$ string matrix
<code>J(<i>r</i>, 0, "")</code>	$r \times 0$ string matrix
<code>J(0, <i>c</i>, "")</code>	$0 \times c$ string matrix
<code>J(0, 0, NULL)</code>	$0 \times 0$ pointer matrix
<code>J(<i>r</i>, 0, NULL)</code>	$r \times 0$ pointer matrix
<code>J(0, <i>c</i>, NULL)</code>	$0 \times c$ pointer matrix

## Description

Mata allows  $0 \times 0$ ,  $r \times 0$ , and  $0 \times c$  matrices. These matrices are called *void matrices*.

## Remarks

Remarks are presented under the following headings:

[Void matrices, vectors, row vectors, and column vectors](#)  
[How to read conformability charts](#)

### Void matrices, vectors, row vectors, and column vectors

Void matrices contain nothing, but they have dimension information (they are  $0 \times 0$ ,  $r \times 0$ , or  $0 \times c$ ) and have an *eltype* (which is real, complex, string, or pointer):

1. A matrix is said to be void if it is  $0 \times 0$ ,  $r \times 0$ , or  $0 \times c$ .
2. A vector is said to be void if it is  $0 \times 1$  or  $1 \times 0$ .
3. A column vector is said to be void if it is  $0 \times 1$ .
4. A row vector is said to be void if it is  $1 \times 0$ .
5. A scalar cannot be void because it is, by definition,  $1 \times 1$ .

The function `J(r, c, val)` creates  $r \times c$  matrices containing *val*; see [M-5] `J()`. `J()` can be used to manufacture void matrices by specifying *r* and/or *c* as 0. The value of the third argument does not matter, but its *eltype* does:

1. `J(0,0,.)` creates a real  $0 \times 0$  matrix, as will `J(0,0,1)` and as will `J()` with any real third argument.
2. `J(0,0,1i)` creates a  $0 \times 0$  complex matrix, as will `J()` with any complex third argument.
3. `J(0,0,"")` creates  $0 \times 0$  string matrices, as will `J()` with any string third argument.
4. `J(0,0,NULL)` creates  $0 \times 0$  pointer matrices, as will `J()` with any pointer third argument.

In fact, one rarely needs to manufacture such matrices because they arise naturally in extreme cases. Similarly, one rarely needs to include special code to handle void matrices because such matrices handle themselves. Loops vanish when the number of rows or columns are zero.

## How to read conformability charts

In general, not only is no emphasis placed on how functions and operators deal with void matrices, no mention is even made of the fact. Instead, the information is buried in the *Conformability* section located near the end of the function's or operator's manual entry.

For instance, the conformability chart for some function might read

```
somefunction(A, B, v):
  A:      r × c
  B:      c × k
  v:      1 × k  or  k × 1
  result:  r × k
```

Among other things, the chart above is stating how `somefunction()` handles void matrices. *A* must be  $r \times c$ . That chart does not say

*A*:  $r \times c, r > 0, c > 0$

and that is what it would have said if `somefunction()` did not allow *A* to be void. Hence, *A* may be  $0 \times 0$ ,  $0 \times c$ , or  $r \times 0$ .

Similarly, *B* may be void as long as `rows(B)==cols(A)`. *v* may be void if `cols(B)==0`. The returned result will be void if `rows(A)==0` or `cols(B)==0`.

Interestingly, `somefunction()` can produce a nonvoid result from void input. For instance, if *A* were  $5 \times 0$  and *B*,  $0 \times 3$ , a  $5 \times 3$  result would be produced. It is interesting to speculate what would be in that  $5 \times 3$  result. Probably, if we knew what `somefunction()` did, it would be obvious to us, but if it were not, the following section, *Diagnostics*, would state what the surprising result would be.

As a real example, see [M-5] `trace()`. `trace()` will take the trace of a  $0 \times 0$  matrix. The result is 0. Or see multiplication (\*) in [M-2] `op_arith`. One can multiply a  $k \times 0$  matrix by a  $0 \times m$  matrix to produce a  $k \times m$  result. The matrix will contain zeros.

## Also see

[M-2] [intro](#) — Language definition

# Title

[M-2] **while** — while (*exp*) *stmt*

## Syntax

```
while (exp) stmt
```

```
while (exp) {  
    stmts  
}
```

where *exp* must evaluate to a real scalar.

## Description

**while** executes *stmt* or *stmts* zero or more times. The loop continues as long as *exp* is not equal to zero.

## Remarks

To understand **while**, enter the following program

```
function example(n)  
{  
    i = 1  
    while (i<=n) {  
        printf("i=%g\n", i)  
        i++  
    }  
    printf("done\n")  
}
```

and run `example(3)`, `example(2)`, `example(1)`, `example(0)`, and `example(-1)`.

One common use of **while** is to loop until convergence:

```
while (mreldif(a, lasta)>1e-10) {  
    lasta = a  
    a = ...  
}
```

## Also see

[M-2] [semicolons](#) — Use of semicolons

[M-2] [do](#) — do ... while (exp)

[M-2] [for](#) — for (exp1; exp2; exp3) stmt

[M-2] [break](#) — Break out of for, while, or do loop

[M-2] [continue](#) — Continue with next iteration of for, while, or do loop

[M-2] [intro](#) — Language definition



## **[M-3] Commands for controlling Mata**





Contents

Command for invoking Mata from Stata:

[M-3] Entry	Command	Description
<b>mata</b>	. mata	invoke Mata

Once you are running Mata, you can use the following commands from the colon prompt:

[M-3] Entry	Command	Description
<b>mata help</b>	: mata help	execute help command
<b>mata clear</b>	: mata clear	clear Mata
<b>mata describe</b>	: mata describe	describe contents of Mata’s memory
<b>mata memory</b>	: mata memory	display memory-usage report
<b>mata rename</b>	: mata rename	rename matrix or function
<b>mata drop</b>	: mata drop	remove from memory matrix or function
<b>mata mosave</b>	: mata mosave	create object file
<b>mata mlib</b>	: mata mlib	create function library
<b>mata matsave</b>	: mata matsave	save matrices
<b>mata matsave</b>	: mata matuse	restore matrices
<b>mata matsave</b>	: mata matdescribe	describe contents of matrix file
<b>mata which</b>	: mata which	identify function
<b>mata set</b>	: mata query	display values of settable parameters
<b>mata set</b>	: mata set	set parameters
<b>mata stata</b>	: mata stata	execute Stata command
<b>end</b>	: end	exit Mata and return to Stata

Description

When you type something at the Mata prompt, it is assumed to be a Mata statement—something that can be compiled and executed—such as

```
: 2+3
5
```

The `mata` command, however, is different. When what you type is prefixed by the word `mata`, think of yourself as standing outside of Mata and giving an instruction that affects the Mata environment and the way Mata works. For instance, typing

```
: mata clear
```

says that Mata is to be cleared. Typing

```
: mata set matastrict on
```

says that Mata is to require that programs explicitly declare their arguments and their working variables; see [M-2] [declarations](#).

## Remarks

The `mata` command cannot be used inside functions. It would make no sense to code

```
function foo(...)
{
    ...
    mata query
    ...
}
```

because `mata query` is something that can be typed only at the Mata colon prompt:

```
: mata query
(output omitted)
```

See [M-1] [how](#).

## Also see

[M-0] [intro](#) — Introduction to the Mata manual

# Title

[M-3] **end** — Exit Mata and return to Stata

## Syntax

`: end`

## Description

`end` exits Mata and returns to Stata.

## Remarks

When you exit from Mata back into Stata, Mata does not clear itself; so if you later return to Mata, you will be right back where you were. See [\[M-3\] `mata`](#).

## Also see

[\[M-3\] `intro`](#) — Commands for controlling Mata

Syntax

The `mata` command documented here is for use from Stata. It is how you enter Mata. You type `mata` at a Stata dot prompt, not a Mata colon prompt.

Syntax 1	Comment
<code>mata</code>	no colon following <code>mata</code>
<i>istmt</i>	
<i>istmt</i>	if an error occurs, you stay in
<code>..</code>	mata mode
<i>istmt</i>	
<code>end</code>	you exit when you type <code>end</code>

Syntax 1 is the best way to use Mata interactively.

Syntax 2	Comment
<code>mata:</code>	colon following <code>mata</code>
<i>istmt</i>	
<i>istmt</i>	if an error occurs, you are
<code>..</code>	dumped from <code>mata</code>
<i>istmt</i>	
<code>end</code>	otherwise, you exit when you type <code>end</code>

Syntax 2 is mostly used by programmers in ado-files.  
Programmers want errors to stop everything.

Syntax 3	Comment
<code>mata <i>istmt</i></code>	rarely used

Syntax 3 is the single-line variant of syntax 1, but it is not useful.

Syntax 4	Comment
<code>mata: <i>istmt</i></code>	for use by programmers

Syntax 4 is the single-line variant of syntax 2, and it exists for the same reason as syntax 2: for use by programmers in ado-files.

## Description

The `mata` command invokes Mata. An *istmt* is something Mata understands; *istmt* stands for interactive statement of Mata.

## Remarks

Remarks are presented under the following headings:

*Introduction*

*The fine distinction between syntaxes 3 and 4*

*The fine distinction between syntaxes 1 and 2*

## Introduction

For interactive use, use [syntax 1](#). Type `mata` (no colon), press *Enter*, and then use Mata freely. Type `end` to return to Stata. (When you exit from Mata back into Stata, Mata does not clear itself; so if you later type `mata`-followed-by-*enter* again, you will be right back where you were.)

For programming use, use [syntax 2](#) or [syntax 4](#). Inside a program or an ado-file, you can just call a Mata function

```
program myprog
...
mata: utility("varlist")
...
end
```

and you can even include that Mata function in your ado-file

---

```
begin myprog.ado

program myprog
...
mata: utility("varlist")
...
end

mata:
function utility(string scalar varlist)
{
...
}
end
```

---

```
end myprog.ado
```

---

or you could separately compile `utility()` and put it in a `.mo` file or in a Mata library.

## The fine distinction between syntaxes 3 and 4

Syntaxes [3](#) and [4](#) are both single-line syntaxes. You type `mata`, perhaps a colon, and follow that with the Mata *istmt*.

The differences between the two syntaxes is whether they allow continuation lines. With a colon, no continuation line is allowed. Without a colon, you may have continuation lines.

For instance, let's consider

```
function renorm(scalar a, scalar b)
{
    ...
}
```

No matter how long the function, it is one *istmt*. Using `mata:`, if you were to try to enter that *istmt*, here is what would happen:

```
. mata: function renorm(scalar a, scalar b)
<istmt> incomplete
r(197);
```

When you got to the end of the first line and pressed *Enter*, you got an error message. Using the `mata:` command, the *istmt* must all fit on one line.

Now try the same thing using `mata` without the colon:

```
. mata function renorm(scalar a, scalar b)
> {
>     ...
> }
.
```

That worked! Single-line `mata` without the colon allows continuation lines and, on this score at least, seems better than single-line `mata` with the colon. In programming contexts, however, this feature can bite. Consider the following program fragment:

```
program example
    ...
    mata utility("`varlist'"
    replace `x' = ...
    ...
end
```

We used `mata` without the colon, and we made an error: we forgot the close parenthesis. `mata` without the colon will be looking for that close parenthesis and so will eat the next line—a line not intended for Mata. Here we will get an error message because “`replace `x' = ...`” will make no sense to Mata, but that error will be different from the one we should have gotten. In the unlikely worse case, that next line will make sense to Mata.

Ergo, programmers want to include the colon. It will make your programs easier to debug.

There is, however, a programmer’s use for single-line `mata` without the colon. In our sample `ado`-file above when we included the routine `utility()`, we bound it in `mata:` and `end`. It would be satisfactory if instead we coded

---

```

program myprog
    ...
    mata: utility("`varlist'")
    ...
end

mata function utility(string scalar varlist)
{
    ...
}

```

---

begin myprog.ado

end myprog.ado

Using `mata` without the colon, we can omit the `end`. We admit we sometimes do that.

## The fine distinction between syntaxes 1 and 2

Nothing said above about continuation lines applies to syntaxes 1 and 2. The multiline `mata`, with or without colon, always allows continuation lines because where the Mata session ends is clear enough: `end`.

The difference between the two multiline syntaxes is whether Mata tolerates errors or instead dumps you back into Stata. Interactive users appreciate tolerance. Programmers want strictness. Programmers, consider the following (using `mata` without the colon):

```
program example2
    ...
    mata
        result = myfunc("`varlist'")
        st_local("n" result)          /* <- mistake here */
        result = J(0,0,"")
    end
    ...
end
```

In the above example, we omitted the comma between `"n"` and `result`. We also used multiline `mata` without the colon. Therefore, the incorrect line will be tolerated by Mata, which will merrily continue executing our program until the `end` statement, at which point Mata will return control to Stata and not tell Stata that anything went wrong! This could have serious consequences, all of which could be avoided by substituting multiline `mata` with the colon.

## Also see

[\[M-3\] intro](#) — Commands for controlling Mata

# Title

[M-3] **mata clear** — Clear Mata's memory

## Syntax

```
: mata clear
```

This command is for use in Mata mode following Mata's colon prompt. To use this command from Stata's dot prompt, type

```
. mata: mata clear
```

## Description

`mata clear` clears Mata's memory, in effect resetting Mata. All functions, matrices, etc., are freed.

## Remarks

Stata can call Mata which can call Stata, which can call Mata, etc. In such cases, `mata clear` releases only resources that are not in use by prior invocations of Mata.

Stata's `clear all` command (see [\[D\] clear](#)) performs a `mata clear`, among other things.

See [\[M-3\] mata drop](#) for clearing individual matrices or functions.

## Also see

[\[M-3\] mata drop](#) — Drop matrix or function

[\[M-3\] intro](#) — Commands for controlling Mata



# Title

[M-3] **mata describe** — Describe contents of Mata’s memory

## Syntax

```
: mata describe [namelist] [, all]  
  
: mata describe using libname
```

where *namelist* is as defined in [M-3] **namelists**. If *namelist* is not specified, “\* \*()” is assumed.  
This command is for use in Mata mode following Mata’s colon prompt. To use this command from Stata’s dot prompt, type

```
. mata: mata describe ...
```

## Description

`mata describe` lists the names of the matrices and functions in memory, including the amount of memory consumed by each.  
`mata describe` using *libname* describes the contents of the specified .mlib library; see [M-3] **mata mlib**.

## Option

`all` specifies that automatically loaded library functions that happen to be in memory are to be included in the output.

## Remarks

`mata describe` is often issued without arguments, and then everything in memory is described:

```
: mata describe
```

# bytes	type	name and extent
50	real matrix	foo()
1,600	real matrix	X[10,20]
8	real scalar	x

`mata describe` using *libname* lists the functions stored in a .mlib library:

```
: mata describe using lmatbase
```

# bytes	type	name and extent
984	auto numeric vector	Corr()
340	auto real matrix	Hilbert()
(output omitted)		
528	auto transmorphic colvector	vech()

## Diagnostics

The reported memory usage does not include overhead, which usually amounts to 64 bytes, but can be less (as small as zero for recently used scalars).

The reported memory usage in the case of pointer matrices reflects the memory used to store the matrix itself and does not include memory consumed by siblings.

## Also see

[\[M-5\] `sizeof\(\)`](#) — Number of bytes consumed by object

[\[M-3\] `intro`](#) — Commands for controlling Mata

# Title

[M-3] **mata drop** — Drop matrix or function

## Syntax

```
: mata drop namelist
```

where *namelist* is as defined in [\[M-3\] \*\*namelists\*\*](#).

This command is for use in Mata mode following Mata's colon prompt. To use this command from Stata's dot prompt, type

```
. mata: mata drop ...
```

## Description

**mata drop** clears from memory the specified matrices and functions.

## Remarks

Use **mata describe** (see [\[M-3\] \*\*mata describe\*\*](#)) to determine what is in memory. Use **mata clear** (see [\[M-3\] \*\*mata clear\*\*](#)) to drop all matrices and functions, or use Stata's **clear mata** command (see [\[D\] \*\*clear\*\*](#)).

To drop a matrix named A, type

```
: mata drop A
```

To drop a function named `foo()`, type

```
: mata drop foo()
```

To drop a matrix named A and a function named `foo()`, type

```
: mata drop A foo()
```

## Also see

[\[M-3\] \*\*mata clear\*\*](#) — Clear Mata's memory

[\[M-3\] \*\*intro\*\*](#) — Commands for controlling Mata

# Title

[M-3] **mata help** — Obtain online help

## Syntax

```
: mata help ...
```

```
: help ...
```

help need not be preceded by mata.

## Description

mata help is Stata's help command. Thus you do not have to exit Mata to use help.

## Remarks

See [M-1] [help](#).

You need not type the Mata prefix:

```
: mata help mata
(output appears in Stata's Viewer)
: help mata
(same result)
```

help can be used to obtain help for Mata or Stata:

```
: help mata missing()
: help missing()
```

## Also see

[M-1] [help](#) — Obtaining online help

[R] [help](#) — Display online help

[M-3] [intro](#) — Commands for controlling Mata

# Title

[M-3] **mata matsave** — Save and restore matrices

## Syntax

```
: mata matsave filename namelist [ , replace ]
```

```
: mata matuse filename [ , replace ]
```

```
: mata matdescribe filename
```

where *namelist* is a list of matrix names as defined in [M-3] **namelists**.

If *filename* is specified without a suffix, `.mmat` is assumed.

These commands are for use in Mata mode following Mata's colon prompt. To use these commands from Stata's dot prompt, type

```
. mata: mata matsave ...
```

## Description

`mata matsave` saves the specified global matrices in *filename*.

`mata matuse` loads the matrices stored in *filename*.

`mata matdescribe` describes the contents of *filename*.

## Option for mata matsave

`replace` specifies that *filename* may be replaced if it already exists.

## Option for mata matuse

`replace` specifies that any matrices in memory with the same name as those stored in *filename* can be replaced.

## Remarks

These commands are for interactive use; they are not for use inside programs. See [M-5] **fopen()** for Mata's programming functions for reading and writing files. In the programming environment, if you have a matrix *X* and want to write it to file `mymatrix.myfile`, you code

```
fh = fopen("mymatrix.myfile", "w")
fputmatrix(fh, X)
fclose(fh)
```

Later, you can read it back by coding

```
fh = fopen("mymatrix.myfile", "r")
X = fgetmatrix(fh)
fclose(fh)
```

`mata matsave`, `mata matuse`, and `mata matdescribe` are for use outside programs, when you are working interactively. You can save your global matrices

```
: mata matsave mywork *
(saving A, X, Z, beta)
file mywork.mmat saved
```

and then later get them back:

```
: mata matuse mywork
(loading A, X, Z, beta)
```

`mata matdescribe` will tell you the contents of a file:

```
: mata matdescribe mywork
file mywork.mmat saved on 4 Apr 2009 08:46:39 contains
X, X, Z, beta
```

## Diagnostics

`mata matsave` saves the contents of view matrices. Thus when they are restored by `mata matuse`, the contents will be correct regardless of the data Stata has loaded in memory.

## Also see

[\[M-3\] intro](#) — Commands for controlling Mata

# Title

## Syntax

```
: mata memory
```

This command is for use in Mata mode following Mata’s colon prompt. To use this command from Stata’s dot prompt, type

```
. mata: mata memory
```

## Description

`mata memory` provides a summary of Mata’s current memory utilization.

## Remarks

: mata memory	#	bytes
autoloaded functions	15	5,514
ado functions	0	0
defined functions	0	0
matrices & scalars	14	8,256
overhead		1,972
total	29	15,742

Functions are divided into three types:

1. *Autoloaded functions*, which refer to library functions currently loaded (and which will be automatically unloaded) from `.mlib` library files and `.mo` object files.
2. *Ado-functions*, which refer to functions currently loaded whose source appears in ado-files themselves. These functions will be cleared when the ado-file is automatically cleared.
3. *Defined functions*, which refer to functions that have been defined either interactively or from do-files and which will be cleared only when a `mata clear`, `mata drop`, `Stata clear mata`, or `Stata clear` all command is executed; see [\[M-3\] mata clear](#), [\[M-3\] mata drop](#), or [\[D\] clear](#).

## Also see

- [\[M-1\] limits](#) — Limits and memory utilization
- [\[M-3\] mata clear](#) — Clear Mata’s memory
- [\[M-3\] intro](#) — Commands for controlling Mata

## Syntax

```
: mata mlib create libname      [ , dir(path) replace size(#) ]  
  
: mata mlib add libname fcnlist() [ , dir(path) complete ]  
  
: mata mlib index  
  
: mata mlib query
```

where *fcnlist*() is a *namelist* containing only function names, such as

*fcnlist*() examples

---

```
myfunc()  
myfunc() myotherfunc() foo()  
f*() g*()  
*()
```

---

see [\[M-3\] namelists](#)

and where *libname* is the name of a library. You must start *libname* with the letter l and do not add the .mlib suffix as it will be added for you. Examples of *libnames* include

<i>libname</i>	Corresponding filename
lmath	lmath.mlib
lmoremath	lmoremath.mlib
lnjc	lnjc.mlib

Also *libnames* that begin with the letters lmata, such as lmatabase, are reserved for the names of official libraries.

This command is for use in Mata mode following Mata’s colon prompt. To use this command from Stata’s dot prompt, type

```
. mata: mata mlib ...
```

## Description

`mata mlib` creates, adds to, and causes Mata to index .mlib files, which are libraries containing the object-code functions.

`mata mlib create` creates a new, empty library.



`mata mlib add` adds new members to a library.

`mata mlib index` causes Mata to build a new list of libraries to be searched.

`mata mlib query` lists the libraries to be searched.

A library may contain up to 1,024 functions by default.

## Options

`dir(path)` specifies the directory (folder) into which the file should be written. `dir(.)` is the default, meaning that if `dir()` is not specified, the file is written into the current (working) directory. *path* may be a directory name or may be the `sysdir` shorthand STATA, UPDATES, BASE, SITE, PLUS, PERSONAL, or OLDPLACE; see [P] [sysdir](#). `dir(PERSONAL)` is recommended.

`complete` is for use when saving class definitions. It specifies that the definition be saved only if it is complete; otherwise, an error message is to be issued. See [M-2] [class](#).

`replace` specifies that the file may be replaced if it already exists.

`size(#)`, used with `mlib create`, specifies the maximum number of members the newly created library will be able to contain,  $2 \leq \# \leq 2048$ . The default is `size(1024)`.

## Remarks

Remarks are presented under the following headings:

*Background*  
*Outline of the procedure for dealing with libraries*  
*Creating a .mlib library*  
*Adding members to a .mlib library*  
*Listing the contents of a library*  
*Making it so Mata knows to search your libraries*  
*Advice on organizing your source code*

Also see [M-1] [how](#) for an explanation of object code.

## Background

.mlib files contain the object code for one or more functions. Functions which happen to be stored in libraries are called library functions, and Mata's library functions are also stored in .mlib libraries. You can create your own libraries, too.

Mata provides two ways to store object code:

1. In a .mo file, which contains the code for one function
2. In a .mlib library file, which may contain the code for up to 500 functions

.mo files are easier to use and work just as well as .mlib libraries; see [M-3] [mata mosave](#). .mlib libraries, however, are easier to distribute to others when you have many functions, because they are combined into one file.

## Outline of the procedure for dealing with libraries

Working with libraries is easy:

1. First, choose a name for your library. We will choose the name `lpersonal`.
2. Next, create an empty library by using the `mata mlib create` command.
3. After that, you can add new members to the library at any time, using `mata mlib add`.

`.mlib` libraries contain object code, not the original source code, so you need to keep track of the source code yourself. Also, if you want to update the object code in a function stored in a library, you must re-create the entire library; there is no way to replace or delete a member once it is added.

We begin by showing you the mechanical steps, and then we will tell you how we manage libraries and source code.

### Creating a `.mlib` library

If you have not read [M-3] [mata mosave](#), please do so.

To create a new, empty library named `lpersonal.mlib` in the current directory, type

```
: mata mlib create lpersonal
file lpersonal.mlib created
```

If `lpersonal.mlib` already exists and you want to replace it, either erase the existing file first or type

```
: mata mlib create lpersonal, replace
file lpersonal.mlib created
```

To create a new, empty library named `lpersonal.mlib` in your `PERSONAL` (see [P] [sysdir](#)) directory, type

```
: mata mlib create lpersonal, dir(PERSONAL)
file c:\ado\personal\lpersonal.mlib created
```

or

```
: mata mlib create lpersonal, dir(PERSONAL) replace
file c:\ado\personal\lpersonal.mlib created
```

### Adding members to a `.mlib` library

Once a library exists, whether you have just created it and it is empty, or it already existed and contains some functions, you can add new functions to it by typing

```
: mata mlib add libname fcname()
```

So, if we wanted to add function `example()` to library `lpersonal.mlib`, we would type

```
: mata mlib add lpersonal example()
(1 function added)
```

In doing this, we do not have to say where `lpersonal.mlib` is stored; Mata searches for it along the `ado-path`.

Before you can add `example()` to the library, however, you must compile it:

```
: function example(...)
> {
>     ...
> }

: mata mlib add lpersonal example()
(1 function added)
```

You can add many functions to a library in one command:

```
: mata mlib add lpersonal example2() example3()
(2 functions added)
```

You can add all the functions currently in memory by typing

```
: mata mlib add lanother *()
(3 functions added)
```

In the above example, we added to `lanother.mlib` because we had already added `example()`, `example2()`, and `example3()` to `lpersonal.mlib` and trying to add them again would result in an error. (Before adding `*`, we could verify that we are adding what we want to add by typing `mata describe *`; see [\[M-3\] mata describe](#).)

## Listing the contents of a library

Once a library exists, you can list its contents (the names of the functions it contains) by typing

```
: mata describe using libname
```

Here we would type

```
: mata describe using lpersonal
(library contains 3 members)
```

# bytes	type	name and extent
32	auto transmorphic matrix	example()
32	auto transmorphic matrix	example2()
32	auto transmorphic matrix	example3()

`mata describe` usually lists the contents of memory, but `mata describe using` lists the contents of a library.

## Making it so Mata knows to search your libraries

Mata automatically finds the `.mlib` libraries on your computer. It does this when Mata is invoked for the first time during a session. Thus everything is automatic except that Mata will know nothing about any new libraries created during the Stata session, so after creating a new library, you must tell Mata about it. You do this by asking Mata to rebuild its library index:

```
: mata mlib index
```

You do not specify the name of your new library. That name does not matter because Mata rebuilds its entire library index.

You can issue the `mata mlib index` command right after creating the new library

```
: mata mlib create lpersonal, dir(PERSONAL)
: mata mlib index
```

or after you have created and added to the library:

```
: mata mlib create lpersonal, dir(PERSONAL)
: mata mlib add lpersonal *()
: mata mlib index
```

It does not matter. Mata does not need to rebuild its index after a known library is updated; Mata needs to be told to rebuild only when a new library is added during the session.

## Advice on organizing your source code

Say you wish to create and maintain `lpersonal.mlib`. Our preferred way is to use a do-file:

```
----- begin lpersonal.do -----
mata:
mata clear
function definitions appear here
mata mlib create lpersonal, dir(PERSONAL) replace
mata mlib add lpersonal *()
mata mlib index
end
----- end lpersonal.do -----
```

This way, all we have to do to create or re-create the library is enter Stata, change to the directory containing our source code, and type

```
. do lpersonal
```

For large libraries, we like to put the source code for different parts in different files:

```
----- begin lpersonal.do -----
mata: mata clear
do function1.mata
do function2.mata
...
mata:
mata mlib create lpersonal, dir(PERSONAL) replace
mata mlib add lpersonal *()
mata mlib index
end
----- end lpersonal.do -----
```

The function files contain the source code, which might include one function, or it might include more than one function if the primary function had subroutines:

---

```
begin function1.mata
mata:
  function definitions appear here
end
```

---

```
end function1.mata
```

---

We name our component files ending in `.mata`, but they are still just do-files.

## Also see

[M-3] [mata mosave](#) — Save function's compiled code in object file

[M-3] [intro](#) — Commands for controlling Mata

# Title

**[M-3] mata mosave** — Save function's compiled code in object file

## Syntax

```
: mata mosave fcname() [ , dir(path) complete replace ]
```

This command is for use in Mata mode following Mata's colon prompt. To use this command from Stata's dot prompt, type

```
. mata: mata mosave ...
```

## Description

`mata mosave` saves the object code for the specified function in the file `fcname.mo`.

## Options

`dir(path)` specifies the directory (folder) into which the file should be written. `dir(.)` is the default, meaning that if `dir()` is not specified, the file is written into the current (working) directory. *path* may be a directory name or may be the `sysdir` shorthand STATA, UPDATES, BASE, SITE, PLUS, PERSONAL, or OLDPLACE; see [\[P\] sysdir](#). `dir(PERSONAL)` is recommended.

`complete` is for use when saving class definitions. It specifies that the definition be saved only if it is complete; otherwise, an error message is to be issued. See [\[M-2\] class](#).

`replace` specifies that the file may be replaced if it already exists.

## Remarks

See [\[M-1\] how](#) for an explanation of object code.

Remarks are presented under the following headings:

*Example of use*

*Where to store .mo files*

*Use of .mo files versus .mlib files*

## Example of use

.mo files contain the object code for one function. If you store a function's object code in a .mo file, then in future Mata sessions, you can use the function without recompiling the source. The function will appear to become a part of Mata just as all the other functions documented in this manual are. The function can be used because the object code will be automatically found and loaded when needed.

For example,

```
: function add(a,b) return(a+b)
: add(1,2)
3
```

```

: mata mosave add()
(file add.mo created)
: mata clear
: add(1,2)
3

```

In the example above, function `add()` was saved in file `add.mo` stored in the current directory. After clearing Mata, we could still use the function because Mata found the stored object code.

## Where to store .mo files

Mata could find `add()` because file `add.mo` was in the current directory, and our `ado-path` included `..`:

```

. adopath
[1] (UPDATES) "C:\Program Files\Stata12\ado\updates\"
[2] (BASE)    "C:\Program Files\Stata12\ado\base\"
[3] (SITE)    "C:\Program Files\Stata12\ado\site\"
[4]          "."
[5] (PERSONAL) "C:\ado\personal\"
[6] (PLUS)    "C:\ado\plus\"
[7] (OLDPLACE) "C:\ado\"

```

If later we were to change our current directory

```

. cd ..\otherdir

```

Mata would no longer be able to find the file `add.mo`. Thus the best place to store your personal `.mo` files is in your `PERSONAL` directory. Thus rather than typing

```

: mata mosave example()

```

we would have been better off typing

```

: mata mosave example(), dir(PERSONAL)

```

## Use of .mo files versus .mlib files

Use of `.mo` files is heartily recommended. The alternative for saving compiled object code are `.mlib` libraries; see [\[M-3\] mata mlib](#) and [\[M-1\] ado](#).

Libraries are useful when you have many functions and want to tie them together into one file, especially if you want to share those functions with others, because then you have only one file to distribute. The disadvantage of libraries is that you must rebuild them whenever you wish to remove or change the code of one. If you have only a few object files, or if you have many but sharing is not an issue, `.mo` libraries are easier to manage.

## Also see

[\[M-3\] mata mlib](#) — Create function library

[\[M-3\] intro](#) — Commands for controlling Mata

# Title

[M-3] **mata rename** — Rename matrix or function

## Syntax

`: mata rename oldmatrixname newmatrixname`

`: mata rename oldfcnname() newfcnname()`

This command is for use in Mata mode following Mata’s colon prompt. To use this command from Stata’s dot prompt, type

`. mata: mata rename ...`

## Description

`mata rename` changes the names of functions and global matrices.

## Also see

[\[M-3\] intro](#) — Commands for controlling Mata



## Syntax

```

: mata query

: mata set matakache      # [ , permanently]
: mata set matalnum       { off | on }
: mata set mataoptimize   { on | off }
: mata set matafavor      { space | speed } [ , permanently]
: mata set matastrict     { off | on } [ , permanently]
: mata set matalibs       "libname;libname;..."
: mata set matamofirst     { off | on } [ , permanently]
    
```

These commands are for use in Mata mode following Mata's colon prompt. To use these commands from Stata's dot prompt, type

```

. mata: mata query
. mata: mata set ...
    
```

## Description

`mata query` shows the values of Mata's system parameters.

`mata set` sets the value of the system parameters:

`mata set matakache` specifies the maximum amount of memory, in kilobytes, that may be consumed before Mata starts looking to drop autoloaded functions that are not currently being used. The default value is 400, meaning 400 kilobytes. This parameter affects the efficiency with which Stata runs. Larger values cannot hurt, but once `matakache` is large enough, larger values will not improve performance.

`mata set matalnum` turns program line-number tracing on or off. The default setting is off. This setting modifies how programs are compiled. Programs compiled when `matalnum` is turned on include code so that, if an error occurs during execution of the program, the line number is also reported. Turning `matalnum` on prevents Mata from being able to optimize programs, so they will run more slowly. Except when debugging, the recommended setting for this is off.

`mata set mataoptimize` turns compile-time code optimization on or off. The default setting is on. Programs compiled when `mataoptimize` is switched off will run more slowly and, sometimes, much more slowly. The only reason to set `mataoptimize` off is if a bug in the optimizer is suspected.

`mata set matafavor` specifies whether, when executing code, Mata should favor conserving memory (space) or running quickly (speed). The default setting is `space`. Switching to `speed` will make Mata, in a few instances, run a little quicker but consume more memory. Also see [M-5] [favorspeed\(\)](#).

`mata set matastrict` sets whether declarations can be omitted inside the body of a program. The default is `off`. If `matastrict` is switched on, compiling programs that omit the declarations will result in a compile-time error; see [M-2] [declarations](#). `matastrict` acts unexpectedly but pleasingly when set/reset inside `ado`-files; see [M-1] [ado](#).

`mata set matalibs` sets the names and order of the `.mlib` libraries to be searched; see [M-1] [how](#). `matalibs` usually is set to `"lmatabase;lmataopt;lmataado"`. However it is set, it is probably set correctly, because Mata automatically searches for libraries the first time it is invoked in a Stata session. If, during a session, you erase or copy new libraries along the `ado-path`, the best way to reset `matalibs` is with the `mata mlib index` command; see [M-3] [mata mlib](#). The only reason to set `matalibs` by hand is to modify the order in which libraries are searched.

`mata set matamofirst` states whether `.mo` files or `.mlib` libraries are searched first. The default is `off`, meaning libraries are searched first.

## Option

`permanently` specifies that, in addition to making the change right now, the setting be remembered and become the default setting when you invoke Stata in the future.

## Remarks

Remarks are presented under the following headings:

*Relationship between Mata's `mata set` and Stata's `set` commands  
c() values*

### Relationship between Mata's `mata set` and Stata's `set` commands

The command

```
: mata set ...
```

issued from Mata's colon prompt and the command

```
. set ...
```

issued from Stata's dot prompt are the same command, so you may set Mata's (or even Stata's) system parameters either way.

The command

```
: mata query
```

issued from Mata's colon prompt and the command

```
. query mata
```

issued from Stata's dot prompt are also the same command.

## c() values

The following concerns Stata more than Mata.

Stata's c-class, `c()`, contains the values of system parameters and settings along with certain other constants. `c()` values may be referred to in Stata, either via macro substitution (`'c(current_date)'`, for example) or in expressions (in which case the macro quoting characters may be omitted). Stata's `c()` is also available in Mata via Mata's `c()` function; see [\[M-5\] c\(\)](#).

Most everything set by `set` is available via `c()`, including Mata's set parameters:

`c(matacache)` returns a numeric scalar equal to the cache size.

`c(matalnum)` returns a string equal to "on" or "off".

`c(mataoptimize)` returns a string equal to "on" or "off".

`c(matafavor)` returns a string equal to "space" or "speed".

`c(matastrict)` returns a string equal to "on" or "off".

`c(matalibs)` returns a string of library names separated by semicolons.

`c(matamofirst)` returns a string equal to "on" or "off".

The above is in StataSpeak. Rather than referring to `c(matacache)`, we would refer to `c("matacache")` if we were using Mata's function. The real use of these values, however, is in Stata.

## Also see

[\[R\] query](#) — Display system parameters

[\[R\] set](#) — Overview of system parameters

[\[P\] creturn](#) — Return c-class values

[\[M-3\] intro](#) — Commands for controlling Mata

# Title

[M-3] **mata stata** — Execute Stata command

## Syntax

```
: mata stata stata_command
```

This command is for use in Mata mode following Mata's colon prompt.

## Description

`mata stata stata_command` passes *stata\_command* to Stata for execution.

## Remarks

`mata stata` is a convenience tool to keep you from having to exit Mata:

```
: st_view(V=., 1\5, ("mpg", "weight"))
no variables defined
      st_view(): 3598  Stata returned error
      <istmt>:    -  function returned error
r(3598);
: mata stata sysuse auto
(1978 Automobile Data)
: st_view(V=., 1\5, ("mpg", "weight"))
```

`mata stata` is for interactive use. If you wish to execute a Stata command from a function, see [\[M-5\] `stata\(\)`](#).

## Also see

[\[M-5\] `stata\(\)`](#) — Execute Stata command

[\[M-3\] `intro`](#) — Commands for controlling Mata

# Title

[M-3] **mata which** — Identify function

## Syntax

```
: mata which fcname()
```

This command is for use in Mata mode following Mata's colon prompt. To use this command from Stata's dot prompt, type

```
. mata: mata which ...
```

## Description

`mata which fcname` looks for *fcname*() and reports whether it is built in, stored in a `.mlib` library, or stored in a `.mo` file.

## Remarks

`mata which fcname()` looks for *fcname*() and reports where it is found:

```
: mata which I()
I(): built-in
: mata which assert()
assert(): lmatbase
: mata which myfcn()
userfunction(): .\myfcn.mo
: mata which nosuchfunction()
function nosuchfunction() not found
r(111);
```

Function `I()` is built in; it was written in C and is a part of Mata itself.

Function `assert()` is a library function and, as a matter of fact, its executable object code is located in the official function library `lmatbase.mlib`.

Function `myfcn()` exists and has its executable object code stored in file `myfcn.mo`, located in the current directory.

Function `nosuchfunction()` does not exist.

Going back to `mata which assert()`, which was found in `lmatbase.mlib`, if you wanted to know where `lmatbase.mlib` was stored, you could type `findfile lmatbase.mlib` at the Stata prompt; see [\[P\] findfile](#).

## Also see

[\[M-3\] intro](#) — Commands for controlling Mata

## Syntax

Many `mata` commands allow or require a *namelist*, such as

```
: mata describe [ namelist ] [ , all ]
```

A *namelist* is defined as a list of matrix and/or function names, such as

```
alpha beta foo()
```

The above *namelist* refers to the matrices `alpha` and `beta` along with the function named `foo()`.

Function names always end in `()`, hence

<code>alpha</code>	refers to the matrix named <code>alpha</code>
<code>alpha()</code>	refers to the function of the same name

Names may also be specified using the `*` and `?` wildcard characters:

<code>*</code>	means zero or more characters go here
<code>?</code>	means exactly one character goes here

hence,

<code>*</code>	means all matrices
<code>*()</code>	means all functions
<code>* *</code>	means all matrices and all functions
<code>s*</code>	means all matrices that start with <i>s</i>
<code>s*()</code>	means all functions that start with <i>s</i>
<code>*e</code>	means all matrices that end with <i>e</i>
<code>*e()</code>	means all functions that end with <i>e</i>
<code>s*e</code>	means all matrices that start with <i>s</i> and end with <i>e</i>
<code>s*e()</code>	means all functions that start with <i>s</i> and end with <i>e</i>
<code>s?e</code>	means all matrices that start with <i>s</i> and end with <i>e</i> and have one character in between
<code>s?e()</code>	means all functions that start with <i>s</i> and end with <i>e</i> and have one character in between

## Description

*Namelists* appear in syntax diagrams.

---

## Remarks

Some *namelists* allow only matrices, and some allow only functions. Even when only functions are allowed, you must include the `()` suffix.

## Also see

[\[M-3\] intro](#) — Commands for controlling Mata





## **[M-4] Index and guide to functions**



# Title

[M-4] intro — Index and guide to functions

## Contents

[M-4] Entry	Description
Mathematical	
<a href="#">matrix</a>	Matrix functions
<a href="#">solvers</a>	Matrix solvers and inverters
<a href="#">scalar</a>	Scalar functions
<a href="#">statistical</a>	Statistical functions
<a href="#">mathematical</a>	Other important functions
Utility & manipulation	
<a href="#">standard</a>	Standard matrices
<a href="#">utility</a>	Matrix utility functions
<a href="#">manipulation</a>	Matrix manipulation functions
Stata interface	
<a href="#">stata</a>	Stata interface functions
String, I/O, & programming	
<a href="#">string</a>	String manipulation functions
<a href="#">io</a>	I/O functions
<a href="#">programming</a>	Programming functions

## Description

The entries in this section provides an index to the functions, grouped according to purpose.

## Remarks

The next section, section [\[M-5\]](#), presents all the Mata functions, in alphabetical order.

Also see

[M-0] [intro](#) — Introduction to the Mata manual

Contents

[M-5] Manual entry	Function	Purpose
Console output		
<b>printf()</b>	printf() sprintf()	display display into string
<b>errprintf()</b>	errprintf()	display error message
<b>display()</b>	display()	display text interpreting SMCL
<b>displayas()</b>	displayas()	set whether output is displayed
<b>displayflush()</b>	displayflush()	flush terminal output buffer
<b>liststruct()</b>	liststruct()	list structure's contents
<b>more()</b>	more() setmore() setmoreonexit()	create --more-- condition query or set more on or off set more on or off on exit
File directories		
<b>direxists()</b>	direxists()	whether directory exists
<b>dir()</b>	dir()	file list
<b>chdir()</b>	pwd() chdir() mkdir() rmdir()	obtain current working directory change current working directory make new directory remove directory
File management		
<b>findfile()</b>	findfile()	find file
<b>fileexists()</b>	fileexists()	whether file exists
<b>cat()</b>	cat()	read file into string matrix
<b>unlink()</b>	unlink()	erase file
<b>adosubdir()</b>	adosubdir()	obtain ado-subdirectory for file

File I/O

---

<b>fopen()</b>	<code>fopen()</code>	open file
	<code>fclose()</code>	close file
	<code>fget()</code>	read line of ASCII file
	<code>fgetnl()</code>	same, but include newline character
	<code>fread()</code>	read <i>k</i> bytes of binary file
	<code>fput()</code>	write line into ASCII file
	<code>fwrite()</code>	write <i>k</i> bytes into binary file
	<code>fgetmatrix()</code>	read matrix
	<code>fputmatrix()</code>	write matrix
	<code>fstatus()</code>	status of last I/O command
	<code>ftell()</code>	report location in file
	<code>fseek()</code>	seek to location in file
	<code>ftruncate()</code>	truncate file at current position
<b>ferrortext()</b>	<code>ferrortext()</code>	error text of file error code
	<code>freturncode()</code>	return code of file error code
<b>bufio()</b>	<code>bufio()</code>	initialize buffer
	<code>bufbyteorder()</code>	reset (specify) byte order
	<code>bufmissingvalue()</code>	reset (specify) missing-value encoding
	<code>bufput()</code>	copy into buffer
	<code>bufget()</code>	copy from buffer
	<code>fbufput()</code>	copy into and write buffer
	<code>fbufget()</code>	read and copy from buffer
	<code>bufbfmtlen()</code>	utility routine
	<code>bufbfmtisnum()</code>	utility routine

Filename & path manipulation

---

<b>pathjoin()</b>	<code>pathjoin()</code>	join paths
	<code>pathsplit()</code>	split paths
	<code>pathbasename()</code>	path basename
	<code>pathsuffix()</code>	file suffix
	<code>pathrmsuffix()</code>	remove file suffix
	<code>pathisurl()</code>	whether path is URL
	<code>pathisabs()</code>	whether path is absolute
	<code>pathasciisuffix()</code>	whether file is ASCII
	<code>pathstatastatusuffix()</code>	whether file is Stata
	<code>pathlist()</code>	process path list
	<code>pathsubsysdir()</code>	substitute for system directories
	<code>pathsearchlist()</code>	path list to search for file

---

## Description

The above functions have to do with

1. Displaying output at the terminal.
2. Reading and writing data in a file.

## Remarks

To display the contents of a scalar, vector, or matrix, it is sufficient merely to code the identity of the scalar, vector, or matrix:

```
: x
      1      2      3      4
1  .1369840784  .643220668  .5578016951  .6047949435
```

You can follow this approach even in programs:

```
function example()
{
    ...
    "i am about to calculate the result"
    ...
    "the result is"
    b
}
```

On the other hand, `display()` and `printf()` (see [M-5] [display\(\)](#) and [M-5] [printf\(\)](#)) will allow you to exercise more control over how the output looks.

Changing the subject: you will find that many I/O functions come in two varieties: with and without an underscore in front of the name, such as `_fopen()` and `fopen()`. As always, functions that begin with an underscore are generally silent about their work and return flags indicating their success or failure. Functions that omit the underscore abort and issue the appropriate error message when things go wrong.

## Reference

Gould, W. W. 2009. [Mata Matters: File processing](#). *Stata Journal* 9: 599–620.

## Also see

[M-4] [intro](#) — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
Transposition		
<b>transposeonly()</b>	transposeonly()	transposition without conjugation
<b>_transpose()</b>	_transpose()	transposition in place
Diagonals		
<b>diag()</b>	diag()	create diagonal matrix from vector
<b>_diag()</b>	_diag()	replace diagonal of matrix
<b>diagonal()</b>	diagonal()	extract diagonal of matrix into vector
Triangular & symmetric		
<b>lowertriangle()</b>	lowertriangle() uppertriangle()	extract lower triangle extract upper triangle
<b>sublowertriangle()</b>	sublowertriangle()	generalized lowertriangle()
<b>makesymmetric()</b>	makesymmetric()	make matrix symmetric (Hermitian)
Sorting		
<b>sort()</b>	sort() jumble() order() unorder() _collate()	sort rows of matrix randomize order of rows of matrix permutation vector for ordered rows permutation vector for randomized rows order matrix on permutation vector
<b>uniqrows()</b>	uniqrows()	sorted, unique rows



## Editing

<code>_fillmissing()</code>	<code>_fillmissing()</code>	change matrix to contain missing values
<code>editmissing()</code>	<code>editmissing()</code>	replace missing values in matrix
<code>editvalue()</code>	<code>editvalue()</code>	replace values in matrix
<code>edittozero()</code>	<code>edittozero()</code> <code>edittozerotol()</code>	edit matrix for roundoff error (zeros) same, absolute tolerance
<code>edittoint()</code>	<code>edittoint()</code> <code>edittointtol()</code>	edit matrix for roundoff error (integers) same, absolute tolerance

## Permutation vectors

<code>invorder()</code>	<code>invorder()</code> <code>revorder()</code>	inverse of permutation vector reverse of permutation vector
-------------------------	--	--

## Matrices into vectors &amp; vice versa

<code>vec()</code>	<code>vec()</code> <code>vech()</code> <code>invvech()</code>	convert matrix into column vector convert symmetric matrix into column vector convert column vector into symmetric matrix
<code>rowshape()</code>	<code>rowshape()</code> <code>colshape()</code>	reshape matrix to have $r$ rows reshape matrix to have $c$ columns

## Associative arrays

<code>asarray()</code>	<code>asarray()</code> <code>asarray_*</code>	store or retrieve element in array utility routines
------------------------	--	--

## Description

The above functions manipulate matrices, such as extracting the diagonal and sorting.

## Remarks

There is a thin line between manipulation and utility; also see

[M-4] [utility](#)      Matrix utility functions

## Also see

[M-4] [intro](#) — Index and guide to functions

Contents

[M-5]

Manual entry	Function	Purpose
Basics (also see <a href="#">[M-4] scalar</a> )		
<a href="#">sum()</a>	<code>rowsum()</code>	sum of each row
	<code>colsum()</code>	sum of each column
	<code>sum()</code>	overall sum
	<code>quadrowsum()</code>	quad-precision sum of each row
	<code>quadcolsum()</code>	quad-precision sum of each column
	<code>quadsum()</code>	quad-precision overall sum
<a href="#">runningsum()</a>	<code>runningsum()</code>	running sum of vector
	<code>quadrunningsum()</code>	quad-precision <code>runningsum()</code>
<a href="#">minmax()</a>	<code>rowmin()</code>	minimum, by row
	<code>colmin()</code>	minimum, by column
	<code>min()</code>	minimum, overall
	<code>rowmax()</code>	maximum, by row
	<code>colmax()</code>	maximum, by column
	<code>max()</code>	maximum, overall
	<code>rowminmax()</code>	minimum and maximum, by row
	<code>colminmax()</code>	minimum and maximum, by column
	<code>minmax()</code>	minimum and maximum, overall
	<code>rowmaxabs()</code>	<code>rowmax(abs())</code>
	<code>colmaxabs()</code>	<code>colmax(abs())</code>
<a href="#">deriv()</a>	<code>deriv()</code>	numerical derivatives
	<code>deriv_init()</code>	begin derivatives
	<code>deriv_init_*</code>	set details
	<code>deriv()</code>	compute derivatives
	<code>deriv_result_*</code>	access results
	<code>deriv_query()</code>	report settings
<a href="#">optimize()</a>	<code>optimize()</code>	function maximization and minimization
	<code>optimize_init()</code>	begin optimization
	<code>optimize_init_*</code>	set details
	<code>optimize()</code>	perform optimization
	<code>optimize_result_*</code>	access results
	<code>optimize_query()</code>	report settings

---

Basics, *continued*

---

<b>moptimize()</b>	<code>moptimize()</code>	function optimization
	<code>moptimize_ado_cleanup()</code>	perform cleanup after ado
	<code>moptimize_evaluate()</code>	evaluate function at initial values
	<code>moptimize_init()</code>	begin setup of optimization problem
	<code>moptimize_init_*</code>	set details
	<code>moptimize_result_*</code>	access <code>moptimize()</code> results
	<code>moptimize_query()</code>	report settings
	<code>moptimize_util_*</code>	utility functions for writing evaluators and processing results

---

Fourier transform

---

<b>fft()</b>	<code>fft()</code>	fast Fourier transform
	<code>invfft()</code>	inverse fast Fourier transform
	<code>convolve()</code>	convolution
	<code>deconvolve()</code>	inverse of <code>convolve()</code>
	<code>Corr()</code>	correlation
	<code>ftperiodogram()</code>	power spectrum
	<code>ftpad()</code>	pad to power-of-2 length
	<code>ftwrap()</code>	convert to frequency-wraparound order
	<code>ftunwrap()</code>	convert from frequency-wraparound order
	<code>ftretime()</code>	change time scale of signal
	<code>ftfreqs()</code>	frequencies of transform

---

Cubic splines

---

<b>spline3()</b>	<code>spline3()</code>	fit cubic spline
	<code>spline3eval()</code>	evaluate cubic spline

---

Polynomials

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<b>polyeval()</b>	<code>polyeval()</code>	evaluate polynomial
	<code>polysolve()</code>	solve for polynomial
	<code>polytrim()</code>	trim polynomial
	<code>polyderiv()</code>	derivative of polynomial
	<code>polyinteg()</code>	integral of polynomial
	<code>polyadd()</code>	add polynomials
	<code>polymult()</code>	multiply polynomials
	<code>polydiv()</code>	divide polynomials
	<code>polyroots()</code>	find roots of polynomial

Number-theoretic point sets

<b>halton()</b>	halton()	generate a Halton or Hammersley set
	ghalton()	generate a generalized Halton sequence

Base conversion

<b>inbase()</b>	inbase()	convert to specified base
	frombase()	convert from specified base

## Description

The above functions are important mathematical functions that most people would not call either matrix functions or scalar functions, but that use matrices and scalars.

## Remarks

For other mathematical functions, see

[M-4] <b>matrix</b>	Matrix mathematical functions
[M-4] <b>scalar</b>	Scalar mathematical functions
[M-4] <b>statistical</b>	Statistical functions

## Also see

[M-4] **intro** — Index and guide to functions

Contents

[M-5] Manual entry Function		Purpose
Characteristics		
<code>trace()</code>	<code>trace()</code>	trace of matrix
<code>det()</code>	<code>det()</code>	determinant
	<code>dettriangular()</code>	determinant of triangular matrix
<code>norm()</code>	<code>norm()</code>	matrix and vector norms
<code>cond()</code>	<code>cond()</code>	matrix condition number
<code>rank()</code>	<code>rank()</code>	rank of matrix
Cholesky decomposition, solvers, & inverters		
<code>cholesky()</code>	<code>cholesky()</code>	Cholesky square-root decomposition $A = GG'$
<code>cholsolve()</code>	<code>cholsolve()</code>	solve $AX = B$ for $X$
<code>cholinv()</code>	<code>cholinv()</code>	inverse of pos. def. symmetric matrix
<code>invsym()</code>	<code>invsym()</code>	real symmetric matrix inversion
LU decomposition, solvers, & inverters		
<code>lud()</code>	<code>lud()</code>	LU decomposition $A = PLU$
<code>lusolve()</code>	<code>lusolve()</code>	solve $AX = B$ for $X$
<code>luinv()</code>	<code>luinv()</code>	inverse of square matrix

QR decomposition, solvers, & inverters

<b>qrd()</b>	<code>qrd()</code>	QR decomposition $A = QR$
	<code>qrdp()</code>	QR decomposition $A = QRP'$
	<code>hqrd()</code>	QR decomposition $A = f(H)R_1$
	<code>hqrdp()</code>	QR decomposition $A = f(H, \tau)R_1P'$
	<code>hqrmultq()</code>	return $QX$ or $Q'X$ , $Q = f(H, \tau)$
	<code>hqrmultq1t()</code>	return $Q'_1X$ , $Q_1 = f(H, \tau)$
	<code>hqrdaq()</code>	return $Q = f(H, \tau)$
	<code>hqrdaq1()</code>	return $Q_1 = f(H, \tau)$
	<code>hqrdr()</code>	return $R$
	<code>hqrdr1()</code>	return $R_1$
<b>qrsolve()</b>	<code>qrsolve()</code>	solve $AX = B$ for $X$
<b>qrinv()</b>	<code>qrinv()</code>	generalized inverse of matrix

Hessenberg decomposition & generalized Hessenberg decomposition

<b>hessenbergd()</b>	<code>hessenbergd()</code>	Hessenberg decomposition $T = Q'XQ$
<b>ghessenbergd()</b>	<code>ghessenbergd()</code>	gen. Hessenberg decomp. $T = Q'XQ$

Schur decomposition & generalized Schur decomposition

<b>schurd()</b>	<code>schurd()</code>	Schur decomposition $T = U'AV$ ; $R = U'BA$
	<code>schurdgroupby()</code>	Schur decomp. with grouping of results
<b>gschurd()</b>	<code>gschurd()</code>	gen. Schur decomposition $T = U'AV$ ; $R = U'BA$
	<code>gschurdgroupby()</code>	gen. Schur decomp. with grouping of results

Singular value decomposition, solvers, & inverters

<b>svd()</b>	<code>svd()</code>	singular value decomposition $A = UDV'$
	<code>svdsv()</code>	singular values $s$
<b>fullsvd()</b>	<code>fullsvd()</code>	singular value decomposition $A = USV'$
	<code>fullsdiag()</code>	convert $s$ to $S$
<b>svsolve()</b>	<code>svsolve()</code>	solve $AX = B$ for $X$
<b>pinv()</b>	<code>pinv()</code>	Moore–Penrose pseudoinverse

---

Triangular solvers

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<b>solverlower()</b>	<code>solverlower()</code>	solve $AX = B$ for $X$ , $A$ lower triangular
	<code>solverupper()</code>	solve $AX = B$ for $X$ , $A$ upper triangular

---

Eigensystems, powers, & transcendental

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<b>eigensystem()</b>	<code>eigensystem()</code>	eigenvectors and eigenvalues
	<code>eigenvalues()</code>	eigenvalues
	<code>lefteigensystem()</code>	left eigenvectors and eigenvalues
	<code>syemeigensystem()</code>	eigenvectors/eigenvalues of symmetric matrix
	<code>syemeigenvalues()</code>	eigenvalues of symmetric matrix
<b>eigensystemselect()</b>	<code>eigensystemselect*()</code>	selected eigenvectors/eigenvalues etc.
<b>geigensystem()</b>	<code>geigensystem()</code>	generalized eigenvectors/eigenvalues etc.
<b>matpowersym()</b>	<code>matpowersym()</code>	powers of symmetric matrix
<b>matexpsym()</b>	<code>matexpsym()</code>	exponentiation of symmetric matrix
	<code>matlogsym()</code>	logarithm of symmetric matrix

---

Equilibration

---

<b>_equilrc()</b>	<code>_equilrc()</code>	row/column equilibration
	<code>_equilr()</code>	row equilibration
	<code>_equilc()</code>	column equilibration
	<code>_perhapsequilrc()</code>	row/column equilibration if necessary
	<code>_perhapsequilr()</code>	row equilibration if necessary
	<code>_perhapsequiloc()</code>	column equilibration if necessary
	<code>rowscalefactors()</code>	row-scaling factors for equilibration
	<code>colscalefactors()</code>	column-scaling factors for equilibration

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LAPACK

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<b>lapack()</b>	<code>LA_*</code>	LAPACK linear-algebra functions
	<code>_flopint()</code>	convert matrix order from row major to column major
	<code>_flopout()</code>	convert matrix order from column major to row major

---

## Description

The above functions are what most people would call mathematical matrix functions.

## Remarks

For other mathematical functions, see

[M-4] <a href="#">scalar</a>	Scalar mathematical functions
[M-4] <a href="#">mathematical</a>	Important mathematical functions

## Also see

[M-4] [intro](#) — Index and guide to functions



Contents

[M-5] Manual entry	Function	Purpose
Argument and caller-preference processing		
<code>args()</code>	<code>args()</code>	number of arguments
<code>isfleeting()</code>	<code>isfleeting()</code>	whether argument is temporary
<code>callersversion()</code>	<code>callersversion()</code>	obtain version number of caller
<code>favorspeed()</code>	<code>favorspeed()</code>	whether speed or space is to be favored
Advanced parsing		
<code>tokenget()</code>	<code>tokeninit()</code>	initialize parsing environment
	<code>tokeninitstata()</code>	initialize environment as Stata would
	<code>tokenset()</code>	set/reset string to be parsed
	<code>tokengetall()</code>	parse entire string
	<code>tokenget()</code>	parse next element of string
	<code>tokenpeek()</code>	peek at next <code>tokenget()</code> result
	<code>tokenrest()</code>	return yet-to-be-parsed portion
	<code>tokenoffset()</code>	query/reset offset in string
	<code>tokenwchars()</code>	query/reset white-space characters
	<code>tokenpchars()</code>	query/reset parsing characters
	<code>tokenqchars()</code>	query/reset quote characters
	<code>tokenallownum()</code>	query/reset number parsing
	<code>tokenallowhex()</code>	query/reset hex-number parsing
Accessing externals		
<code>findexternal()</code>	<code>findexternal()</code>	find global
	<code>crexternal()</code>	create global
	<code>rmexternal()</code>	remove global
	<code>nameexternal()</code>	name of external
<code>direxternal()</code>	<code>direxternal()</code>	obtain list of existing globals
<code>valofexternal()</code>	<code>valofexternal()</code>	obtain value of global

Break key

<code>setbreakintr()</code>	<code>setbreakintr()</code> <code>querybreakintr()</code> <code>breakkey()</code> <code>breakkeyreset()</code>	turn off/on break-key interrupt whether break-key interrupt is off/on whether break key has been pressed reset break key
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Associative arrays

<code>asarray()</code>	<code>asarray()</code> <code>asarray_*(())</code>	store or retrieve element in array utility routines
<code>hash1()</code>	<code>hash1()</code>	Jenkins' one-at-a-time hash

Miscellaneous

<code>assert()</code>	<code>assert()</code> <code>asserteq()</code>	abort execution if not true abort execution if not equal
<code>c()</code>	<code>c()</code>	access <code>c()</code> value
<code>sizeof()</code>	<code>sizeof()</code>	number of bytes consumed by object
<code>swap()</code>	<code>swap()</code>	interchange contents of variables

System info

<code>byteorder()</code>	<code>byteorder()</code>	byte order used by computer
<code>stataversion()</code>	<code>stataversion()</code> <code>statasetversion()</code>	version of Stata being used version of Stata set

Exiting

<code>exit()</code>	<code>exit()</code>	terminate execution
<code>error()</code>	<code>error()</code> <code>_error()</code>	issue standard Stata error message issue error message with traceback log

Also see

[M-4] [intro](#) — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
Complex		
Re()	Re()	real part
	Im()	imaginary part
C()	C()	make complex
Sign related		
abs()	abs()	absolute value (length if complex)
sign()	sign()	sign function
	quadrant()	quadrant of value
dsign()	dsign()	FORTRAN-like DSIGN function
conj()	conj()	complex conjugate
Transcendental & square root		
exp()	exp()	exponentiation
	ln(), log()	natural logarithm
	log10()	base-10 logarithm
sqrt()	sqrt()	square root
sin()	sin()	sine
	cos()	cosine
	tan()	tangent
	asin()	arcsine
	acos()	arccosine
	atan()	arctangent
	arg()	arctangent of complex
	atan2()	two-argument arctangent
	sinh()	hyperbolic sine
	cosh()	hyperbolic cosine
	tanh()	hyperbolic tangent
	asinh()	inverse-hyperbolic sine
	acosh()	inverse-hyperbolic cosine
	atanh()	inverse-hyperbolic tangent
	pi()	value of $\pi$

Factorial & gamma

factorial()	factorial()	factorial
	lnfactorial()	natural logarithm of factorial
	gamma()	gamma function
	lngamma()	natural logarithm of gamma function
	digamma()	derivative of lngamma()
	trigamma()	second derivative of lngamma()

Modulus & integer rounding

mod()	mod()	modulus
trunc()	trunc()	truncate to integer
	floor()	round down to integer
	ceil()	round up to integer
	round()	round to closest integer or multiple

Dates

date()	clock()	%tc of string
	mdyhms()	%tc of month, day, year, hour, minute, and second
	dhms()	%tc of %td, hour, minute, and second
	hms()	%tc of hour, minute, and second
	hh()	hour of %tc
	mm()	minute of %tc
	ss()	second of %tc
	dofc()	%td of %tc
	Cofc()	%tC of %tc
	Clock()	%tC of string
	Cmdyhms()	%tC of month, day, year, hour, minute, and second
	Cdhms()	%tC of %td, hour, minute, and second
	Chms()	%tC of hour, minute, and second
	hhC()	hour of %tC
	mmC()	minute of %tC
	ssC()	second of %tC
	dofC()	%td of %tC
	date()	%td of string
	mdy()	%td of month, day, and year
	yw()	%tw of year and week
	ym()	%tm of year and month
	yq()	%tq of year and quarter
	yh()	%th of year and half
	cofd()	%tc of %td
	Cofd()	%tC of %td

---

<b>date()</b> , <i>continued</i>	dofb()	%td of %tb
	bofd()	%tb of %td
	month()	month of %td
	day()	day-of-month of %td
	year()	year of %td
	dow()	day-of-week of %td
	week()	week of %td
	quarter()	quarter of %td
	halfyear()	half-of-year of %td
	doy()	day-of-year of %td
	yearly()	%ty of string
	yofd()	%ty of %td
	dofy()	%td of %ty
	halfyearly()	%th of string
	hofd()	%th of %td
	dofh()	%td of %th
	quarterly()	%tq of string
	qofd()	%tq of %td
	dofq()	%td of %tq
	monthly()	%tm of string
	mofd()	%tm of %td
	dofm()	%td of %tm
	weekly()	%tw of string
	wofd()	%tw of %td
	dofw()	%td of %tw
	hours()	hours of milliseconds
	minutes()	minutes of milliseconds
	seconds()	seconds of milliseconds
	msofhours()	milliseconds of hours
	msofminutes()	milliseconds of minutes
	msofseconds()	milliseconds of seconds

---

## Description

With a few exceptions, the above functions are what most people would consider scalar functions, although in fact all will work with matrices, in an element-by-element fashion.

## Remarks

For other mathematical functions, see

[M-4] <a href="#">matrix</a>	Matrix functions
[M-4] <a href="#">mathematical</a>	Important mathematical functions
[M-4] <a href="#">statistical</a>	Statistical functions

## Also see

[M-4] [intro](#) — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
Solvers		
<code>cholsolve()</code>	<code>cholsolve()</code>	$A$ positive definite; symmetric or Hermitian
<code>lusolve()</code>	<code>lusolve()</code>	$A$ full rank, square, real or complex
<code>qrsolve()</code>	<code>qrsolve()</code>	$A$ general; $m \times n$ , $m \geq n$ , real or complex; least-squares generalized solution
<code>svsolve()</code>	<code>svsolve()</code>	generalized; $m \times n$ , real or complex; minimum norm, least-squares solution
Inverters		
<code>invsym()</code>	<code>invsym()</code>	generalized; real symmetric
<code>cholin()</code>	<code>cholin()</code>	positive definite; symmetric or Hermitian
<code>luinv()</code>	<code>luinv()</code>	full rank; square; real or complex
<code>qrinv()</code>	<code>qrinv()</code>	generalized; $m \times n$ , $m \geq n$ ; real or complex
<code>pinv()</code>	<code>pinv()</code>	generalized; $m \times n$ , real or complex Moore–Penrose pseudoinverse

Description

The above functions solve  $AX = B$  for  $X$  and solve for  $A^{-1}$ .

Remarks

Matrix solvers can be used to implement matrix inverters, and so the two nearly always come as a pair.

Solvers solve  $AX = B$  for  $X$ . One way to obtain  $A^{-1}$  is to solve  $AX = I$ . If `f(A, B)` solves  $AX=B$ , then `f(A, I(rows(A)))` solves for the inverse. Some matrix inverters are in fact implemented this way, although usually custom code is written because memory savings are possible when it is known that  $B = I$ .

The pairings of inverter and solver are

inverter	solver
<code>invsym()</code>	<code>(none)</code>
<code>cholinv()</code>	<code>cholsolve()</code>
<code>luinv()</code>	<code>lusolve()</code>
<code>qrinv()</code>	<code>qrsolve()</code>
<code>pinv()</code>	<code>svsolve()</code>

**Also see**

[M-4] [intro](#) — Index and guide to functions



Contents

[M-5] Manual entry	Function	Purpose
Unit & constant matrices		
<b>I()</b>	<b>I()</b>	identity matrix
<b>e()</b>	<b>e()</b>	unit vectors
<b>J()</b>	<b>J()</b>	matrix of constants
<b>designmatrix()</b>	<b>designmatrix()</b>	design matrices
Block-diagonal matrices		
<b>blockdiag()</b>	<b>blockdiag()</b>	block-diagonal matrix
Ranges		
<b>range()</b>	<b>range()</b> <b>rangen()</b>	vector over specified range vector of $n$ over specified range
<b>unitcircle()</b>	<b>unitcircle()</b>	unit circle on complex plane
Random		
<b>runiform()</b>	<b>runiform()</b> <b>rseed()</b>	uniformly distributed random numbers obtain or set random variate generator seed
Named matrices		
<b>Hilbert()</b>	<b>Hilbert()</b> <b>invHilbert()</b>	Hilbert matrices inverse Hilbert matrices
<b>Toeplitz()</b>	<b>Toeplitz()</b>	Toeplitz matrices
<b>Vandermonde()</b>	<b>Vandermonde()</b>	Vandermonde matrices

vec() & vech() transform

<b>Dmatrix()</b>	<code>Dmatrix()</code>	duplication matrices
<b>Kmatrix()</b>	<code>Kmatrix()</code>	commutation matrices
<b>Lmatrix()</b>	<code>Lmatrix()</code>	elimination matrices

## Description

The functions above create standard matrices such as the identity matrix, etc.

## Remarks

For other mathematical functions, see

[M-4] <b>matrix</b>	Matrix mathematical functions
[M-4] <b>scalar</b>	Scalar mathematical functions
[M-4] <b>mathematical</b>	Important mathematical functions

## Also see

[M-4] **intro** — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
<div>Access to data</div>		
<code>st_nvar()</code>	<code>st_nvar()</code>	number of variables
	<code>st_nobs()</code>	number of observations
<code>st_data()</code>	<code>st_data()</code>	load numeric data from Stata into matrix
	<code>st_sdata()</code>	load string data from Stata into matrix
<code>st_store()</code>	<code>st_store()</code>	store numeric data in Stata dataset
	<code>st_sstore()</code>	store string data in Stata dataset
<code>st_view()</code>	<code>st_view()</code>	make view onto num. data in Stata dataset
	<code>st_sview()</code>	make view onto string data in Stata dataset
<code>st_subview()</code>	<code>st_subview()</code>	make view from view
<code>st_viewvars()</code>	<code>st_viewvars()</code>	identify variables and observations
	<code>st_viewobs()</code>	corresponding to view
<div>Variable names &amp; indices</div>		
<code>st_varindex()</code>	<code>st_varindex()</code>	variable indices from variable names
<code>st_varname()</code>	<code>st_varname()</code>	variable names from variable indices

## Variable characteristics

<b>st_varrename()</b>	st_varrename()	rename Stata variable
<b>st_vartype()</b>	st_vartype()	storage type of Stata variable
	st_isnumvar()	whether variable is numeric
	st_isstrvar()	whether variable is string
<b>st_varformat()</b>	st_varformat()	obtain/set format of Stata variable
	st_varlabel()	obtain/set variable label
	st_varvaluelabel()	obtain/set value label
<b>st_vlexists()</b>	st_vlexists()	whether value label exists
	st_vldrop()	drop value
	st_vlmap()	map values
	st_vlsearch()	map text
	st_vlload()	load value label
	st_vlmodify()	create or modify value label

## Temporary variables &amp; time-series operators

<b>st_tempname()</b>	st_tempname()	temporary variable name
	st_tempfilename()	temporary filename
<b>st_tsrevar()</b>	st_tsrevar()	create time-series op.varname
	_st_tsrevar()	same

## Adding &amp; removing variables &amp; observations

<b>st_addobs()</b>	st_addobs()	add observations to Stata dataset
<b>st_addvar()</b>	st_addvar()	add variable to Stata dataset
<b>st_dropvar()</b>	st_dropvar()	drop variables
	st_dropobsin()	drop specified observations
	st_dropobsif()	drop selected observations
	st_keepvar()	keep variables
	st_keepobsin()	keep specified observations
	st_keepobsif()	keep selected observations
<b>st_updata()</b>	st_updata()	query/set data-have-changed flag

## Executing Stata commands

<b>stata()</b>	stata()	execute Stata command
<b>st_macroexpand()</b>	st_macroexpand()	expand Stata macros

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Accessing e(), r(), s(), macros, matrices, etc.

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<b>st_global()</b>	<code>st_global()</code>	obtain/set Stata global
	<code>st_global_hcat()</code>	obtain hidden/historical status
<b>st_local()</b>	<code>st_local()</code>	obtain/set local Stata macro
<b>st_numscalar()</b>	<code>st_numscalar()</code>	obtain/set Stata numeric scalar
	<code>st_numscalar_hcat()</code>	obtain hidden/historical status
	<code>st_strscalar()</code>	obtain/set Stata string scalar
<b>st_matrix()</b>	<code>st_matrix()</code>	obtain/set Stata matrix
	<code>st_matrix_hcat()</code>	obtain hidden/historical status
	<code>st_matrixrowstripe()</code>	obtain/set row labels
	<code>st_matrixcolstripe()</code>	obtain/set column labels
	<code>st_replacematrix()</code>	replace existing Stata matrix
<b>st_dir()</b>	<code>st_dir()</code>	obtain list of Stata objects
<b>st_rclear()</b>	<code>st_rclear()</code>	clear <code>r()</code>
	<code>st_eclear()</code>	clear <code>e()</code>
	<code>st_sclear()</code>	clear <code>s()</code>

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Parsing & verification

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<b>st_isname()</b>	<code>st_isname()</code>	whether valid Stata name
	<code>st_islmmname()</code>	whether valid local macro name
<b>st_isfmt()</b>	<code>st_isfmt()</code>	whether valid <code>%fmt</code>
	<code>st_isnumfmt()</code>	whether valid numeric <code>%fmt</code>
	<code>st_isstrfmt()</code>	whether valid string <code>%fmt</code>
<b>abbrev()</b>	<code>abbrev()</code>	abbreviate strings
<b>strtoname()</b>	<code>strtoname()</code>	translate strings to Stata names

---

## Description

The above functions interface with Stata.

## Remarks

The following manual entries have to do with getting data from or putting data into Stata:

[M-5] <b>st_data()</b>	Load copy of current Stata dataset
[M-5] <b>st_view()</b>	Make matrix that is a view onto current Stata dataset
[M-5] <b>st_store()</b>	Modify values stored in current Stata dataset
[M-5] <b>st_nvar()</b>	Numbers of variables and observations

In some cases, you may find yourself needing to translate variable names into variable indices and vice versa:

[M-5] <a href="#">st_varname()</a>	Obtain variable names from variable indices
[M-5] <a href="#">st_varindex()</a>	Obtain variable indices from variable names
[M-5] <a href="#">st_tsrevar()</a>	Create time-series op.varname variables

The other functions mostly have to do with getting and putting Stata's scalars, matrices, and returned results:

[M-5] <a href="#">st_local()</a>	Obtain strings from and put strings into Stata
[M-5] <a href="#">st_global()</a>	Obtain strings from and put strings into global macros
[M-5] <a href="#">st_numscalar()</a>	Obtain values from and put values into Stata scalars
[M-5] <a href="#">st_matrix()</a>	Obtain and put Stata matrices

The `stata()` function, documented in

[M-5] <a href="#">stata()</a>	Execute Stata command
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allows you to cause Stata to execute a command that you construct in a string.

## Reference

Gould, W. W. 2008. [Mata Matters: Macros](#). *Stata Journal* 8: 401–412.

## Also see

[M-4] [intro](#) — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
Pseudorandom variates		
runiform()	runiform()	uniform pseudorandom variates
	rseed()	obtain or set the random-variate generator seed
	rbeta()	beta pseudorandom variates
	rbinomial()	binomial pseudorandom variates
	rchi2()	chi-squared pseudorandom variates
	rdiscrete()	discrete pseudorandom variates
	rgamma()	gamma pseudorandom variates
	rhypergeometric()	hypergeometric pseudorandom variates
	rnbinomial()	negative binomial pseudorandom variates
	rnormal()	normal (Gaussian) pseudorandom variates
	rpoisson()	Poisson pseudorandom variates
	rt()	Student's <i>t</i> pseudorandom variates
Means, variances, & correlations		
mean()	mean()	mean
	variance()	variance
	quadvariance()	quad-precision variance
	meanvariance()	mean and variance
	quadmeanvariance()	quad-precision mean and variance
	correlation()	correlation
	quadcorrelation()	quad-precision correlation
cross()	cross()	$X'X$ , $X'Z$ , $X'\text{diag}(w)Z$ , etc.
corr()	corr()	make correlation from variance matrix
crossdev()	crossdev()	$(X: -x)'(X: -x)$ , $(X: -x)'(Z: -z)$ , etc.
quadcross()	quadcross()	quad-precision cross()
	quadcrossdev()	quad-precision crossdev()

Factorial & combinations

<b>factorial()</b>	factorial() lnfactorial() gamma() lngamma() digamma() trigamma()	factorial natural logarithm of factorial gamma function natural logarithm of gamma function derivative of lngamma() second derivative of lngamma()
<b>comb()</b>	comb()	combinatorial function $n$ choose $k$
<b>cvpermute()</b>	cvpermutesetup() cvpermute()	permutation setup return permutations, one at a time

Densities & distributions

<b>normal()</b>	normalden() normal() invnormal() lnnormalden() lnnormal()  binormal()  betaden() ibeta()  ibetatail() invibeta() invibetatail()  binomialp() binomial() binomialtail() invbinomial() invbinomialtail()  chi2() chi2tail() invchi2() invchi2tail()  dunnettprob()  invdunnettprob()	normal density cumulative normal dist. inverse cumulative normal logarithm of the normal density logarithm of the cumulative normal dist.  cumulative binormal dist.  beta density cumulative beta dist.; a.k.a. incomplete beta function reverse cumulative beta dist. inverse cumulative beta inverse reverse cumulative beta  binomial probability cumulative of binomial reverse cumulative of binomial inverse binomial (lower tail) inverse binomial (upper tail)  cumulative chi-squared dist. reverse cumulative chi-squared dist. inverse cumulative chi-squared inverse reverse cumulative chi-squared  cumulative multiple range distribution; used in Dunnett's multiple comparison inverse cumulative multiple range distribution; used in Dunnett's multiple comparison
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<b>normal()</b> , <i>continued</i>	
<code>Fden()</code>	$F$ density
<code>F()</code>	cumulative $F$ dist.
<code>Ftail()</code>	reverse cumulative $F$ dist.
<code>invF()</code>	inverse cumulative $F$
<code>invFtail()</code>	inverse reverse cumulative $F$
<hr/>	
<code>gammaden()</code>	gamma density
<code>gammap()</code>	cumulative gamma dist.;
	a.k.a. incomplete gamma function
<code>gammaptail()</code>	reverse cumulative gamma dist.;
<code>invgammap()</code>	inverse cumulative gamma
<code>invgammaptail()</code>	inverse reverse cumulative gamma
<code>dgammapda()</code>	$\partial P(a, x)/\partial a$ , where $P(a, x) = \text{gammap}(a, x)$
<code>dgammapdx()</code>	$\partial P(a, x)/\partial x$ , where $P(a, x) = \text{gammap}(a, x)$
<code>dgammapdada()</code>	$\partial^2 P(a, x)/\partial a^2$ , where $P(a, x) = \text{gammap}(a, x)$
<code>dgammapdadx()</code>	$\partial^2 P(a, x)/\partial a \partial x$ , where $P(a, x) = \text{gammap}(a, x)$
<code>dgammapdxdx()</code>	$\partial^2 P(a, x)/\partial x^2$ , where $P(a, x) = \text{gammap}(a, x)$
<hr/>	
<code>hypergeometricp()</code>	hypergeometric probability
<code>hypergeometric()</code>	cumulative of hypergeometric
<hr/>	
<code>nbetaden()</code>	noncentral beta density
<code>nibeta()</code>	cumulative noncentral beta dist.
<code>invnibeta()</code>	inverse cumulative noncentral beta
<hr/>	
<code>nbinomialp()</code>	negative binomial probability
<code>nbinomial()</code>	negative binomial cumulative
<code>nbinomialtail()</code>	reverse cumulative of negative binomial
<code>invnbinomial()</code>	inverse negative binomial (lower tail)
<code>invnbinomialtail()</code>	inverse negative binomial (upper tail)
<hr/>	
<code>nchi2()</code>	noncentral cumulative chi-squared dist.
<code>invnchi2()</code>	inverse noncentral cumulative chi-squared
<code>npnchi2()</code>	noncentrality parameter of <code>nchi2()</code>
<hr/>	
<code>nFden()</code>	noncentral $F$ density
<code>nFtail()</code>	reverse cumulative noncentral $F$ dist.
<code>invnFtail()</code>	inverse reverse cumulative noncentral $F$
<hr/>	
<code>poissonp()</code>	Poisson probability
<code>poisson()</code>	cumulative of Poisson
<code>poissontail()</code>	reverse cumulative of Poisson
<code>invpoisson()</code>	inverse Poisson (lower tail)
<code>invpoissontail()</code>	inverse Poisson (upper tail)
<hr/>	

<b>normal()</b> , <i>continued</i>	<code>tdden()</code>	Student $t$ density
	<code>ttail()</code>	reverse cumulative $t$ dist.
	<code>invttail()</code>	inverse reverse cumulative $t$
	<code>tukeyprob()</code>	cumulative multiple range distribution; used in Tukey's multiple comparison
	<code>invtukeyprob()</code>	inverse cumulative multiple range distribution; used in Tukey's multiple comparison

Maximization & minimization

<b>optimize()</b>	<code>optimize()</code>	function maximization and minimization
	<code>optimize_evaluate()</code>	evaluate function at initial values
	<code>optimize_init()</code>	begin optimization
	<code>optimize_init_*</code>	set details
	<code>optimize_result_*</code>	access results
<b>moptimize()</b>	<code>optimize_query()</code>	report settings
	<code>moptimize()</code>	function optimization
	<code>moptimize_evaluate()</code>	evaluate function at initial values
	<code>moptimize_init()</code>	begin setup of optimization problem
	<code>moptimize_init_*</code>	set details
	<code>moptimize_result_*</code>	access <code>moptimize()</code> results
	<code>moptimize_ado_cleanup()</code>	perform cleanup after ado
	<code>moptimize_query()</code>	report settings
	<code>moptimize_util_*</code>	utility functions for writing evaluators and processing results

Logits, odds, & related

<b>logit()</b>	<code>logit()</code>	log of the odds ratio
	<code>invlogit()</code>	inverse log of the odds ratio
	<code>cloglog()</code>	complementary log-log
	<code>invcloglog()</code>	inverse complementary log-log

Multivariate normal

<b>ghk()</b>	<code>ghk()</code>	GHK multivariate normal (MVN) simulator
	<code>ghk_init()</code>	GHK MVN initialization
	<code>ghk_init_*</code>	set details
	<code>ghk()</code>	perform simulation
	<code>ghk_query_npts()</code>	return number of simulation points
<b>ghkfast()</b>	<code>ghkfast()</code>	GHK MVN simulator
	<code>ghkfast_init()</code>	GHK MVN initialization
	<code>ghkfast_init_*</code>	set details
	<code>ghkfast()</code>	perform simulation
	<code>ghkfast_i()</code>	results for the $i$ th observation
	<code>ghk_query_*</code>	display settings

## Description

The above functions are statistical, probabilistic, or designed to work with data matrices.

## Remarks

Concerning data matrices, see

[M-4] **stata**                Stata interface functions

and especially

[M-5] **st\_data()**            Load copy of current Stata dataset

[M-5] **st\_view()**            Make matrix that is a view onto current Stata dataset

For other mathematical functions, see

[M-4] **matrix**                Matrix mathematical functions

[M-4] **scalar**                Scalar mathematical functions

[M-4] **mathematical**       Important mathematical functions

## Also see

[M-4] **intro** — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
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Parsing

<code>tokens()</code>	<code>tokens()</code>	obtain tokens (words) from string
<code>invtokens()</code>	<code>invtokens()</code>	concatenate string vector into string scalar
<code>strmatch()</code>	<code>strmatch()</code>	pattern matching
<code>tokenget()</code>	<code>...</code>	advanced parsing

Length & position

<code>strlen()</code>	<code>strlen()</code>	length of string
<code>fmtwidth()</code>	<code>fmtwidth()</code>	width of <i>%fmt</i>
<code>strpos()</code>	<code>strpos()</code>	find substring within string
<code>indexnot()</code>	<code>indexnot()</code>	find character not in list

Editing

<code>substr()</code>	<code>substr()</code>	extract substring
<code>strupper()</code>	<code>strupper()</code>	convert to uppercase
	<code>strlower()</code>	convert to lowercase
	<code>strproper()</code>	convert to proper case
<code>strtrim()</code>	<code>stritrim()</code>	replace multiple, consecutive internal blanks with one blank
	<code>strltrim()</code>	remove leading blanks
	<code>strrrtrim()</code>	remove trailing blanks
	<code>strtrim()</code>	remove leading and trailing blanks
<code>subinstr()</code>	<code>subinstr()</code>	substitute text
	<code>subinword()</code>	substitute word
<code>_substr()</code>	<code>_substr()</code>	substitute into string

---

Editing, *continued*


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<b>strdup()</b>	*	duplicate string
<b>strreverse()</b>	strreverse()	reverse string
<b>soundex()</b>	soundex()	convert to soundex code
	soundex_nara()	convert to U. S. Census soundex code

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Stata

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<b>abbrev()</b>	abbrev()	abbreviate strings
<b>strtoname()</b>	strtoname()	translate strings to Stata names

---

ASCII translation

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<b>stroofreal()</b>	stroofreal()	convert real to string
<b>strtoreal()</b>	strtoreal()	convert string to real
<b>ascii()</b>	ascii()	obtain ASCII codes of string
	char()	make string from ASCII codes

---

## Description

The above functions are for manipulating strings. Strings in Mata are strings of ASCII characters, usually the printable characters, but Mata enforces no such restriction. In particular, strings may contain binary 0.

## Remarks

In addition to the above functions, two operators are especially useful for dealing with strings.

The first is `+`. Addition is how you concatenate strings:

```
: "abc" + "def"
abcdef
: command = "list"
: args = "mpg weight"
: result = command + " " + args
: result
list mpg weight
```

The second is `*`. Multiplication is how you duplicate strings:

```
: 5*"a"
aaaaa
: "b"*3
bbb
```

```
: indent = 20
: title = indent*" " + "My Title"
: title
           My Title
```

## Also see

[\[M-4\] intro](#) — Index and guide to functions

Contents

[M-5] Manual entry	Function	Purpose
Complex		
<b>Re()</b>	Re() Im()	real part imaginary part
<b>C()</b>	C()	make complex
Shape & type		
<b>rows()</b>	rows() cols() length()	number of rows number of columns number of elements of vector
<b>eltype()</b>	eltype() orgtype()	element type of object organizational type of object
<b>isreal()</b>	isreal() iscomplex() isstring() ispointer()	object is real matrix object is complex matrix object is string matrix object is pointer matrix
<b>isrealvalues()</b>	isrealvalues()	whether matrix contains only real values
<b>isview()</b>	isview()	whether matrix is view
Properties		
<b>issymmetric()</b>	issymmetric() issymmetriconly()	whether matrix is symmetric (Hermitian) whether matrix is mechanically symmetric
<b>isdiagonal()</b>	isdiagonal()	whether matrix is diagonal
<b>diag0cnt()</b>	diag0cnt()	count 0s on diagonal
Selection		
<b>select()</b>	select() st_select()	select rows or columns select rows or columns of view

## Missing values

<b>missing()</b>	<code>missing()</code>	count of missing values
	<code>rowmissing()</code>	count of missing values, by row
	<code>colmissing()</code>	count of missing values, by column
	<code>nonmissing()</code>	count of nonmissing values
	<code>rownonmissing()</code>	count of nonmissing values, by row
	<code>colnonmissing()</code>	count of nonmissing values, by column
	<code>hasmissing()</code>	whether matrix has missing values
<b>missingof()</b>	<code>missingof()</code>	appropriate missing value

## Range, sums, &amp; cross products

<b>minmax()</b>	<code>rowmin()</code>	minimum, by row
	<code>colmin()</code>	minimum, by column
	<code>min()</code>	minimum, overall
	<code>rowmax()</code>	maximum, by row
	<code>colmax()</code>	maximum, by column
	<code>max()</code>	maximum, overall
	<code>rowminmax()</code>	minimum and maximum, by row
	<code>colminmax()</code>	minimum and maximum, by column
	<code>minmax()</code>	minimum and maximum, overall
	<code>rowmaxabs()</code>	<code>rowmax(abs())</code>
	<code>colmaxabs()</code>	<code>colmax(abs())</code>
<b>minindex()</b>	<code>minindex()</code>	indices of minimums
	<code>maxindex()</code>	indices of maximums
<b>sum()</b>	<code>rowsum()</code>	sum of each row
	<code>colsum()</code>	sum of each column
	<code>sum()</code>	overall sum
	<code>quadrowsum()</code>	quad-precision sum of each row
	<code>quadcolsum()</code>	quad-precision sum of each column
	<code>quadsum()</code>	quad-precision overall sum
<b>runningsum()</b>	<code>runningsum()</code>	running sum of vector
	<code>quadrunningsum()</code>	quad-precision runningsum()
<b>cross()</b>	<code>cross()</code>	$X'X$ , $X'Z$ , etc.
<b>crossdev()</b>	<code>crossdev()</code>	$(X: -x)'(X: -x)$ , $(X: -x)'(Z: -z)$ , etc.
<b>quadcross()</b>	<code>quadcross()</code>	quad-precision <code>cross()</code>
	<code>quadcrossdev()</code>	quad-precision <code>crossdev()</code>



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Programming

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<b>reldif()</b>	<code>reldif()</code> <code>mreldif()</code> <code>mreldifsym()</code> <code>mreldifre()</code>	relative difference max. relative difference between matrices max. relative difference from symmetry max. relative difference from real
<b>all()</b>	<code>all()</code> <code>any()</code> <code>allof()</code> <code>anyof()</code>	<code>sum(!L)==0</code> <code>sum(L)!=0</code> <code>all(P==s)</code> <code>any(P==s)</code>
<b>panelsetup()</b>	<code>panelsetup()</code> <code>panelstats()</code> <code>panelsubmatrix()</code> <code>panelsubview()</code>	initialize panel-data processing summary statistics on panels obtain matrix for panel $i$ obtain view matrix for panel $i$
<b>_negate()</b>	<code>_negate()</code>	fast negation of matrix

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Constants & tolerances

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<b>mindouble()</b>	<code>mindouble()</code> <code>maxdouble()</code> <code>smallestdouble()</code>	minimum nonmissing value maximum nonmissing value smallest $e > 0$
<b>epsilon()</b>	<code>epsilon()</code>	unit roundoff error
<b>floatround()</b>	<code>floatround()</code>	round to float precision
<b>solve_tol()</b>	<code>solve_tol()</code>	tolerance used by solvers and inverters

---

## Description

Matrix utility functions tell you something about the matrix, such as the number of rows or whether it is diagonal.

## Remarks

There is a thin line between utility and manipulation; also see

[M-4] **manipulation**

Matrix manipulation functions

## Also see

[M-4] [intro](#) — Index and guide to functions

## **[M-5] Mata functions**



# Title

[M-5] intro — Mata functions

## Contents

See [\[M-4\] intro](#)

## Description

The following entries, alphabetically arranged, document all the Mata functions.

## Remarks

See [\[M-4\] intro](#) for an index, grouped logically, of the functions presented in this section.

## Also see

[\[M-4\] intro](#) — Index and guide to functions

[\[M-0\] intro](#) — Introduction to the Mata manual

# Title

[M-5] **abbrev()** — Abbreviate strings

## Syntax

*string matrix* abbrev(*string matrix s*, *real matrix n*)

## Description

abbrev(*s*, *n*) returns the *n*-character abbreviation of *s* such that the abbreviation uniquely identifies a variable in Stata, if there are variables in Stata. Otherwise, abbrev(*s*, *n*) returns the *n*-character abbreviation of *s*.

1. *n* is the abbreviation length and is assumed to contain integer values in the range 5, 6, ..., 32.
2. If *s* contains a period, ., and *n* < 8, then the value *n* defaults to 8. Otherwise, if *n* < 5, then *n* defaults to 5.
3. If *n* is missing, the entire string (up to the first binary 0) is returned.

If there is a binary 0 in *s*, the abbreviation is derived from the beginning of the string up to but not including the binary 0.

When arguments are not scalar, abbrev() returns element-by-element results.

## Conformability

abbrev( <i>s</i> , <i>n</i> ):	
<i>s</i> :	$r_1 \times c_1$
<i>n</i> :	$r_2 \times c_2$ ; <i>s</i> and <i>n</i> r-conformable
<i>result</i> :	$\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

abbrev() returns "" if *s* is "". abbrev() aborts with error if *s* is not a string.

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] **abs()** — Absolute value (length)

## Syntax

*real matrix*    `abs(numeric matrix Z)`

## Description

For  $Z$  real, `abs(Z)` returns the elementwise absolute values of  $Z$ .

For  $Z$  complex, `abs(Z)` returns the elementwise length of each element. If  $Z = a + bi$ , returned is `sqrt(a2 + b2)`, although the calculation is not made in that way. The method actually used prevents overflow.

## Conformability

`abs(Z)`:  
           $Z$ :         $r \times c$   
           $result$ :     $r \times c$

## Diagnostics

`abs(.)` returns `.` (missing).

## Also see

[M-4] **scalar** — Scalar mathematical functions

# Title

[M-5] **adosubdir()** — Determine ado-subdirectory for file

## Syntax

*string scalar* **adosubdir**(*string scalar filename*)

## Description

**adosubdir**(*filename*) returns the subdirectory in which Stata would search for *filename*. Typically, the subdirectory will be simply the first letter of *filename*. However, certain files may result in a different subdirectory, depending on their extension.

## Remarks

**adosubdir**("xyz.ado") returns "x" because Stata ado-files are located in subdirectories with name given by their first letter.

**adosubdir**("xyz.style") returns "style" because Stata style files are located in subdirectories named `style`.

## Conformability

**adosubdir**(*filename*):

*filename*:      1 × 1

*result*:        1 × 1

## Diagnostics

**adosubdir**() returns the first letter of the filename if the filetype is unknown to Stata, thus treating unknown filetypes as if they were ado-files.

**adosubdir**() aborts with error if the filename is too long for the operating system; nothing else causes abort with error.

## Also see

[M-4] **io** — I/O functions



## Syntax

*real scalar*    all(*real matrix L*)

*real scalar*    any(*real matrix L*)

*real scalar*    allof(*transmorphic matrix P, transmorphic scalar s*)

*real scalar*    anyof(*transmorphic matrix P, transmorphic scalar s*)

## Description

all(*L*) is equivalent to sum(!*L*)==0 but is significantly faster.

any(*L*) is equivalent to sum(*L*)!=0 but is slightly faster.

allof(*P, s*) returns 1 if every element of *P* equals *s* and returns 0 otherwise. allof(*P, s*) is faster and consumes less memory than the equivalent construction all(*P*==*s*).

anyof(*P, s*) returns 1 if any element of *P* equals *s* and returns 0 otherwise. anyof(*P, s*) is faster and consumes less memory than the equivalent any(*P*==*s*).

## Remarks

These functions are fast, so their use is encouraged over alternative constructions.

all() and any() are typically used with logical expressions to detect special cases, such as

```
if (any(x :< 0)) {
    ...
}
```

or

```
if (all(x :>= 0)) {
    ...
}
```

allof() and anyof() are used to look for special values:

```
if (allof(x, 0)) {
    ...
}
```

or

```
if (anyof(x, 0)) {
    ...
}
```

Do not use `allof()` and `anyof()` to check for missing values—for example, `anyof(x, .)`—because to really check, you would have to check not only `.` but also `.a`, `.b`, `...`, `.z`. Instead use `missing()`; see [M-5] [missing\(\)](#).

## Conformability

```
all(L), any(L):  
    L:      r × c  
    result: 1 × 1  
  
allof(P, s), anyof(P, s):  
    P:      r × c  
    s:      1 × 1  
    result: 1 × 1
```

## Diagnostics

- `all(L)` and `any(L)` treat missing values in *L* as true.
- `all(L)` and `any(L)` return 0 (false) if *L* is  $r \times 0$ ,  $0 \times c$ , or  $0 \times 0$ .
- `allof(P, s)` and `anyof(P, s)` return 0 (false) if *P* is  $r \times 0$ ,  $0 \times c$ , or  $0 \times 0$ .

## Also see

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] **args()** — Number of arguments

## Syntax

*real scalar*    **args()**

## Description

**args()** returns the number of arguments actually passed to the function; see [M-2] **optargs**.

## Conformability

**args()**:  
*result:*         $1 \times 1$

## Diagnostics

None.

## Also see

[M-4] **programming** — Programming functions

## Syntax

$A =$	<code>asarray_create([ keytype                   [, keydim                   [, minsize                   [, minratio                   [, maxratio ]]]])</code>	<i>declare A</i>
	<code>asarray(A, key, a)</code>	$A[key] = a$
$a =$	<code>asarray(A, key)</code>	$a = A[key]$ or $a = \text{notfound}$
	<code>asarray_remove(A, key)</code>	delete $A[key]$ if it exists
$bool =$	<code>asarray_contains(A, key)</code>	$A[key]$ exists?
$N =$	<code>asarray_elements(A)</code>	# of elements in A
$keys =$	<code>asarray_keys(A)</code>	all keys in A
$loc =$	<code>asarray_first(A)</code>	location of first element or NULL
$loc =$	<code>asarray_next(A, loc)</code>	location of next element or NULL
$key =$	<code>asarray_key(A, loc)</code>	key at loc
$a =$	<code>asarray_contents(A, loc)</code>	contents a at loc
	<code>asarray_notfound(A, notfound)</code>	set notfound value
$notfound =$	<code>asarray_notfound(A)</code>	query notfound value

where

$A$ : Associative array  $A[key]$ . Created by `asarray_create()` and passed to the other functions. If  $A$  is declared, it is declared *transmorphic*.

*keytype*: Element type of keys; "string", "real", "complex", or "pointer".  
Optional; default "string".

*keydim*: Dimension of key;  $1 \leq keydim \leq 50$ . Optional; default 1.

- minsize*: Initial size of hash table used to speed locating keys in *A*; *real scalar*;  
 $5 \leq minsize \leq 1,431,655,764$ . Optional; default 100.
- minratio*: Fraction filled at which hash table is automatically downsized; *real scalar*;  
 $0 \leq minratio \leq 1$ . Optional; default 0.5.
- maxratio*: Fraction filled at which hash table is automatically upsized; *real scalar*;  
 $1 < maxratio \leq .$  (meaning infinity). Optional; default 1.5.
- key*: Key under which an element is stored in *A*; *string scalar* by default; type and dimension are declared using `asarray_create()`.
- a*: Element of *A*; *transmorphic*; may be anything of any size; different *A[key]* elements may have different types of contents.
- bool*: Boolean logic value; *real scalar*; equal to 1 or 0 meaning true or false.
- N*: Number of elements stored in *A*; *real scalar*;  $0 \leq N \leq 2,147,483,647$ .
- keys*: List of all keys that exist in *A*. Each row is a key. Thus *keys* is a *string colvector* if keys are *string scalars*, a *string matrix* if keys are *string vectors*, a *real colvector* if keys are *real scalars*, etc. Note that `rows(keys) = N`.
- loc*: A location in *A*; *transmorphic*. The first location is returned by `asarray_first()`, subsequent locations by `asarray_next()`. *loc*==NULL when there are no more elements.
- notfound*: Value returned by `asarray(A, key)` when *key* does not exist in *A*.  
*notfound* = `J(0,0,.)` by default.

## Description

`asarray()` provides one- and multi-dimensional associative arrays, also known as containers, maps, dictionaries, indices, and hash tables. In associative arrays, rather than being dense integers, the indices can be anything, even strings. So you might have *A*["Frank Smith"] equal to something and *A*["Mary Jones"] equal to something else. In Mata, you write that as `asarray(A, "Frank Smith", something)` and `asarray(A, "Mary Jones", somethingelse)` to define the elements and `asarray(A, "Frank Smith")` and `asarray(A, "Mary Jones")` to obtain their values.

*A* = `asarray_create()` declares (creates) an associative array. The function allows arguments, but they are optional. Without arguments, `asarray_create()` declares an associative array with string scalar keys, corresponding to the *A*["Frank Smith"] and *A*["Mary Jones"] example above.

*A* = `asarray_create(keytype, keydim)` declares an associative array with *keytype* keys each of dimension  $1 \times keydim$ . `asarray_create("string", 1)` is equivalent to `asarray_create()` without arguments. `asarray_create("string", 2)` declares the keys to be string, as before, but now they are  $1 \times 2$  rather than  $1 \times 1$ , so array elements would be of the form *A*["Frank Smith", "Mary Jones"]. *A*["Mary Jones", "Frank Smith"] would be a different element. `asarray_create("real", 2)` declares the keys to be real  $1 \times 2$ , which would somewhat correspond to our ordinary idea of a matrix, namely *A*[*i,j*]. The difference would be that to store, say, *A*[100,980], it would not be necessary to store the interior elements, and in addition to storing *A*[100,980], we could store *A*[3.14159,2.71828].

`asarray_create()` has three more optional arguments: *minsize*, *minratio*, and *maxratio*. We recommend that you do not specify them. They are discussed in [Setting the efficiency parameters](#) under *Remarks* below.

`asarray(A, key, a)` sets or resets element  $A[key] = a$ . Note that if you declare *key* to be  $1 \times 2$ , you must use the parentheses vector notation to specify key literals, such as `asarray(A, (100,980), 2.15)`. Alternatively, if  $k = (100,980)$ , then you can omit the parentheses in `asarray(A, k, 2.15)`.

`asarray(A, key)` returns element  $A[key]$  or it returns *notfound* if the element does not exist. By default, *notfound* is `J(0,0,.)`, but you can change that using `asarray_notfound()`. If you redefined *notfound* to be 0 and defined keys to be real  $1 \times 2$ , you would be on your way to recording sparse matrices efficiently.

`asarray_remove(A, key)` removes  $A[key]$ , or it does nothing if  $A[key]$  is already undefined.

`asarray_contains(A, key)` returns 1 if  $A[key]$  is defined, and it returns 0 otherwise.

`asarray_elements(A)` returns the number of elements stored in *A*.

`asarray_keys(A)` returns a vector or matrix containing all the keys, one to a row. The keys are not in alphabetical or numerical order. If you want them that way, code `sort(asarray_keys(A), 1)` if your keys are scalar, or in general, code `sort(asarray_keys(A), idx)`; see [M-5] `sort()`.

`asarray_first(A)` and `asarray_next(A, loc)` provide a way of obtaining the names one at a time. Code

```
for (loc=asarray_first(A); loc!=NULL; loc=asarray_next(A, loc)) {
    ...
}
```

`asarray_key(A, loc)` and `asarray_contents(A, loc)` return the key and contents at *loc*, so the loop becomes

```
for (loc=asarray_first(A); loc!=NULL; loc=asarray_next(A, loc)) {
    ...
    ... asarray_key(A, loc) ...
    ...
    ... asarray_contents(A, loc) ...
    ...
}
```

`asarray_notfound(A, notfound)` defines what `asarray(A, key)` returns when the element does not exist. By default, *notfound* is `J(0,0,.)`, which is to say, a  $0 \times 0$  real matrix. You can reset *notfound* at any time. `asarray_notfound(A)` returns the current value of *notfound*.

## Remarks

Before writing a program using `asarray()`, you should try it interactively. Remarks are presented under the following headings:

*Example 1: Scalar keys and scalar contents*

*Example 2: Scalar keys and matrix contents*

*Example 3: Vector keys and scalar contents; sparse matrix*

*Setting the efficiency parameters*

### Example 1: Scalar keys and scalar contents

```

: A = asarray_create()
: asarray(A, "bill", 1.25)
: asarray(A, "mary", 2.75)
: asarray(A, "dan", 1.50)
: asarray(A, "bill")
1.25
: asarray(A, "mary")
2.75
: asarray(A, "mary", 3.25)
: asarray(A, "mary")
3.25
: sum = 0
: for (loc=asarray_first(A); loc!=NULL; loc=asarray_next(A, loc)) {
>     sum = sum + asarray_contents(A, loc)
> }
: sum
6
: sum/asarray_elements(A)
2

```

### Example 2: Scalar keys and matrix contents

```

: A = asarray_create()
: asarray(A, "Count", (1,2\3,4))
: asarray(A, "Hilbert", Hilbert(3))
: asarray(A, "Count")
  1  2
1  1  2
2  3  4

```

1	1	2
2	3	4

```

: asarray(A, "Hilbert")
[symmetric]
      1      2      3
1      1
2      .5   .333333333
3   .333333333      .25      .2

```

1	1		
2	.5	.333333333	
3	.333333333	.25	.2

### Example 3: Vector keys and scalar contents; sparse matrix

```

: A = asarray_create("real", 2)
: asarray_notfound(A, 0)
: asarray(A, ( 1, 1), 1)
: asarray(A, (1000, 999), .5)
: asarray(A, (1000, 1000), 1)
: asarray(A, (1000, 1001), .5)
: asarray(A, (1,1))
1

```

```

: asarray(A, (2,2))
0
: // one way to get the trace:
: trace = 0
: for (i=1; i<=1000; i++) trace = trace + asarray(A, (i,i))
: trace
2
: // another way to get the trace
: trace = 0
: for (loc=asarray_first(A); loc!=NULL; loc=asarray_next(A, loc)) {
>     index = asarray_key(A, loc)
>     if (index[1]==index[2]) {
>         trace = trace + asarray_contents(A, loc)
>     }
> }
: trace
2

```

## Setting the efficiency parameters

The syntax `asarray_create()` is

```
A = asarray_create(keytype, keydim, minsize, minratio, maxratio)
```

All arguments are optional. The first two specify the characteristics of the key and their use has already been illustrated. The last three are efficiency parameters. In most circumstances, we recommend you do not specify them. The default values have been chosen to produce reasonable execution times with reasonable memory consumption.

`asarray()` works via hash tables. Say we wish to record  $n$  entries. The idea is to allocate a hash table of  $N$  rows, where  $N$  can be less than, equal to, or greater than  $n$ . When one needs to find the element corresponding to a key, one calculates a function of the key, called a hash function, that returns an integer  $h$  from 1 to  $N$ . One first looks in row  $h$ . If row  $h$  is already in use and the keys are different, we have a collision. In that case, we have to allocate a duplicates list for the  $h$ th row and put the duplicate keys and contents into that list. Collisions are bad because, when they occur, `asarray()` has to allocate a duplicates list, requiring both time and memory, though it does not require much. When fetching results, if row  $h$  has a duplicates list, `asarray()` has to search the list, which it does sequentially, and that takes extra time, too. Hash tables work best when collisions happen rarely.

Obviously, collisions are certain to occur if  $N < n$ . Note, however, that although performance suffers, the method does not break. A hash table of  $N$  can hold any number of entries, even if  $N < n$ .

Performance depends on details of implementation. We have examined the behavior of `asarray()` and discovered that collisions rarely occur when  $n/N \leq 0.75$ . When  $n/N = 1.5$ , performance suffers, but not by as much as you might expect. Around  $n/N = 2$ , performance degrades considerably.

When you add or remove an element, `asarray()` examines  $n/N$  and considers rebuilding the table with a larger or smaller  $N$ ; it rebuilds the table when  $n/N$  is large to preserve efficiency. It rebuilds the table when  $n/N$  is small to conserve memory. Rebuilding the table is a computer-intensive operation, and so should not be performed too often.



In making these decisions, `asarray()` uses three parameters:

*maxratio*: When  $n/N \geq \text{maxratio}$ , the table is upsize to  $N = 1.5n$ .

*minratio*: When  $n/N \leq \text{minratio}/1.5$ , the table is downsize to  $N = 1.5n$ . (For an exception, see *minsize*.)

*minsize*: If the new  $N < 1.5\text{minsize}$ , the table is downsize to  $N = 1.5\text{minsize}$  if it is not already that size.

The default values of the three parameters are 1.5, 0.5, and 100. You can reset them, though you are unlikely to improve on the default values of *minratio* and *maxratio*.

You can improve on *minsize* when you know the number of elements that will be in the table and that number is greater than 100. For instance, if you know the table will contain at least 1,000 elements, starting *minsize* at 1,000, which implies  $N = 1,500$ , will prevent two rescalings, namely, from 150 to 451, and from 451 to 1,354. This saves a little time.

You can also turn off the resizing features. Setting *minratio* to 0 turns off downsizing. Setting *maxratio* to . (missing) turns off upsizing. You might want to turn off both downsizing and upsizing if you set *minsize* sufficiently large for your problem.

We would never recommend turning off upsizing alone, and we seldom would recommend turning off downsizing alone. In a program where it is known that the array will exist for only a short time, however, turning off downsizing can be efficient. In a program where the array might exist for a considerable time, turning off downsizing is dangerous because then the array could only grow (and probably will).

## Conformability

`asarray_create(keytype, keydim, minsize, minratio, maxratio):`

<i>keytype</i> :	$1 \times 1$ (optional)
<i>keydim</i> :	$1 \times 1$ (optional)
<i>minsize</i> :	$1 \times 1$ (optional)
<i>minratio</i> :	$1 \times 1$ (optional)
<i>maxratio</i> :	$1 \times 1$ (optional)
<i>result</i> :	<i>transmorphic</i>

`asarray(A, key, a):`

<i>A</i> :	<i>transmorphic</i>
<i>key</i> :	$1 \times \text{keydim}$
<i>a</i> :	$r_{\text{key}} \times c_{\text{key}}$
<i>result</i> :	<i>void</i>

`asarray(A, key):`

<i>A</i> :	<i>transmorphic</i>
<i>key</i> :	$1 \times \text{keydim}$
<i>result</i> :	$r_{\text{key}} \times c_{\text{key}}$

`asarray_remove(A, key):`

*A:*        *transmorphic*  
*key:*      $1 \times \text{keydim}$   
*result:*   *void*

`asarray_contains(A, key), asarray_elements(A, key):`

*A:*        *transmorphic*  
*key:*      $1 \times \text{keydim}$   
*result:*    $1 \times 1$

`asarray_keys(A, key):`

*A:*        *transmorphic*  
*key:*      $1 \times \text{keydim}$   
*result:*    $n \times \text{keydim}$

`asarray_first(A):`

*A:*        *transmorphic*  
*result:*   *transmorphic*

`asarray_first(A, loc):`

*A:*        *transmorphic*  
*loc:*      *transmorphic*  
*result:*   *transmorphic*

`asarray_key(A, loc):`

*A:*        *transmorphic*  
*loc:*      *transmorphic*  
*result:*    $1 \times \text{keydim}$

`asarray_contents(A, loc):`

*A:*        *transmorphic*  
*loc:*      *transmorphic*  
*result:*    $r_{\text{key}} \times c_{\text{key}}$

`asarray_notfound(A, notfound):`

*A:*        *transmorphic*  
*notfound:*  $r \times c$   
*result:*   *void*

`asarray_notfound(A):`

*A:*        *transmorphic*  
*result:*    $r \times c$

## Diagnostics

None.

---

## Also see

[M-5] [hash1\(\)](#) — Jenkins' one-at-a-time hash function

[M-4] [manipulation](#) — Matrix manipulation

[M-4] [programming](#) — Programming functions

# Title

[M-5] **ascii()** — Manipulate ASCII codes

## Syntax

*real rowvector*    **ascii**(*string scalar s*)

*string scalar*    **char**(*real rowvector c*)

## Description

**ascii**(*s*) returns a row vector containing the ASCII codes corresponding to *s*. For instance, **ascii**("abc") returns (97, 98, 99).

**char**(*c*) returns a string consisting of the specified ASCII codes. For instance, **char**((97, 98, 99)) returns "abc".

## Conformability

**ascii**(*s*):

<i>s</i> :	$1 \times 1$
<i>result</i> :	$1 \times \text{strlen}(s)$

**char**(*c*):

<i>c</i> :	$1 \times n, \quad n \geq 0$
<i>result</i> :	$1 \times 1$

## Diagnostics

**ascii**(*s*) returns J(1,0,.) if **strlen**(*s*)==0.

In **char**(*c*), if any element of *c* is outside the range 0 to 255, the returned string is terminated at that point. For instance, **char**((97,98,99,1000,97,98,99))="abc".

**char**(J(1,0,.)) returns "".

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] `assert()` — Abort execution if false

## Syntax

```
void  assert(real scalar r)

void  asserteq(transmorphic matrix A, transmorphic matrix B)
```

## Description

`assert(r)` produces the error message “assertion is false” and aborts with error if  $r == 0$ .

`asserteq(A, B)` is logically equivalent to `assert(A==B)`. If the assertion is false, however, information is presented on the number of mismatches.

## Remarks

In the midst of complicated code, you know that a certain calculation must produce a result greater than 0, but you worry that perhaps you have an error in your code:

```
...
assert(n>0)
...
```

In another spot, you have produced matrix A and know every element of A should be positive or zero:

```
...
assert(A:>=0)
...
```

Once you are convinced that your function works, these verifications should be removed. In a third part of your code, however, the problem is different if the number of rows  $r$  exceed the number of columns  $c$ . In all the cases you need to use it, however,  $r$  will be less than  $c$ , so you are not much interested in programming the alternative solution:

```
...
assert(rows(PROBLEM) < cols(PROBLEM))
...
```

Leave that one in.

## Conformability

```
assert(r):
    r:      1 × 1
    result: void
```

```
asserteq(A, B):
```

<i>A:</i>	$r_1 \times c_1$
<i>B:</i>	$r_2 \times c_2$
<i>result:</i>	<i>void</i>

## Diagnostics

`assert(r)` aborts with error if  $r == 0$ .

`asserteq(A, B)` aborts with error if  $A \neq B$ .

## Also see

[\[M-4\] programming](#) — Programming functions

# Title

[M-5] **blockdiag()** — Block-diagonal matrix

## Syntax

*numeric matrix*   **blockdiag**(*numeric matrix*  $Z_1$ , *numeric matrix*  $Z_2$ )

## Description

**blockdiag**( $Z_1$ ,  $Z_2$ ) returns a block-diagonal matrix with  $Z_1$  in the upper-left corner and  $Z_2$  in the lower right, that is,

$$\begin{bmatrix} Z_1 & \mathbf{0} \\ \mathbf{0} & Z_2 \end{bmatrix}$$

$Z_1$  and  $Z_2$  may be either real or complex and need not be of the same type.

## Remarks

To create a block diagonal matrix of  $Z_1$ ,  $Z_2$ ,  $Z_3$ , code

`: blockdiag( $Z1$ , blockdiag( $Z2,Z3$ ))`

## Conformability

**blockdiag**( $Z_1$ ,  $Z_2$ ):

$Z_1$ :	$r_1 \times c_1$
$Z_2$ :	$r_2 \times c_2$
<i>result</i> :	$r_1 + r_2 \times c_1 + c_2$

## Diagnostics

None. Either or both  $Z_1$  and  $Z_2$  may be void.

## Also see

[M-4] **standard** — Functions to create standard matrices

## Syntax

```

colvector C = bufio()

real scalar      bufbyteorder(C)
void             bufbyteorder(C, real scalar byteorder)
real scalar      bufmissingvalue(C)
void             bufmissingvalue(C, real scalar version)

void             bufput(C, B, offset, bfmt, X)

scalar           bufget(C, B, offset, bfmt)
rowvector       bufget(C, B, offset, bfmt, c)
matrix          bufget(C, B, offset, bfmt, r, c)

void             fbufput(C, fh, bfmt, X)

scalar           fbufget(C, fh, bfmt)
rowvector       fbufget(C, fh, bfmt, c)
matrix          fbufget(C, fh, bfmt, r, c)

real scalar      bufbfmtlen(string scalar bfmt)
real scalar      bufbfmtisnum(string scalar bfmt)
    
```

where

*C*: *colvector* returned by bufio()  
*B*: *string scalar* (buffer)  
*offset*: *real scalar* (buffer position, starts at 0)  
*fh*: file handle returned by [fopen\(\)](#)  
*bfmt*: *string scalar* (binary format; see [below](#))  
*r*: *string scalar*  
*c*: *string scalar*  
*X*: value to be written; see [Remarks](#)



*bfmt* may contain

<i>bfmt</i>	meaning
<code>%{ 8 4 }z</code>	8-byte double or 4-byte double
<code>%{ 4 2 1 }b[ s u ]</code>	4-, 2-, or 1-byte integer; <i>Stata</i> , signed or unsigned
<code>%#s</code>	text string
<code>%#S</code>	binary string

## Description

These functions manipulate buffers (string scalars) containing binary data and, optionally, perform I/O.

`bufio()` returns a control vector, *C*, that you pass to the other buffer functions. *C* specifies the byte order of the buffer and specifies how missing values are to be encoded. Despite its name, `bufio()` opens no files and performs no I/O. `bufio()` merely returns a vector of default values for use with the remaining buffer functions.

`bufbyteorder()` and `bufmissingvalue()` allow changing the defaults in *C*.

`bufput()` and `bufget()` copy elements into and out of buffers. No I/O is performed. Buffers can then be written by using `fwrite()` and read by using `fread()`; see [M-5] [fopen\(\)](#).

`fbufput()` and `fbufget()` do the same, and they perform the corresponding I/O by using `fwrite()` or `fread()`.

`bufbfmtlen(bfmt)` and `bufbfmtisnum(bfmt)` are utility routines for processing *bfmts*; they are rarely used. `bufbfmtlen(bfmt)` returns the implied length, in bytes, of the specified *bfmt*, and `bufbfmtisnum(bfmt)` returns 1 if the *bfmt* is numeric, 0 if string.

## Remarks

If you wish simply to read and write matrices, etc., see `fgetmatrix()` and `fputmatrix()` and the other functions in [M-5] [fopen\(\)](#).

The functions documented here are of interest if

1. you wish to create your own binary-data format because you are writing routines in low-level languages such as FORTRAN or C and need to transfer data between your new routines and *Stata*, or
2. you wish to write a program to read and write the binary format of another software package.

These are advanced and tedious programming projects.

Remarks are presented under the following headings:

- Basics
- Argument C
- Arguments B and offset
- Argument fh
- Argument bfmt
- bfmts for numeric data
- bfmts for string data
- Argument X
- Arguments r and c
- Advanced issues

Basics

Let’s assume that you wish to write a matrix to disk so you can move it back and forth from FORTRAN. You settle on a file format in which the number of rows and number of columns are first written as 4-byte integers, and then the values of the matrix are written as 8-byte doubles, by row:

# rows	# cols	X[1,1]	X[1,2]	...
4 bytes	4 bytes	8 bytes	8 bytes	

One solution to writing matrices in such a format is

```
fh = fopen("filename", "w")
C = bufio()
fbufput(C, fh, "%4b", rows(X))
fbufput(C, fh, "%4b", cols(X))
fbufput(C, fh, "%8z", X)
fclose(fh)
```

The code to read the matrix back is

```
fh = fopen("filename", "r")
C = bufio()
rows = fbufget(C, fh, "%4b")
cols = fbufget(C, fh, "%4b")
X = fbufget(C, fh, "%8z", rows, cols)
fclose(fh)
```

Another solution, which would be slightly more efficient, is

```
fh = fopen("filename", "w")
C = bufio()
buf = 8*char(0)
bufput(C, buf, 0, "%4b", rows(X))
bufput(C, buf, 4, "%4b", cols(X))
fwrite(C, buf)
fbufput(C, fh, "%8z", X)
fclose(fh)
```

and

```
fh  = fopen("filename", "r")
C   = bufio()
buf = fread(fh, 8)
rows = bufget(C, buf, 0, "%4b")
cols = bufget(C, buf, 4, "%4b")
X    = fbufget(C, fh, "%8z", rows, cols)
fclose(fh)
```

What makes the above approach more efficient is that, rather than writing 4 bytes (the number of rows), and 4 bytes again (the number of columns), we created one 8-byte buffer and put the two 4-byte fields in it, and then we wrote all 8 bytes at once. We did the reverse when we read back the data: we read all 8 bytes and then broke out the fields. The benefit is minuscule here but, in general, writing longer buffers results in faster I/O.

In all the above examples, we wrote and read the entire matrix with one function call,

```
fbufput(C, fh, "%8z", X)
```

and

```
X    = fbufget(C, fh, "%8z", rows, cols)
```

Perhaps you would have preferred our having coded

```
for (i=1; i<=rows(X); i++) {
    for (j=1; j<=cols(X); j++) {
        fbufput(C, fh, "%8z", X[i,j])
    }
}
```

and perhaps you would have preferred our having coded something similar to read back the matrix. Had we done so, the results would have been the same.

If you are familiar with FORTRAN, you know that it records matrices in column-dominant order, rather than the row-dominant order used by Mata. It would be a little easier to code the FORTRAN side of things if we changed our file-format design to write columns first:

# rows	# cols	X[1,1]	X[2,1]	...
4 bytes	4 bytes	8 bytes	8 bytes	

One way we could do that would be to write the loops ourselves:

```
fh = fopen("filename", "w")
C  = bufio()
fbufput(C, fh, "%4b", rows(X))
fbufput(C, fh, "%4b", cols(X))
for (j=1; j<=cols(X); j++) {
    for (i=1; i<=rows(X); i++) {
        fbufput(C, fh, "%8z", X[i,j])
    }
}
```

and

```
fh  = fopen("filename", "r")
C   = bufio()
rows = fbufget(C, fh, "%4b")
cols = fbufget(C, fh, "%4b")
X    = J(rows, cols, .)
for (j=1; j<=cols(X); j++) {
    for (i=1; i<=rows(X); i++) {
        X[i,j] = fbufget(C, fh, "%8z")
    }
}
```

We could do that, but there are more efficient and easier ways to proceed. For instance, we could simply transpose the matrix before writing and after reading, and if we do that transposition in place, our code will use no extra memory:

```
fh = fopen("filename", "w")
C  = bufio()
fbufput(C, fh, "%4b", rows(X))
fbufput(C, fh, "%4b", cols(X))
_transpose(X)
fbufput(C, fh, "%8z", X)
_transpose(X)
fclose(fh)
```

The code to read the matrices back is

```
fh  = fopen("filename", "r")
C   = bufio()
rows = fbufget(C, fh, "%4b")
cols = fbufget(C, fh, "%4b")
X    = fbufget(C, fh, "%8z", cols, rows)
_transpose(X)
fclose(fh)
```

## Argument C

Argument *C* in

```
bufput(C, B, offset, bfmt, X),
bufget(C, B, offset, bfmt, ...),
fbufput(C, fh, bfmt, X), and
fbufget(C, fh, bfmt, ...)
```

specifies the control vector. You obtain *C* by coding

```
C = bufio()
```

`bufio()` returns *C*, which is nothing more than a vector filled in with default values. The other buffer routines know how to interpret the vector. The vector contains two pieces of information:

1. The byte order to be used
2. The missing-value coding scheme to be used

Some computer hardware writes numbers left to right (for example, Sun), and other computer hardware writes numbers right to left (for example, Intel); see [M-5] `byteorder()`. If you are going to write binary files, and if you want your files to be readable on all computers, you must write code to deal with this issue.

Many programmers ignore the issue because the programs they write are intended for one computer or on computers like the one they use. If that is the case, you can ignore the issue, too. The default byte order filled in by `bufio()` is the byte order of the computer you are using.

If you intend to read and write files across different computers, however, you will need to concern yourself with byte order, and how you do that is described in [Advanced issues](#) below.

The other issue you may need to consider is missing values. If you are writing a binary format that is intended to be used outside Stata, it is best if the values you write simply do not include missing values. Not all packages have them, and the packages that do don't agree on how they are encoded. In such cases, if the data you are writing include missing values, change the values to another value such as `-1`, `99`, `999`, or `-9999`.

If, however, you are writing binary files in Stata to be read back in Stata, you can allow Stata's missing values `.`, `.a`, `.b`, `...`, `.z`. No special action is required. The missing-value scheme in *C* specifies how those missing values are encoded, and there is only one way right now, so there is in fact no issue at all. *C* includes the extra information in case Stata ever changes the way it encodes missing values so that you will have a way to read and write old-format files. How this process works is described in [Advanced issues](#).

## Arguments **B** and **offset**

### Functions

`bufput(C, B, offset, bfmt, X)` and

`bufget(C, B, offset, bfmt, ...)`

do not perform I/O; they copy values into and out of the buffer. *B* specifies the buffer, and *offset* specifies the position within it.

*B* is a string scalar.

*offset* is an integer scalar specifying the position within *B*. Offset 0 specifies the first byte of *B*.

For `bufput()`, *B* must already exist and be long enough to receive the result, and it is usual to code something like

```
B = (4 + 4 + rows(X)*cols(X)*8) * char(0)
bufput(C, B, 0, "%4b", rows(X))
bufput(C, B, 4, "%4b", cols(X))
bufput(C, B, 8, "%8z", X)
```

## Argument fh

Argument *fh* in

```
fbufput(C, fh, bfmt, X) and  
fbufget(C, fh, bfmt, ...)
```

plays the role of arguments *B* and *offset* in `bufput()` and `bufget()`. Rather than copy into or out of a buffer, data are written to, or read from, file *fh*. *fh* is obtained from `fopen()`; see [M-5] `fopen()`.

## Argument bfmt

Argument *bfmt* in

```
bufput(C, B, offset, bfmt, X),  
bufget(C, B, offset, bfmt, ...),  
fbufput(C, fh, bfmt, X), and  
fbufget(C, fh, bfmt, ...)
```

specifies how the elements are to be written or read.

## bfmts for numeric data

The numeric *bfmts* are

<i>bfmt</i>	Interpretation
%8z	8-byte floating point
%4z	4-byte floating point
%4bu	4-byte unsigned integer
%4bs	4-byte signed integer
%4b	4-byte Stata integer
%2bu	2-byte unsigned integer
%2bs	2-byte signed integer
%2b	2-byte Stata integer
%1bu	1-byte unsigned integer
%1bs	1-byte signed integer
%1b	1-byte Stata integer

A Stata integer is the same as a signed integer, except that the largest 27 values are given the interpretation `., .a, .b, ..., .z`.

## bfmts for string data

The string *bfmts* are

<i>bfmt</i>	Interpretation
<code>%#s</code>	text string
<code>%#S</code>	binary string

where *#* represents the length of the string field. Examples include `%8s` and `%639876S`.

When writing, it does not matter whether you use `%#s` or `%#S`, the same actions are taken:

1. If the string being written is shorter than *#*, the field is padded with `char(0)`.
2. If the string being written is longer than *#*, only the first *#* bytes of the string are written.

When reading, the distinction between `%#s` and `%#S` is important:

1. When reading with `%#s`, if `char(0)` appears within the first *#* bytes, the returned result is truncated at that point.
2. When reading with `%#S`, a full *#* bytes are returned in all cases.

## Argument X

Argument *X* in

```
bufput(C, B, offset, bfmt, X) and
fbufput(C, fh, bfmt, X)
```

specifies the value to be written. *X* may be real or string and may be a scalar, vector, or matrix. If *X* is a vector, the elements are written one after the other. If *X* is a matrix, the elements of the first row are written one after the other, followed by the elements of the second row, and so on.

In

```
X = bufget(C, B, offset, bfmt, ...) and
X = fbufget(C, fh, bfmt, ...)
```

*X* is returned.

## Arguments r and c

Arguments *r* and *c* are optional in the following:

```
X = bufget(C, B, offset, bfmt),
X = bufget(C, B, offset, bfmt, c),
X = bufget(C, B, offset, bfmt, r, c),
X = fbufget(C, fh, bfmt),
```

$X = \text{fbufget}(C, fh, bfmt, c)$ , and

$X = \text{fbufget}(C, fh, bfmt, r, c)$ .

If  $r$  is not specified, results are as if  $r = 1$ .

If  $c$  is not specified, results are as if  $c = 1$ .

Thus

$X = \text{bufget}(C, B, offset, bfmt)$  and

$X = \text{fbufget}(C, fh, bfmt)$

read one element and return it, whereas

$X = \text{bufget}(C, B, offset, bfmt, c)$  and

$X = \text{fbufget}(C, fh, bfmt, c)$

read  $c$  elements and return them in a column vector, and

$X = \text{bufget}(C, B, offset, bfmt, r, c)$  and

$X = \text{fbufget}(C, fh, bfmt, r, c)$

read  $r * c$  elements and return them in an  $r \times c$  matrix.

## Advanced issues

A properly designed binary-file format includes a signature line first thing in the file:

```
fh = fopen(filename, "w")
fwrite(fh, "MyFormat For Mats v. 1.0")
/* ----+----1----+----2---- */
```

and

```
fh = fopen(filename, "r")
if (fread(fh, 24) != "MyFormat For Mats v. 1.0") {
    fprintf("%s not My-Format file\n", filename)
    exit(610)
}
```

If you are concerned with byte order and mapping of missing values, you should write the byte order and missing-value mapping in the file, write in natural byte order, and be prepared to read back in either byte order.

The code for writing is

```
fh = fopen(filename, "w")
fwrite(fh, "MyFormat For Mats v. 1.0")

C = bufio()
fbufput(C, fh, "%1bu", bufbyteorder(C))
fbufput(C, fh, "%2bu", bufmissingvalue(C))
```



and the corresponding code for reading the file is

```
fh = fopen(filename, "r")
if (fread(fh, 24) != "MyFormat For Mats v. 1.0") {
    fprintf("%s not My-Format file\n", filename)
    exit(610)
}

C = bufio()
bufbyteorder(C, fbufget(C, "%1bu"))
bufmissingvalue(C, fbufget(C, "%2bu"))
```

All we write in the file before recording the byte order are strings and bytes. This way, when we read the file later, we can set the byte order before reading any 2-, 4-, or 8-byte fields.

`bufbyteorder(C)`—`bufbyteorder()` with one argument—returns the byte-order encoding recorded in *C*. It returns 1 (meaning HILO) or 2 (meaning LOHI).

`bufbyteorder(C, value)`—`bufbyteorder()` with two arguments—resets the byte order recorded in *C*. Once reset, all buffer functions will automatically reverse bytes if necessary.

`bufmissingvalue()` works the same way. With one argument, it returns a code for the encoding scheme recorded in *C* (said code being the Stata release number multiplied by 100). With two arguments, it resets the code. Once the code is reset, all buffer routines used will automatically take the appropriate action.

## Conformability

`bufio()`:

*result:*      *colvector*

`bufbyteorder(C)`:

*C:*            *colvector*      made by `bufio()`  
*result:*       $1 \times 1$           containing 1 (HILO) or 2 (LOHI)

`bufbyteorder(C, byteorder)`:

*C:*            *colvector*      made by `bufio()`  
*byteorder:*    $1 \times 1$           containing 1 (HILO) or 2 (LOHI)  
*result:*      *void*

`bufmissingvalue(C)`:

*C:*            *colvector*      made by `bufio()`  
*result:*       $1 \times 1$

`bufmissingvalue(C, version)`:

*C:*            *colvector*      made by `bufio()`  
*version:*       $1 \times 1$   
*result:*      *void*

bufput(*C*, *B*, *offset*, *bfmt*, *X*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>B</i> :	$1 \times 1$	
<i>offset</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>X</i> :	$r \times c$	
<i>result</i> :	<i>void</i>	

bufget(*C*, *B*, *offset*, *bfmt*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>B</i> :	$1 \times 1$	
<i>offset</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>result</i> :	$1 \times 1$	

bufget(*C*, *B*, *offset*, *bfmt*, *r*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>B</i> :	$1 \times 1$	
<i>offset</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>r</i> :	$1 \times 1$	
<i>result</i> :	$1 \times c$	

bufget(*C*, *B*, *offset*, *bfmt*, *r*, *c*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>B</i> :	$1 \times 1$	
<i>offset</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>r</i> :	$1 \times 1$	
<i>c</i> :	$1 \times 1$	
<i>result</i> :	$r \times c$	

fbufput(*C*, *fh*, *bfmt*, *X*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>fh</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>X</i> :	$r \times c$	
<i>result</i> :	<i>void</i>	

fbufget(*C*, *fh*, *bfmt*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>fh</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>result</i> :	$1 \times 1$	

fbufget(*C*, *fh*, *bfmt*, *r*):

<i>C</i> :	<i>colvector</i>	made by bufio()
<i>fh</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>r</i> :	$1 \times 1$	
<i>result</i> :	$1 \times c$	

`fbufget(C, fh, bfmt, r, c):`

<i>C</i> :	<i>colvector</i>	made by <code>bufio()</code>
<i>fh</i> :	$1 \times 1$	
<i>bfmt</i> :	$1 \times 1$	
<i>r</i> :	$1 \times 1$	
<i>c</i> :	$1 \times 1$	
<i>result</i> :	$r \times c$	

`bufbfmtlen(bfmt):`

<i>bfmt</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

`bufbfmtisnum(bfmt):`

<i>bfmt</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

`bufio()` cannot fail.

`bufbyteorder(C)` cannot fail. `bufbyteorder(C, byteorder)` aborts with error if *byteorder* is not 1 or 2.

`bufmissingvalue(C)` cannot fail. `bufmissingvalue(C, version)` aborts with error if *version* < 100 or *version* > `stataversion()`.

`bufput(C, B, offset, bfmt, X)` aborts with error if *B* is too short to receive the result, *offset* is out of range, *bfmt* is invalid, or *bfmt* is a string format and *X* is numeric or vice versa. Putting a void matrix results in 0 bytes being inserted into the buffer and is not an error.

`bufget(C, B, offset, bfmt, ...)` aborts with error if *B* is too short, *offset* is out of range, or *bfmt* is invalid. Reading zero rows or columns results in a void returned result and is not an error.

`fbufput(C, fh, bfmt, X)` aborts with error if *fh* is invalid, *bfmt* is invalid, or *bfmt* is a string format and *X* is numeric or vice versa. Putting a void matrix results in 0 bytes being written and is not an error. I/O errors are possible; use `fstatus()` to detect them.

`fbufget(C, fh, bfmt, ...)` aborts with error if *fh* is invalid or *bfmt* is invalid. Reading zero rows or columns results in a void returned result and is not an error. End-of-file and I/O errors are possible; use `fstatus()` to detect them.

`bufbfmtlen(bfmt)` and `bufbfmtisnum(bfmt)` abort with error if *bfmt* is invalid.

## Also see

[M-5] `fopen()` — File I/O

[M-4] `io` — I/O functions

# Title

**[M-5] `byteorder()`** — Byte order used by computer

## Syntax

*real scalar* `byteorder()`

## Description

`byteorder()` returns 1 if the computer is HILO (records most significant byte first) and returns 2 if LOHI (records least significant byte first).

## Remarks

Pretend that the values 00 and 01 are recorded at memory positions 58 and 59 and that you know what is recorded there is a 2-byte integer. How should the 2-byte number be interpreted: as 0001 (meaning 1) or 0100 (meaning 256 in decimal)? For different computers, the answer varies. For HILO computers, the number is to be interpreted as 0001. For LOHI computers, the number is interpreted as 0100.

Regardless, it does not matter because the computer is consistent with itself. An issue arises, however, when we write binary files that may be read on computers using a different byte order or when we read files from computers that used a different byte order.

Stata and Mata automatically handle these problems for you, so you may wish to stop reading. `byteorder()`, however, is the tool on which the solution is based. If you intend to write code based on your own binary-file format or to write code to process the binary files of others, then you may need to use it.

There are two popular solutions to the byte-order problem: (1) write the file in a known byte order or (2) write the file by using whichever byte order is convenient and record the byte order used in the file. StataCorp tends to use the second, but others who have worried about this problem have used both solutions.

In solution (1), it is usually agreed that the file will be written in HILO order. If you are using a HILO computer, you write and read files the ordinary way, ignoring the problem altogether. If you are on a LOHI computer, however, you must reverse the bytes before placing them in the file. If you are writing code designed to execute on both kinds of computers, you must write code for both circumstances, and you must consider the problem when both reading and writing the files.

In solution (2), files are written LOHI or HILO, depending solely on the computer being used. Early in the file, however, the byte order is recorded. When reading the file, you compare the order in which the file is recorded with the order of the computer and, if they are different, you reverse bytes.

Mata-buffered I/O utilities will automatically reverse bytes for you. See [\[M-5\] `bufio\(\)`](#).

## Conformability

`byteorder()`:  
*result*:  $1 \times 1$

## Diagnostics

None.

## Also see

[M-5] [bufio\(\)](#) — Buffered (binary) I/O

[M-5] [stataversion\(\)](#) — Version of Stata being used

[M-4] [programming](#) — Programming functions

# Title

[M-5] C() — Make complex

## Syntax

*complex matrix*    `C(numeric matrix A)`

*complex matrix*    `C(real matrix R, real matrix I)`

## Description

`C(A)` returns  $A$  converted to complex. `C(A)` returns  $A$  if  $A$  is already complex. If  $A$  is real, `C(A)` returns  $A+0i$ — $A$  cast up to complex. Coding `C(A)` is thus how you ensure that the matrix is treated as complex.

`C(R, I)` returns the complex matrix  $R+Ii$  and is faster than the alternative  $R + I:*1i$ .

## Remarks

Many of Mata's functions are overloaded, meaning they return a real when given real arguments and a complex when given complex arguments. Given real arguments, if the result cannot be expressed as a real, missing value is returned. Thus `sqrt(-1)` evaluates to missing, whereas `sqrt(-1+0i)` is  $1i$ .

`C()` is the fast way to make arguments that might be real into complex. You can code

```
result = sqrt(C(x))
```

If  $x$  already is complex, `C()` does nothing; if  $x$  is real, `C(x)` returns the complex equivalent.

The two-argument version of `C()` is less frequently used. `C(R, I)` is literally equivalent to  $R :+ I*1i$ , meaning that  $R$  and  $I$  need only be c-conformable.

For instance, `C(1, (1,2,3))` evaluates to  $(1+1i, 1+2i, 1+3i)$ .

## Conformability

`C(A)`:

<i>A</i> :	$r \times c$
<i>result</i> :	$r \times c$

`C(R, I)`:

<i>R</i> :	$r_1 \times c_1$
<i>I</i> :	$r_2 \times c_2$ , $R$ and $I$ c-conformable
<i>result</i> :	$\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

`C(Z)`, if  $Z$  is complex, literally returns  $Z$  and not a copy of  $Z$ . This makes execution of `C()` applied to complex arguments instant.

In `C(R, I)`, the  $i, j$  element of the result will be missing anywhere  $R[i, j]$  or  $I[i, j]$  is missing. For instance, `C((1,3,.), (. ,2,4))` results in `(. , 3+2i, .)`. If  $R[i, j]$  and  $I[i, j]$  are both missing, then the  $R[i, j]$  value will be used; for example, `C(.a, .b)` results in `.a`.

## Also see

[M-5] **Re()** — Extract real or imaginary part

[M-4] **scalar** — Scalar mathematical functions

[M-4] **utility** — Matrix utility functions

# Title

[M-5] `c()` — Access `c()` value

## Syntax

*scalar* `c(string scalar name)`

returned is either a real or string scalar, depending on the value of *name*.

## Description

`c(name)` returns Stata's c-class value; see [P] [creturn](#).

Do not confuse `c()` with `C()`, which makes complex out of real arguments; see [M-5] [C\(\)](#).

## Remarks

See [P] [creturn](#) or, in Stata, type

```
. creturn list
```

to see what is stored in `c()`. Among the especially useful `c()` values are

```
string c("current_date")
string c("current_time")
string c("os")
string c("dirsep")
```

## Conformability

`c(name)`:

<i>name</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

`c(name)` returns a string or real depending on the particular `c()` value requested. If *name* is an invalid name or contains a name for which no `c()` value is defined, returned is "".

## Also see

[M-4] [programming](#) — Programming functions



# Title

[M-5] **callersversion()** — Obtain version number of caller

## Syntax

*real scalar*    **callersversion()**

## Description

**callersversion()** returns the version set by the caller (see [M-2] [version](#)), or if the caller did not set the version, it returns the version of Stata under which the caller was compiled.

## Remarks

**callersversion()** is how [M-2] [version](#) is made to work. Say that you have written function

*real matrix* **useful**(*real matrix* *A*, *real scalar* *k*)

and assume that **useful()** aborts with error if *A* is void. You wrote **useful()** in the days of Stata 11. For Stata 12, you want to change **useful()** so that it returns **J(0,0,.)** if *A* is void, but you want to maintain the current behavior for old Stata 11 callers and programs. You do that as follows:

```
real matrix useful(real matrix A, real scalar k)
{
    ...
    if (callersversion()>=12) {
        if (rows(A)==0 | cols(A)==0) return(J(0,0,.))
    }
    ...
}
```

## Conformability

**callersversion()**:  
*result:*      $1 \times 1$

## Diagnostics

None.

## Also see

[M-2] [version](#) — Version control

[M-4] [programming](#) — Programming functions

# Title

[M-5] **cat()** — Load file into string matrix

## Syntax

*string colvector* **cat**(*string scalar filename* [ , *real scalar line1* [ , *real scalar line2* ] ])

## Description

**cat**(*filename*) returns a column vector containing the lines from ASCII file *filename*.

**cat**(*filename*, *line1*) returns a column vector containing the lines from ASCII file *filename* starting with line number *line1*.

**cat**(*filename*, *line1*, *line2*) returns a column vector containing the lines from ASCII file *filename* starting with line number *line1* and ending with line number *line2*.

## Remarks

**cat**(*filename*) removes new-line characters at the end of lines.

## Conformability

**cat**(*filename*, *line1*, *line2*):

<i>filename</i> :	$1 \times 1$	
<i>line1</i> :	$1 \times 1$	(optional)
<i>line2</i> :	$1 \times 1$	(optional)
<i>result</i> :	$r \times 1$ ,	$r \geq 0$

## Diagnostics

**cat**(*filename*) aborts with error if *filename* does not exist.

**cat**() returns a  $0 \times 1$  result if *filename* contains 0 bytes.

## Also see

[M-4] **io** — I/O functions

## Syntax

```
string scalar  pwd()

void           chdir(string scalar dirpath)
real scalar   _chdir(string scalar dirpath)

void           mkdir(string scalar dirpath)
void           mkdir(string scalar dirpath, real scalar public)
real scalar   _mkdir(string scalar dirpath)
real scalar   _mkdir(string scalar dirpath, real scalar public)

void           rmdir(string scalar dirpath)
real scalar   _rmdir(string scalar dirpath)
```

## Description

`pwd()` returns the full name (path) of the current working directory.

`chdir(dirpath)` changes the current working directory to *dirpath*. `chdir()` aborts with error if the directory does not exist or the operating system cannot change to it.

`_chdir(dirpath)` does the same thing but returns 170 (a return code) when `chdir()` would abort. `_chdir()` returns 0 if it is successful.

`mkdir(dirpath)` and `mkdir(dirpath, public)` create directory *dirpath*. `mkdir()` aborts with error if the directory already exists or cannot be created. If *public*  $\neq$  0 is specified, the directory is given permissions so that everyone can read it; otherwise, it is given the usual permissions.

`_mkdir(dirpath)` and `_mkdir(dirpath, public)` do the same thing but return 693 (a return code) when `mkdir()` would abort. `_mkdir()` returns 0 if it is successful.

`rmdir(dirpath)` removes directory *dirpath*. `rmdir()` aborts with error if the directory does not exist, is not empty, or the operating system refuses to remove it.

`_rmdir(dirpath)` does the same thing but returns 693 (a return code) when `rmdir()` would abort. `_rmdir()` returns 0 if it is successful.

## Conformability

`pwd()`:

*result:*         $1 \times 1$

`chdir(dirpath)`:

*dirpath:*        $1 \times 1$

*result:*        *void*

`_chdir(dirpath)`:

*dirpath:*        $1 \times 1$

*result:*         $1 \times 1$

`mkdir(dirpath, public)`:

*dirpath:*        $1 \times 1$

*public:*         $1 \times 1$     (optional)

*result:*        *void*

`_mkdir(dirpath, public)`:

*dirpath:*        $1 \times 1$

*public:*         $1 \times 1$     (optional)

*result:*         $1 \times 1$

`rmdir(dirpath)`:

*dirpath:*        $1 \times 1$

*result:*        *void*

`_rmdir(dirpath)`:

*dirpath:*        $1 \times 1$

*result:*         $1 \times 1$

## Diagnostics

`pwd()` never aborts with error, but it can return "" if the operating system does not know or does not have a name for the current directory (which happens when another process removes the directory in which you are working).

`chdir(dirpath)` aborts with error if the directory does not exist or the operating system cannot change to it.

`_chdir(dirpath)` never aborts with error; it returns 0 on success and 170 on failure.

`mkdir(dirpath)` and `mkdir(dirpath, public)` abort with error if the directory already exists or the operating system cannot change to it.

`_mkdir(dirpath)` and `_mkdir(dirpath, public)` never abort with error; they return 0 on success and 693 on failure.

`rmdir(dirpath)` aborts with error if the directory does not exist, is not empty, or the operating system cannot remove it.

`_rmdir(dirpath)` never aborts with error; it returns 0 on success and 693 on failure.

---

## Also see

[M-4] [io](#) — I/O functions

# Title

[M-5] **cholesky()** — Cholesky square-root decomposition

## Syntax

*numeric matrix*      **cholesky**(*numeric matrix A*)

*void*                    **\_cholesky**(*numeric matrix A*)

## Description

**cholesky**(*A*) returns the Cholesky decomposition *G* of symmetric ([Hermitian](#)), positive-definite matrix *A*. **cholesky**() returns a lower-triangular matrix of missing values if *A* is not positive definite.

**\_cholesky**(*A*) does the same thing, except that it overwrites *A* with the Cholesky result.

## Remarks

The Cholesky decomposition *G* of a symmetric, positive-definite matrix *A* is

$$A = GG'$$

where *G* is lower triangular. When *A* is complex, *A* must be Hermitian, and *G'*, of course, is the conjugate transpose of *G*.

Decomposition is performed via [\[M-1\]](#) **LAPACK**.

## Conformability

**cholesky**(*A*):

*A*:  $n \times n$

*result*:  $n \times n$

**\_cholesky**(*A*):

*input*:

*A*:  $n \times n$

*output*:

*A*:  $n \times n$

## Diagnostics

**cholesky**() returns a lower-triangular matrix of missing values if *A* contains missing values or if *A* is not positive definite.

**\_cholesky**(*A*) overwrites *A* with a lower-triangular matrix of missing values if *A* contains missing values or if *A* is not positive definite.

Both functions use the elements from the lower triangle of  $A$  without checking whether  $A$  is symmetric or, in the complex case, Hermitian.

André-Louis Cholesky (1875–1918) was born near Bordeaux in France. He studied at the Ecole Polytechnique and then joined the French army. Cholesky served in Tunisia and Algeria and then worked in the Geodesic Section of the Army Geographic Service, where he invented his now-famous method. In the war of 1914–1918, he served in the Vosges and in Romania but after return to the Western front was fatally wounded. Cholesky's method was written up posthumously by one of his fellow officers but attracted little attention until the 1940s.

## Reference

Chabert, J.-L., É. Barbin, J. Borowczyk, M. Guillemot, and A. Michel-Pajus. 1999. *A History of Algorithms: From the Pebble to the Microchip*. Trans. C. Weeks. Berlin: Springer.

## Also see

[M-5] `lud()` — LU decomposition

[M-4] `matrix` — Matrix functions

# Title

[M-5] **cholinv()** — Symmetric, positive-definite matrix inversion

## Syntax

```
numeric matrix    cholinv(numeric matrix A)
numeric matrix    cholinv(numeric matrix A, real scalar tol)

void              _cholinv(numeric matrix A)
void              _cholinv(numeric matrix A, real scalar tol)
```

## Description

`cholinv(A)` and `cholinv(A, tol)` return the inverse of real or complex, symmetric ([Hermitian](#)), positive-definite, square matrix *A*.

`_cholinv(A)` and `_cholinv(A, tol)` do the same thing except that, rather than returning the inverse matrix, they overwrite the original matrix *A* with the inverse.

In all cases, optional argument *tol* specifies the tolerance for determining singularity; see [Remarks](#) below.

## Remarks

These routines calculate the inverse of a symmetric, positive-definite square matrix *A*. See [\[M-5\] luinv\(\)](#) for the inverse of a general square matrix.

*A* is required to be square and positive definite. See [\[M-5\] qrinv\(\)](#) and [\[M-5\] pinv\(\)](#) for generalized inverses of nonsquare or rank-deficient matrices. See [\[M-5\] invsym\(\)](#) for generalized inverses of real, symmetric matrices.

`cholinv(A)` is logically equivalent to `cholsolve(A, I(rows(A)))`; see [\[M-5\] cholsolve\(\)](#) for details and for use of the optional *tol* argument.

## Conformability

`cholinv(A, tol):`

<i>A</i> :	$n \times n$	
<i>tol</i> :	$1 \times 1$	(optional)
<i>result</i> :	$n \times n$	

`_cholinv(A, tol):`

*input*:

<i>A</i> :	$n \times n$	
<i>tol</i> :	$1 \times 1$	(optional)

*output*:

<i>A</i> :	$n \times n$	
------------	--------------	--



## Diagnostics

The inverse returned by these functions is real if  $A$  is real and is complex if  $A$  is complex. If you use these functions with a non-positive-definite matrix, or a matrix that is too close to singularity, returned will be a matrix of missing values. The determination of singularity is made relative to *tol*. See [Tolerance](#) under *Remarks* in [M-5] `cholsolve()` for details.

`cholinv(A)` and `_cholinv(A)` return a result containing all missing values if  $A$  is not positive definite or if  $A$  contains missing values.

`_cholinv(A)` aborts with error if  $A$  is a view.

See [M-5] `cholsolve()` and [M-1] `tolerance` for information on the optional *tol* argument.

Both functions use the elements from the lower triangle of  $A$  without checking whether  $A$  is symmetric or, in the complex case, Hermitian.

## Also see

[M-5] `invsym()` — Symmetric real matrix inversion

[M-5] `luinv()` — Square matrix inversion

[M-5] `qrinv()` — Generalized inverse of matrix via QR decomposition

[M-5] `pinv()` — Moore–Penrose pseudoinverse

[M-5] `cholsolve()` — Solve  $AX=B$  for  $X$  using Cholesky decomposition

[M-5] `solve_tol()` — Tolerance used by solvers and inverters

[M-4] `matrix` — Matrix functions

[M-4] `solvers` — Functions to solve  $AX=B$  and to obtain  $A$  inverse

# Title

[M-5] **cholsolve()** — Solve  $AX=B$  for  $X$  using Cholesky decomposition

## Syntax

<i>numeric matrix</i>	<code>cholsolve(<i>numeric matrix A</i>, <i>numeric matrix B</i>)</code>
<i>numeric matrix</i>	<code>cholsolve(<i>numeric matrix A</i>, <i>numeric matrix B</i>, <i>real scalar tol</i>)</code>
<i>void</i>	<code>_cholsolve(<i>numeric matrix A</i>, <i>numeric matrix B</i>)</code>
<i>void</i>	<code>_cholsolve(<i>numeric matrix A</i>, <i>numeric matrix B</i>, <i>real scalar tol</i>)</code>

## Description

`cholsolve( $A$ ,  $B$ )` solves  $AX = B$  and returns  $X$  for symmetric ([Hermitian](#)), positive-definite  $A$ . `cholsolve()` returns a matrix of missing values if  $A$  is not positive definite or if  $A$  is singular.

`cholsolve( $A$ ,  $B$ ,  $tol$ )` does the same thing; it allows you to specify the tolerance for declaring that  $A$  is singular; see [Tolerance](#) under *Remarks* below.

`_cholsolve( $A$ ,  $B$ )` and `_cholsolve( $A$ ,  $B$ ,  $tol$ )` do the same thing except that, rather than returning the solution  $X$ , they overwrite  $B$  with the solution, and in the process of making the calculation, they destroy the contents of  $A$ .

## Remarks

The above functions solve  $AX = B$  via Cholesky decomposition and are accurate. When  $A$  is not symmetric and positive definite, [\[M-5\] lusolve\(\)](#), [\[M-5\] qrsolve\(\)](#), and [\[M-5\] svsolve\(\)](#) are alternatives based on the LU decomposition, the QR decomposition, and the singular value decomposition (SVD). The alternatives differ in how they handle singular  $A$ . Then the LU-based routines return missing values, whereas the QR-based and SVD-based routines return generalized (least-squares) solutions.

Remarks are presented under the following headings:

[Derivation](#)  
[Relationship to inversion](#)  
[Tolerance](#)

## Derivation

We wish to solve for  $X$

$$AX = B \tag{1}$$

when  $A$  is symmetric and positive definite. Perform the Cholesky decomposition of  $A$  so that we have  $A = GG'$ . Then (1) can be written

$$GG'X = B \tag{2}$$

Define

$$Z = G'X \quad (3)$$

Then (2) can be rewritten

$$GZ = B \quad (4)$$

It is easy to solve (4) for  $Z$  because  $G$  is a lower-triangular matrix. Once  $Z$  is known, it is easy to solve (3) for  $X$  because  $G'$  is upper triangular.

## Relationship to inversion

See [Relationship to inversion](#) in [M-5] [lusolve\(\)](#) for a discussion of the relationship between solving the linear system and matrix inversion.

## Tolerance

The default tolerance used is

$$\eta = \frac{(1e-13)*\text{trace}(\text{abs}(G))}{n}$$

where  $G$  is the lower-triangular Cholesky factor of  $A$ :  $n \times n$ .  $A$  is declared to be singular if [cholesky\(\)](#) (see [M-5] [cholesky\(\)](#)) finds that  $A$  is not positive definite, or if  $A$  is found to be positive definite, if any diagonal element of  $G$  is less than or equal to  $\eta$ . Mathematically, positive definiteness implies that the matrix is not singular. In the numerical method used, two checks are made: [cholesky\(\)](#) makes one and then the  $\eta$  rule is applied to ensure numerical stability in the use of the result [cholesky\(\)](#) returns.

If you specify  $tol > 0$ , the value you specify is used to multiply  $\eta$ . You may instead specify  $tol \leq 0$  and then the negative of the value you specify is used in place of  $\eta$ ; see [M-1] [tolerance](#).

See [M-5] [lusolve\(\)](#) for a detailed discussion of the issues surrounding solving nearly singular systems. The main point to keep in mind is that if  $A$  is ill conditioned, then small changes in  $A$  or  $B$  can lead to radically large differences in the solution for  $X$ .

## Conformability

[cholsolve](#)( $A$ ,  $B$ ,  $tol$ ):

*input:*

$A$ :	$n \times n$	
$B$ :	$n \times k$	
$tol$ :	$1 \times 1$	(optional)
<i>result:</i>	$n \times k$	

[\\_cholsolve](#)( $A$ ,  $B$ ,  $tol$ ):

*input:*

$A$ :	$n \times n$	
$B$ :	$n \times k$	
$tol$ :	$1 \times 1$	(optional)

*output:*

$A$ :	$0 \times 0$
$B$ :	$n \times k$

## Diagnostics

`cholsolve(A, B, ...)`, and `_cholsolve(A, B, ...)` return a result of all missing values if  $A$  is not positive definite or if  $A$  contains missing values.

`_cholsolve(A, B, ...)` also aborts with error if  $A$  or  $B$  is a view.

All functions use the elements from the lower triangle of  $A$  without checking whether  $A$  is symmetric or, in the complex case, Hermitian.

## Also see

[M-5] `cholesky()` — Cholesky square-root decomposition

[M-5] `cholinv()` — Symmetric, positive-definite matrix inversion

[M-5] `solverlower()` — Solve  $AX=B$  for  $X$ ,  $A$  triangular

[M-5] `lusolve()` — Solve  $AX=B$  for  $X$  using LU decomposition

[M-5] `qrsolve()` — Solve  $AX=B$  for  $X$  using QR decomposition

[M-5] `svsolve()` — Solve  $AX=B$  for  $X$  using singular value decomposition

[M-5] `solve_tol()` — Tolerance used by solvers and inverters

[M-4] `matrix` — Matrix functions

[M-4] `solvers` — Functions to solve  $AX=B$  and to obtain  $A$  inverse

# Title

[M-5] **comb()** — Combinatorial function

## Syntax

*real matrix*   `comb`(*real matrix* *n*, *real matrix* *k*)

## Description

`comb(n, k)` returns the elementwise combinatorial function *n*-choose-*k*, the number of ways to choose *k* items from *n* items, regardless of order.

## Conformability

`comb(n, k)`:

<i>n</i> :	$r_1 \times c_1$
<i>k</i> :	$r_2 \times c_2$ , <i>n</i> and <i>k</i> r-conformable
<i>result</i> :	$\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

`comb(n, k)` returns missing when either argument is missing or when the result would be larger than  $10^{300}$ .

## Also see

[\[M-4\] \*\*statistical\*\*](#) — Statistical functions

# Title

[M-5] cond() — Condition number

## Syntax

*real scalar*    `cond(numeric matrix A)`

*real scalar*    `cond(numeric matrix A, real scalar p)`

## Description

`cond(A)` returns `cond(A, 2)`.

`cond(A, p)` returns the value of the condition number of *A* for the specified [norm](#) *p*, where *p* may be 0, 1, 2, or . (missing).

## Remarks

The condition number of a matrix *A* is

$$cond = \text{norm}(A, p) \times \text{norm}(A^{-1}, p)$$

These functions return missing when *A* is singular.

Values near 1 indicate that the matrix is well conditioned, and large values indicate ill conditioning.

## Conformability

`cond(A)`:  
    *A*:         $r \times c$   
    *result*:     $1 \times 1$

`cond(A, p)`:  
    *A*:         $r \times c$   
    *p*:         $1 \times 1$   
    *result*:     $1 \times 1$

## Diagnostics

`cond(A, p)` aborts with error if *p* is not 0, 1, 2, or . (missing).

`cond(A)` and `cond(A, p)` return missing when *A* is singular or if *A* contains missing values.

`cond(A)` and `cond(A, p)` return 1 when *A* is void.

`cond(A)` and `cond(A, 2)` return missing if the SVD algorithm fails to converge, which is highly unlikely; see [\[M-5\] svd\(\)](#).

---

## Also see

[M-5] [`norm\(\)`](#) — Matrix and vector norms

[M-4] [`matrix`](#) — Matrix functions

# Title

[M-5] conj() — Complex conjugate

## Syntax

*numeric matrix*    conj(*numeric matrix Z*)  
*void*                \_conj(*numeric matrix A*)

## Description

conj(*Z*) returns the elementwise complex conjugate of *Z*, that is,  $\text{conj}(a+bi) = a - bi$ . conj() may be used with real or complex matrices. If *Z* is real, *Z* is returned unmodified.

\_conj(*A*) replaces *A* with conj(*A*). Coding \_conj(*A*) is equivalent to coding *A* = conj(*A*), except that less memory is used.

## Remarks

Given  $m \times n$  matrix *Z*, conj(*Z*) returns an  $m \times n$  matrix; it does not return the transpose. To obtain the conjugate transpose matrix, also known as the adjoint matrix, adjugate matrix, Hermitian adjoint, or Hermitian transpose, code

$$Z'$$

See [M-2] [op\\_transpose](#).

A matrix equal to its conjugate transpose is called Hermitian or self-adjoint, although in this manual, we often use the term symmetric.

## Conformability

conj(*Z*):  
    *Z*:         $r \times c$   
    *result*:     $r \times c$

\_conj(*A*):  
    *input*:  
        *A*:         $r \times c$   
    *output*:  
        *A*:         $r \times c$



## Diagnostics

`conj(Z)` returns a real matrix if  $Z$  is real and a complex matrix if  $Z$  is complex.

`conj(Z)`, if  $Z$  is real, returns  $Z$  itself and not a copy. This makes `conj()` execute instantly when applied to real matrices.

`_conj(A)` does nothing if  $A$  is real (and hence, does not abort if  $A$  is a view).

## Also see

[M-5] `_transpose()` — Transposition in place

[M-4] `scalar` — Scalar mathematical functions

# Title

[M-5] **corr()** — Make correlation matrix from variance matrix

## Syntax

```
real matrix  corr(real matrix V)

void         _corr(real matrix V)
```

## Description

`corr(V)` returns the correlation matrix corresponding to variance matrix  $V$ .  
`_corr(V)` changes the contents of  $V$  from being a variance matrix to being a correlation matrix.

## Remarks

See function `variance()` in [M-5] **mean()** for obtaining a variance matrix from data.

## Conformability

```
corr(V):
  input:
      V:       $k \times k$ 
  result:     $k \times k$ 

_corr(V):
  input:
      V:       $k \times k$ 
  output:
      V:       $k \times k$ 
```

## Diagnostics

`corr()` and `_corr()` abort with error if  $V$  is not square.  $V$  should also be symmetric, but this is not checked.

## Also see

- [M-5] **mean()** — Means, variances, and correlations
- [M-4] **statistical** — Statistical functions

## Syntax

```

real matrix  cross(X, Z)

real matrix  cross(X, w, Z)

real matrix  cross(X, xc, Z, zc)

real matrix  cross(X, xc, w, Z, zc)
    
```

where

```

X:      real matrix X
xc:     real scalar xc
w:      real vector w
Z:      real matrix Z
zc:     real scalar zc
    
```

## Description

`cross()` makes calculations of the form

```

      X'X
      X'Z
      X' diag(w) X
      X' diag(w) Z
    
```

`cross()` is designed for making calculations that often arise in statistical formulas. In one sense, `cross()` does nothing that you cannot easily write out in standard matrix notation. For instance, `cross(X, Z)` calculates  $X'Z$ . `cross()`, however, has the following differences and advantages over the standard matrix-notation approach:

1. `cross()` omits the rows in  $X$  and  $Z$  that contain missing values, which amounts to dropping observations with missing values.
2. `cross()` uses less memory and is especially efficient when used with views.
3. `cross()` watches for special cases and makes calculations in those special cases more efficiently. For instance, if you code `cross(X, X)`, `cross()` observes that the two matrices are the same and makes the calculation for a symmetric matrix result.

`cross(X, Z)` returns  $X'Z$ . Usually `rows(X)==rows(Z)`, but  $X$  is also allowed to be a scalar, which is then treated as if `J(rows(Z), 1, 1)` were specified. Thus `cross(1, Z)` is equivalent to `colsum(Z)`.

`cross(X, w, Z)` returns  $X'\text{diag}(w)Z$ . Usually, `rows(w)==rows(Z)` or `cols(w)==rows(Z)`, but  $w$  is also allowed to be a scalar, which is treated as if `J(rows(Z), 1, w)` were specified. Thus `cross(X, 1, Z)` is the same as `cross(X, Z)`.  $Z$  may also be a scalar, just as in the two-argument case.

`cross(X, xc, Z, zc)` is similar to `cross(X, Z)` in that  $X'Z$  is returned. In the four-argument case, however,  $X$  is augmented on the right with a column of 1s if  $xc \neq 0$  and  $Z$  is similarly augmented if  $zc \neq 0$ . `cross(X, 0, Z, 0)` is equivalent to `cross(X, Z)`.  $Z$  may be specified as a scalar.

`cross(X, xc, w, Z, zc)` is similar to `cross(X, w, Z)` in that  $X'\text{diag}(w)Z$  is returned. As with the four-argument `cross()`,  $X$  is augmented on the right with a column of 1s if  $xc \neq 0$  and  $Z$  is similarly augmented if  $zc \neq 0$ . Both  $Z$  and  $w$  may be specified as scalars. `cross(X, 0, 1, Z, 0)` is equivalent to `cross(X, Z)`.

## Remarks

In the following examples, we are going to calculate linear regression coefficients using  $b = (X'X)^{-1}X'y$ , which means using  $\sum x/n$ , and variances using  $n/(n-1) \times (\sum x^2/n - \text{mean}^2)$ . See [M-5] `crossdev()` for examples of the same calculations made in a more numerically stable way.

The examples use the automobile data. Since we are using the absolute form of the calculation equations, it would be better if all variables had values near 1 (in which case the absolute form of the calculation equations are perfectly adequate). Thus we suggest

```
. sysuse auto
. replace weight = weight/1000
```

Some of the examples use a weight  $w$ . For that, you might try

```
. gen w = int(4*runiform())+1
```

### ► Example 1: Linear regression, the traditional way

```
: y = X = .
: st_view(y, ., "mpg")
: st_view(X, ., "weight foreign")
:
: X = X, J(rows(X),1,1)
: b = invsym(X'X)*X'y
```

**Comments:** Does not handle missing values and uses much memory if  $X$  is large.



### ► Example 2: Linear regression using `cross()`

```
: y = X = .
: st_view(y, ., "mpg")
: st_view(X, ., "weight foreign")
:
: XX = cross(X,1 , X,1)
: Xy = cross(X,1 , y,0)
: b = invsym(XX)*Xy
```

**Comments:** There is still an issue with missing values; `mpg` might not be missing everywhere `weight` and `foreign` are missing.



### ► Example 3: Linear regression using cross() and one view

```

: // We will form
: //
: //      (y' X)'(y X) = (y'y, y'X \ X'y, X'X)
:
: M = .
: st_view(M, ., "mpg weight foreign", 0)
:
: CP = cross(M, 1, M, 1)
: XX = CP[|2, 2 \ ., .|]
: Xy = CP[|2, 1 \ ., 1|]
: b = invsym(XX)*Xy

```

**Comments:** Using one view handles all missing-value issues (we specified fourth argument 0 to `st_view()`; see [M-5] [st\\_view\(\)](#)).

◀

### ► Example 4: Linear regression using cross() and subviews

```

: M = X = y = .
: st_view(M, ., "mpg weight foreign", 0)
: st_subview(y, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: XX = cross(X, 1, X, 1)
: Xy = cross(X, 1, y, 0)
: b = invsym(XX)*Xy

```

**Comments:** Using subviews also handles all missing-value issues; see [M-5] [st\\_subview\(\)](#). The subview approach is a little less efficient than the previous solution but is perhaps easier to understand. The efficiency issue concerns only the extra memory required by the subviews `y` and `X`, which is not much.

Also, this subview solution could be used to handle the missing-value problems of calculating linear regression coefficients the traditional way, shown in [example 1](#):

```

: M = X = y = .
: st_view(M, ., "mpg weight foreign", 0)
: st_subview(y, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: X = X, J(rows(X), 1, 1)
: b = invsym(X'X)*X'y

```

◀

### ► Example 5: Weighted linear regression, the traditional way

```

: M = w = y = X = .
: st_view(M, ., "w mpg weight foreign", 0)
: st_subview(w, M, ., 1)
: st_subview(y, M, ., 2)
: st_subview(X, M, ., (3\..))
:
: X = X, J(rows(X), 1, 1)
: b = invsym(X'diag(w)*X)*X'diag(w)'y

```

**Comments:** The memory requirements are now truly impressive because `diag(w)` is an  $N \times N$  matrix! That is, the memory requirements are truly impressive when  $N$  is large. Part of the power of Mata is that you can write things like `invsym(X'diag(w)*X)*X'diag(w)'y` and obtain solutions. We do not mean to be dismissive of the traditional approach; we merely wish to emphasize its memory requirements and note that there are alternatives.

◀

### ► Example 6: Weighted linear regression using `cross()`

```
: M = w = y = X = .
: st_view(M, ., "w mpg weight foreign", 0)
: st_subview(w, M, ., 1)
: st_subview(y, M, ., 2)
: st_subview(X, M, ., (3\..))
:
: XX = cross(X,1 ,w, X,1)
: Xy = cross(X,1 ,w, y,0)
: b = invsym(XX)*Xy
```

**Comments:** The memory requirements here are no greater than they were in [example 4](#), which this example closely mirrors. We could also have mirrored the logic of [example 3](#):

```
: M = w = M2 = .
: st_view(M, ., "w mpg weight foreign", 0)
: st_subview(w, M, ., 1)
: st_subview(M2, M, ., (2\..))
:
: CP = cross(M,1 ,w, M,1)
: XX = CP[1,3 \ .,.]
: Xy = CP[1,3 \ .,2]
: b = invsym(XX)*Xy
```

Note how similar these solutions are to their unweighted counterparts. The only important difference is the appearance of `w` as the middle argument of `cross()`. Since specifying the middle argument as a scalar 1 is also allowed and produces unweighted estimates, the above code could be modified to produce unweighted or weighted estimates, depending on how `w` is defined.

◀

### ► Example 7: Mean of one variable

```
: x = .
: st_view(x, ., "mpg", 0)
:
: CP = cross(1,0 , x,1)
: mean = CP[1]/CP[2]
```

**Comments:** An easier and every bit as good a solution would be

```
: x = .
: st_view(x, ., "mpg", 0)
:
: mean = mean(x,1)
```

`mean()` (see [\[M-5\] mean\(\)](#)) is implemented in terms of `cross()`. Actually, `mean()` is implemented using the quad-precision version of `cross()`; see [\[M-5\] quadcross\(\)](#). We could implement our solution in terms of `quadcross()`:

```

: x = .
: st_view(x, ., "mpg", 0)
:
: CP = quadcross(1,0 , x,1)
: mean = CP[1]/CP[2]

```

quadcross() returns a double-precision result just as does cross(). The difference is that quadcross() uses quad precision internally in calculating sums.



### ► Example 8: Means of multiple variables

```

: X = .
: st_view(X, ., "mpg weight displ", 0)
:
: CP = cross(1,0 , X,1)
: n = cols(CP)
: means = CP[|1\n-1|] :/ CP[n]

```

**Comments:** The above logic will work for one variable, too. With mean(), the solution would be

```

: X = .
: st_view(X, ., "mpg weight displ", 0)
:
: means = mean(X, 1)

```



### ► Example 9: Weighted means of multiple variables

```

: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: CP = cross(1,0, w, X,1)
: n = cols(CP)
: means = CP[|1\n-1|] :/ CP[n]

```

**Comments:** Note how similar this solution is to the unweighted solution: w now appears as the middle argument of cross(). The line CP = cross(1,0, w, X,1) could also be coded CP = cross(w,0, X,1); it would make no difference.

The mean() solution to the problem is

```

: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: means = mean(X, w)

```



### ► Example 10: Variance matrix, traditional approach 1

```
: X = .
: st_view(X, .., "mpg weight displ", 0)
:
: n      = rows(X)
: means  = mean(X, 1)
: cov    = (X'X/n - means'means)*(n/(n-1))
```

**Comments:** This above is not 100% traditional since we used `mean()` to obtain the means, but that does make the solution more understandable. The solution requires calculating  $X'$ , requiring that the data matrix be duplicated. Also, we have used a numerically poor calculation formula.

◀

### ► Example 11: Variance matrix, traditional approach 2

```
: X = .
: st_view(X, .., "mpg weight displ", 0)
:
: n      = rows(X)
: means  = mean(X, 1)
: cov    = (X:-means)'(X:-means) :/ (n-1)
```

**Comments:** We use a better calculation formula and, in the process, increase our memory usage substantially.

◀

### ► Example 12: Variance matrix using cross()

```
: X = .
: st_view(X, .., "mpg weight displ", 0)
:
: n      = rows(X)
: means  = mean(X, 1)
: XX     = cross(X, X)
: cov    = ((XX:/n)-means'means)*(n/(n-1))
```

**Comments:** The above solution conserves memory but uses the numerically poor calculation formula. A related function, `crossdev()`, will calculate deviation crossproducts:

```
: X = .
: st_view(X, .., "mpg weight displ", 0)
:
: n      = rows(X)
: means  = mean(X, 1)
: xx     = crossdev(X,means, X,means)
: cov    = xx:/(n-1)
```

See [M-5] `crossdev()`. The easiest solution, however, is

```
: X = .
: st_view(X, .., "mpg weight displ", 0)
:
: cov = variance(X, 1)
```



See [M-5] `mean()` for a description of the `variance()` function. `variance()` is implemented in terms of `crossdev()`.



### ► Example 13: Weighted variance matrix, traditional approaches

```
: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: n      = colsum(w)
: means = mean(X, w)
: cov    = (X'diag(w)*X:/n - means'means)*(n/(n-1))
```

**Comments:** Above we use the numerically poor formula. Using the better deviation formula, we would have

```
: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: n      = colsum(w)
: means = mean(X, w)
: cov    = (X:-means)'diag(w)*(X:-means) :/ (n-1)
```

The memory requirements include making a copy of the data with the means removed and making an  $N \times N$  diagonal matrix.



### ► Example 14: Weighted variance matrix using `cross()`

```
: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: n      = colsum(w)
: means = mean(X, w)
: cov    = (cross(X,w,X):/n - means'means)*(n/(n-1))
```

**Comments:** As in [example 12](#), the above solution conserves memory but uses a numerically poor calculation formula. Better is to use `crossdev()`:

```
: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: n      = colsum(w)
: means = mean(X, w)
: cov    = crossdev(X,means, w, X,means) :/ (n-1)
```

and easiest is to use `variance()`:

```
: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: cov = variance(X, w)
```

See [M-5] `crossdev()` and [M-5] `mean()`.



## Comment concerning `cross()` and missing values

`cross()` automatically omits rows containing missing values in making its calculation. Depending on this feature, however, is considered bad style because so many other Mata functions do not provide that feature and it is easy to make a mistake.

The right way to handle missing values is to exclude them when constructing views and subviews, as we have done above. When we constructed a view, we invariably specified fourth argument 0 to `st_view()`. In formal programming situations, you will probably specify the name of the `touse` variable you have previously constructed in your `ado`-file that calls your Mata function.

## Conformability

`cross(X, xc, w, Z, zc):`

<i>X</i> :	$n \times v_1$	or	$1 \times 1$ ,	$1 \times 1$ treated as if $n \times 1$
<i>xc</i> :	$1 \times 1$			(optional)
<i>w</i> :	$n \times 1$	or	$1 \times n$	or $1 \times 1$ (optional)
<i>Z</i> :	$n \times v_2$			
<i>zc</i> :	$1 \times 1$			(optional)
<i>result</i> :	$(v_1 + (xc \neq 0)) \times (v_2 + (zc \neq 0))$			

## Diagnostics

`cross(X, xc, w, Z, zc)` omits rows in *X* and *Z* that contain missing values.

## Also see

[M-5] `crossdev()` — Deviation cross products

[M-5] `quadcross()` — Quad-precision cross products

[M-5] `mean()` — Means, variances, and correlations

[M-4] `statistical` — Statistical functions

[M-4] `utility` — Matrix utility functions

## Syntax

*real matrix*    `crossdev(X, x, Z, z)`

*real matrix*    `crossdev(X, x, w, Z, z)`

*real matrix*    `crossdev(X, xc, x, Z, zc, z)`

*real matrix*    `crossdev(X, xc, x, w, Z, zc, z)`

where

*X*:    *real matrix X*  
*xc*:    *real scalar xc*  
*x*:    *real rowvector x*  
  
*w*:    *real vector w*  
  
*Z*:    *real matrix Z*  
*zc*:    *real scalar zc*  
*z*:    *real rowvector z*

## Description

`crossdev()` makes calculations of the form

$$(X: -x)' (X: -x)$$

$$(X: -x)' (Z: -z)$$

$$(X: -x)' \text{diag}(w) (X: -x)$$

$$(X: -x)' \text{diag}(w) (Z: -z)$$

`crossdev()` is a variation on [\[M-5\] cross\(\)](#). `crossdev()` mirrors `cross()` in every respect except that it has two additional arguments: *x* and *z*. *x* and *z* record the amount by which *X* and *Z* are to be deviated. *x* and *z* usually contain the (appropriately weighted) column means of *X* and *Z*.

## Remarks

*x* usually contains the same number of rows as *X* but, if *xc*  $\neq$  0, *x* may contain an extra element on the right recording the amount from which the constant 1 should be deviated.

The same applies to *z*: it usually contains the same number of rows as *Z* but, if *zc*  $\neq$  0, *z* may contain an extra element on the right.

### ► Example 1: Linear regression using one view

```
: M = .
: st_view(M, ., "mpg weight foreign", 0)
:
: means = mean(M, 1)
: CP = crossdev(M, means, M, means)
: xx = CP[|2,2 \ .,|.]
: xy = CP[|2,1 \ .,|.1]
: b = invsym(xx)*xy
: b = b \ means[1]-means[|2\|.]*b
```

Compare this solution with [example 3](#) in [M-5] `cross()`.



### ► Example 2: Linear regression using subviews

```
: M = X = y = .
: st_view(M, ., "mpg weight foreign", 0)
: st_subview(y, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: xmean = mean(X, 1)
: ymean = mean(y, 1)
: xx = crossdev(X, xmean, X, xmean)
: xy = crossdev(X, xmean, y, ymean)
: b = invsym(xx)*xy
: b = b \ ymean-xmean*b
```

Compare this solution with [example 4](#) in [M-5] `cross()`.



### ► Example 3: Weighted linear regression

```
: M = X = y = w = .
: st_view(M, ., "w mpg weight foreign", 0)
: st_subview(w, M, ., 1)
: st_subview(y, M, ., 2)
: st_subview(X, M, ., (3\..))
:
: xmean = mean(X, w)
: ymean = mean(y, w)
: xx = crossdev(X, xmean, w, X, xmean)
: xy = crossdev(X, xmean, w, y, ymean)
: b = invsym(xx)*xy
: b = b \ ymean-xmean*b
```

Compare this solution with [example 6](#) in [M-5] `cross()`.



### ► Example 4: Variance matrix

```
: X = .
: st_view(X, ., "mpg weight displ", 0)
:
: n      = rows(X)
: means = mean(X, 1)
: xx     = crossdev(X, means, X, means)
: cov    = xx / (n-1)
```

This is exactly what `variance()` does; see [M-5] [mean\(\)](#). Compare this solution with [example 12](#) in [M-5] [cross\(\)](#).

◀

### ► Example 5: Weighted variance matrix

```
: M = w = X = .
: st_view(M, ., "w mpg weight displ", 0)
: st_subview(w, M, ., 1)
: st_subview(X, M, ., (2\..))
:
: n      = colsum(w)
: means = mean(X, w)
: cov    = crossdev(X, means, w, X, means) :/ (n-1)
```

This is exactly what `variance()` does with weighted data; see [M-5] [mean\(\)](#). Compare this solution with [example 14](#) in [M-5] [cross\(\)](#).

◀

## Conformability

`crossdev(X, xc, x, w, Z, zc, z):`

<i>X</i> :	$n \times v_1$	or	$1 \times 1$ ,	$1 \times 1$ treated as if $n \times 1$
<i>xc</i> :	$1 \times 1$			(optional)
<i>x</i> :	$1 \times v_1$	or	$1 \times v_1 + (xc \neq 0)$	
<i>w</i> :	$n \times 1$	or	$1 \times n$ or $1 \times 1$	(optional)
<i>Z</i> :	$n \times v_2$			
<i>zc</i> :	$1 \times 1$			(optional)
<i>z</i> :	$1 \times v_2$	or	$1 \times v_2 + (zc \neq 0)$	
<i>result</i> :	$(v_1 + (xc \neq 0)) \times (v_2 + (zc \neq 0))$			

## Diagnostics

`crossdev(X, xc, x, w, Z, zc, z)` omits rows in *X* and *Z* that contain missing values.

## Also see

[M-5] `cross()` — Cross products

[M-5] `quadcross()` — Quad-precision cross products

[M-4] `utility` — Matrix utility functions

[M-4] `statistical` — Statistical functions

## Title

**[M-5] cvpermute()** — Obtain all permutations

## Syntax

```
info = cvpermutesetup(real colvector V [ , real scalar unique ])  
real colvector cvpermute(info)
```

where *info* should be declared *transmorphic*.

## Description

`cvpermute()` returns all permutations of the values of column vector *V*, one at a time. If  $V = (1\backslash 2\backslash 3)$ , there are six permutations and they are  $(1\backslash 2\backslash 3)$ ,  $(1\backslash 3\backslash 2)$ ,  $(2\backslash 1\backslash 3)$ ,  $(2\backslash 3\backslash 1)$ ,  $(3\backslash 1\backslash 2)$ , and  $(3\backslash 2\backslash 1)$ . If  $V = (1\backslash 2\backslash 1)$ , there are three permutations and they are  $(1\backslash 1\backslash 2)$ ,  $(1\backslash 2\backslash 1)$ , and  $(2\backslash 1\backslash 1)$ .

Vector *V* is specified by calling `cvpermutesetup()`,

```
info = cvpermutesetup(V)
```

*info* holds information that is needed by `cvpermute()` and it is *info*, not *V*, that is passed to `cvpermute()`. To obtain the permutations, repeated calls are made to `cvpermute()` until it returns `J(0,1,.):`

```
info = cvpermutesetup(V)  
while ((p=cvpermute(info)) != J(0,1,.)) {  
    ... p ...  
}
```

Column vector *p* will contain a permutation of *V*.

`cvpermutesetup()` may be specified with one or two arguments:

```
info = cvpermutesetup(V)
```

```
info = cvpermutesetup(V, unique)
```

*unique* is usually not specified. If *unique* is specified, it should be 0 or 1. Not specifying *unique* is equivalent to specifying *unique* = 0. Specifying *unique* = 1 states that the elements of *V* are unique or, at least, are to be treated that way.

When the arguments of *V* are unique—for instance,  $V = (1\backslash 2\backslash 3)$ —specifying *unique* = 1 will make `cvpermute()` run faster. The same permutations will be returned, although usually in a different order.

When the arguments of *V* are not unique—for instance,  $V = (1\backslash 2\backslash 1)$ —specifying *unique* = 1 will make `cvpermute()` treat them as if they were unique. With *unique* = 0, there are three permutations of  $(1\backslash 2\backslash 1)$ . With *unique* = 1, there are six permutations, just as there are with  $(1\backslash 2\backslash 3)$ .

## Remarks

### ► Example 1

You have the following data:

v1	v2
22	29
17	33
21	26
20	32
16	35

You wish to do an exact permutation test for the correlation between v1 and v2.

That is, you wish to (1) calculate the correlation between v1 and v2—call that value  $r$ —and then (2) calculate the correlation between v1 and v2 for all permutations of v1, and count how many times the result is more extreme than  $r$ .

For the first step,

```
: X = (22, 29 \  
>      17, 33 \  
>      21, 26 \  
>      20, 32 \  
>      16, 35)  
:  
: correlation(X)  
[symmetric]
```

	1	2
1	1	
2	-.8468554653	1

The correlation is  $-.846855$ . For the second step,

```
: V1 = X[,1]  
: V2 = X[,2]  
: num = den = 0  
: info = cvpermutesetup(V1)  
: while ((V1=cvpermute(info)) != J(0,1,.)) {  
>     rho = correlation((V1,V2))[2,1]  
>     if (rho<=-.846 | rho>=.846) num++  
>     den++  
> }  
: (num, den, num/den)
```

	1	2	3
1	13	120	.1083333333

Of the 120 permutations, 13 (10.8%) were outside .846855 or  $-.846855$ .



## ► Example 2

You now wish to do the same thing but using the Spearman rank-correlation coefficient. Mata has no function that will calculate that, but Stata has a command that does—see [R] [spearman](#)—so we will use the Stata command as our subroutine.

This time, we will assume that the data have been loaded into a Stata dataset:

```
. list
```

	var1	var2
1.	22	29
2.	17	33
3.	21	26
4.	20	32
5.	16	35

For the first step,

```
. spearman var1 var2
Number of obs =      5
Spearman's rho =   -0.9000
Test of Ho: var1 and var2 are independent
Prob > |t| =      0.0374
```

For the second step,

```
. mata
----- mata (type end to exit) -----
: V1 = st_data(., "var1")
: info = cvpermutesetup(V1)
: num = den = 0
: while ((V1=cvpermute(info)) != J(0,1,.)) {
>   st_store(., "var1", V1)
>   stata("quietly spearman var1 var2")
>   rho = st_numscalar("r(rho)")
>   if (rho<=-.9 | rho>=.9) num++
>   den++
> }
: (num, den, num/den)
               1               2               3
1  

|   |     |            |
|---|-----|------------|
| 2 | 120 | .016666667 |
|---|-----|------------|


```

Only two of the permutations resulted in a rank correlation of at least .9 in magnitude.

In the code above, we obtained the rank correlation from `r(rho)` which, we learned from [R] [spearman](#), is where `spearman` stores it.

Also note how we replaced the contents of `var1` by using `st_store()`. Our code leaves the dataset changed and so could be improved.

## Conformability

cvpermutesetup(*V*, *unique*):

<i>V</i> :	$n \times 1$	
<i>unique</i> :	$1 \times 1$	(optional)
<i>result</i> :	$1 \times L$	

cvpermute(*info*):

<i>info</i> :	$1 \times L$
<i>result</i> :	$n \times 1$ or $0 \times 1$

where

$$L = \begin{cases} 3 & \text{if } n = 0 \\ 4 & \text{if } n = 1 \\ (n + 3)(n + 2)/2 - 6 & \text{otherwise} \end{cases}$$

The value of *L* is not important except that the *info* vector returned by cvpermutesetup() and then passed to cvpermute() consumes memory. For instance,

<i>n</i>	<i>L</i>	Total memory ( $8 * L$ )
5	22	176 bytes
10	72	576
50	1,372	10,560
100	5,247	41,976
1,000	502,497	4,019,976

## Diagnostics

cvpermute() returns J(0,1,.) when there are no more permutations.

## Also see

[M-4] [statistical](#) — Statistical functions

## Syntax

```

tc = clock(strdatetime, pattern [ , year ])
tc = mdyhms(month, day, year, hour, minute, second)
tc = dhms(td, hour, minute, second)
tc = hms(hour, minute, second)
hour = hh(tc)
minute = mm(tc)
second = ss(tc)
td = dofC(tc)

tC = CofC(tc)
tC = Clock(strdatetime, pattern [ , year ])
tC = Cmdyhms(month, day, year, hour, minute, second)
tC = Cdhms(td, hour, minute, second)
tC = Chms(hour, minute, second)
hour = hhC(tC)
minute = mmC(tC)
second = ssC(tC)
td = dofC(tC)
tc = cofC(tC)

td = date(strdate, dpattern [ , year ])
td = mdy(month, day, year)
tw = yw(year, week)
tm = ym(year, month)
tq = yq(year, quarter)
th = yh(year, half)
tc = cofd(td)
tC = Cofd(td)

```

```
td = dofb(tb, "calendar")
```

```
tb = bofd("calendar", td)
```

```
month = month(td)
```

```
day = day(td)
```

```
year = year(td)
```

```
dow = dow(td)
```

```
week = week(td)
```

```
quarter = quarter(td)
```

```
half = halfyear(td)
```

```
doy = doy(td)
```

```
ty = yearly(strydate, ypattern [, year])
```

```
ty = yofd(td)
```

```
td = dofy(ty)
```

```
th = halfyearly(strhdate, hpattern [, year])
```

```
th = hofd(td)
```

```
td = dofh(th)
```

```
tq = quarterly(strqdate, qpattern [, year])
```

```
tq = qofd(td)
```

```
td = dofq(tq)
```

```
tm = monthly(strmdate, mpattern [, year])
```

```
tm = mofd(td)
```

```
td = dofm(tm)
```

```
tw = weekly(strwdate, wpattern [, year])
```

```
tw = wofd(td)
```

```
td = dofww(tw)
```

```

hours = hours(ms)
minutes = minutes(ms)
seconds = seconds(ms)
ms = msofhours(hours)
ms = msofminutes(minutes)
ms = msofseconds(seconds)

```

where

<i>tc</i> :	number of milliseconds from 01jan1960 00:00:00.000, unadjusted for leap seconds
<i>tC</i> :	number of milliseconds from 01jan1960 00:00:00.000, adjusted for leap seconds
<i>strdatetime</i> :	string-format date, time, or date/time, e.g., "15jan1992", "1/15/1992", "15-1-1992", "January 15, 1992", "9:15", "13:42", "1:42 p.m.", "1:42:15.002 pm", "15jan1992 9:15", "1/15/1992 13:42", "15-1-1992 1:42 p.m.", "January 15, 1992 1:42:15.002 pm"
<i>pattern</i> :	order of month, day, year, hour, minute, and seconds in <i>strdatetime</i> , plus optional default century, e.g., "DMYhms" (meaning day, month, year, hour, minute, second), "DMYhm", "MDYhm", "hmMDY", "hms", "hm", "MDY", "MD19Y", "MDY20Yhm"
<i>td</i> :	number of days from 01jan1960
<i>tb</i> :	business date (days)
<i>calendar</i> :	string scalar containing calendar name or %tb format
<i>strdate</i> :	string-format date, e.g., "15jan1992", "1/15/1992", "15-1-1992", "January 15, 1992"
<i>dpattern</i> :	order of month, day, and year in <i>strdate</i> , plus optional default century, e.g., "DMY" (meaning day, month, year), "MDY", "MD19Y"
<i>hour</i> :	hour, 0–23
<i>minute</i> :	minute, 0–59
<i>second</i> :	second, 0.000–59.999 (maximum 60.999 in case of leap second)
<i>month</i> :	month number, 1–12
<i>day</i> :	day-of-month number, 1–31
<i>year</i> :	year, e.g., 1942, 1995, 2008
<i>week</i> :	week within year, 1–52
<i>quarter</i> :	quarter within year, 1–4
<i>half</i> :	half within year, 1–2
<i>dow</i> :	day of week, 0–6, 0 = Sunday
<i>doy</i> :	day within year, 1–366

<i>ty</i> :	calendar year
<i>strydate</i> :	string-format calendar year, e.g., "1980", "80"
<i>ypattern</i> :	pattern of <i>strydate</i> , e.g., "Y", "19Y"
<i>th</i> :	number of halves from 1960h1
<i>strhdate</i> :	string-format <i>hdate</i> , e.g., "1982-1", "1982h2", "2 1982"
<i>hpattern</i> :	pattern of <i>strhdate</i> , e.g., "YH", "19YH", "HY"
<i>tq</i> :	number of quarters from 1960q1
<i>strqdate</i> :	string-format <i>qdate</i> , e.g., "1982-3", "1982q2", "3 1982"
<i>qpattern</i> :	pattern of <i>strqdate</i> , e.g., "YQ", "19YQ", "QY"
<i>tm</i> :	number of months from 1960m1
<i>strmdate</i> :	string-format <i>mdate</i> , e.g., "1982-3", "1982m2", "3/1982"
<i>mpattern</i> :	pattern of <i>strmdate</i> , e.g., "YM", "19YM", "MR"
<i>tw</i> :	number of weeks from 1960w1
<i>strwdate</i> :	string-format <i>wdate</i> , e.g., "1982-3", "1982w2", "1982-15"
<i>wpattern</i> :	pattern of <i>strwdate</i> , e.g., "YW", "19YW", "WY"
<i>hours</i> :	interval of time in hours (positive or negative, real)
<i>minutes</i> :	interval of time in minutes (positive or negative, real)
<i>seconds</i> :	interval of time in seconds (positive or negative, real)
<i>ms</i> :	interval of time in milliseconds (positive or negative, integer)

Functions return an element-by-element result. Functions are usually used with scalars.

All variables are *real matrix* except the *str\** and *\*pattern* variables, which are *string matrix*.

## Description

These functions mirror Stata’s date functions; see [\[D\] `datetime`](#).

## Conformability

`clock(strdatetime, pattern, year)`, `Clock(strdatetime, pattern, year)`:

<i>strdatetime</i> :	$r_1 \times c_1$	
<i>pattern</i> :	$r_2 \times c_2$	(c-conformable with <i>strdatetime</i> )
<i>year</i> :	$r_3 \times c_3$	(optional, c-conformable)
<i>result</i> :	$\max(r_1, r_2, r_3) \times \max(c_1, c_2, c_3)$	

`mdyhms(month, day, year, hour, minute, second)`,  
`Cmdyhms(month, day, year, hour, minute, second)`:

*month*:  $r_1 \times c_1$   
*day*:  $r_2 \times c_2$   
*year*:  $r_3 \times c_3$   
*hour*:  $r_4 \times c_4$   
*minute*:  $r_5 \times c_5$   
*second*:  $r_6 \times c_6$  (all variables c-conformable)  
*result*:  $\max(r_1, r_2, r_3, r_4, r_5, r_6) \times \max(c_1, c_2, c_3, c_4, c_5, c_6)$

`hms(hour, minute, second)`, `Chms(hour, minute, second)`:

*hour*:  $r_1 \times c_1$   
*minute*:  $r_2 \times c_2$   
*second*:  $r_3 \times c_3$   
*result*:  $\max(r_1, r_2, r_3) \times \max(c_1, c_2, c_3)$

`dhms(td, hour, minute, second)`, `Cdhms(td, hour, minute, second)`:

*td*:  $r_1 \times c_1$   
*hour*:  $r_2 \times c_2$   
*minute*:  $r_3 \times c_3$   
*second*:  $r_4 \times c_4$  (all variables c-conformable)  
*result*:  $\max(r_1, r_2, r_3, r_4) \times \max(c_1, c_2, c_3, c_4)$

`hh(x)`, `mm(x)`, `ss(x)`, `hhC(x)`, `mmC(x)`, `ssC(x)`,

*x*:  $r \times c$   
*result*:  $r \times c$

`date(strdate, dpattern, year)`:

*strdate*:  $r_1 \times c_1$   
*dpattern*:  $r_2 \times c_2$  (c-conformable with *strdate*)  
*year*:  $r_3 \times c_3$  (optional, c-conformable)  
*result*:  $\max(r_1, r_2, r_3) \times \max(c_1, c_2, c_3)$

`mdy(month, day, year)`:

*month*:  $r_1 \times c_1$   
*day*:  $r_2 \times c_2$   
*year*:  $r_3 \times c_3$  (all variables c-conformable)  
*result*:  $\max(r_1, r_2, r_3) \times \max(c_1, c_2, c_3)$

`yw(year, detail)`, `ym(year, detail)`, `yq(year, detail)`, `yh(year, detail)`:

*year*:  $r_1 \times c_1$   
*detail*:  $r_2 \times c_2$  (c-conformable with *year*)  
*result*:  $\max(r_1, r_2) \times \max(c_1, c_2)$

`month(td)`, `day(td)`, `year(td)`, `dow(td)`, `week(td)`, `quarter(td)`, `halfyear(td)`, `doy(td)`:

*td*:  $r \times c$   
*result*:  $r \times c$

`yearly(str, pat, year)`, `halfyearly(str, pat, year)`, `quarterly(str, pat, year)`,  
`monthly(str, pat, year)`, `weekly(str, pat, year)`:

*str*:  $r_1 \times c_1$   
*pat*:  $r_2 \times c_2$  (c-conformable with *str*)  
*year*:  $r_3 \times c_3$  (optional, c-conformable)  
*result*:  $\max(r_1, r_2, r_3) \times \max(c_1, c_2, c_3)$

```
Cofc(x), cofC(x), dofC(x), dofC(x), cofd(x), Cofd(x), yofd(x), dofy(x), hofd(x),
dofh(x), qofd(x), dofQ(x), mofd(x), dofM(x), wofd(x), dofW(x):
    x:      r × c
    result: r × c

dofb(tb, "calendar")
    tb:      r × c
    calendar: 1 × 1
    result:  r × c

bofd("calendar", td)
    calendar: 1 × 1
    td:      r × c
    result:  r × c

hours(x), minutes(x), seconds(x), msofhours(x), msofminutes(x): msofseconds(x):
    x:      r × c
    result:  r × c
```

Diagnostics

None.

Also see

[M-4] [scalar](#) — Scalar mathematical functions



## Syntax

```

        D = deriv_init()

(varies)      deriv_init_evaluator(D [, &function()])
(varies)      deriv_init_evaluatortype(D [, evaluatortype])
(varies)      deriv_init_params(D [, real rowvector parameters])
(varies)      deriv_init_argument(D, real scalar k [, X])
(varies)      deriv_init_narguments(D [, real scalar K])
(varies)      deriv_init_weights(D [, real colvector weights])
(varies)      deriv_init_h(D [, real rowvector h])
(varies)      deriv_init_scale(D [, real matrix scale])
(varies)      deriv_init_bounds(D [, real rowvector minmax])
(varies)      deriv_init_search(D [, search])
(varies)      deriv_init_verbose(D [, {"on" | "off"}])

(varies)      deriv(D, {0 | 1 | 2})
real scalar  _deriv(D, {0 | 1 | 2})

real scalar  deriv_result_value(D)
real vector  deriv_result_values(D)
void         _deriv_result_values(D, v)
real rowvector deriv_result_gradient(D)
void         _deriv_result_gradient(D, g)
real matrix  deriv_result_scores(D)
void         _deriv_result_scores(D, S)
real matrix  deriv_result_Jacobian(D)
void         _deriv_result_Jacobian(D, J)
real matrix  deriv_result_Hessian(D)
void         _deriv_result_Hessian(D, H)
    
```

```
real rowvector deriv_result_h(D)
real matrix    deriv_result_scale(D)
real matrix    deriv_result_delta(D)
real scalar    deriv_result_errorcode(D)
string scalar  deriv_result_errortext(D)
real scalar    deriv_result_returncode(D)

void           deriv_query(D)
```

where *D*, if it is declared, should be declared

```
transmorphic D
```

and where *evaluator**type* optionally specified in `deriv_init_evaluator`*type*() is

<i>evaluator</i> <i>type</i>	Description
"d"	<i>function</i> () returns <i>scalar</i> value
"v"	<i>function</i> () returns <i>colvector</i> value
"t"	<i>function</i> () returns <i>rowvector</i> value

The default is "d" if not set.

and where *search* optionally specified in `deriv_init_search`() is

<i>search</i>	Description
"interpolate"	use linear and quadratic interpolation to search for an optimal delta
"bracket"	use a bracketed quadratic formula to search for an optimal delta
"off"	do not search for an optimal delta

The default is "interpolate" if not set.

## Description

These functions compute derivatives of the real function *f*(*p*) at the real parameter values *p*.

`deriv_init`() begins the definition of a problem and returns *D*, a problem-description handle set that contains default values.

The `deriv_init_*`(*D*, ...) functions then allow you to modify those defaults. You use these functions to describe your particular problem: to set the identity of function *f*(), to set parameter values, and the like.

`deriv`(*D*, *todo*) then computes derivatives depending upon the value of *todo*.

`deriv`(*D*, 0) returns the function value without computing derivatives.

`deriv(D, 1)` returns the first derivatives, also known as the gradient vector for scalar-valued functions (type `d` and `v`) or the Jacobian matrix for vector-valued functions (type `t`).

`deriv(D, 2)` returns the matrix of second derivatives, also known as the Hessian matrix; the gradient vector is also computed. This syntax is not allowed for type `t` evaluators.

The `deriv_result_*(D)` functions can then be used to access other values associated with the solution.

Usually you would stop there. In other cases, you could compute derivatives at other parameter values:

```
deriv_init_params(D, p_alt)
deriv(D, todo)
```

Aside: The `deriv_init_*(D, ...)` functions have two modes of operation. Each has an optional argument that you specify to set the value and that you omit to query the value. For instance, the full syntax of `deriv_init_params()` is

```
void deriv_init_params(D, real rowvector parameters)
real rowvector deriv_init_params(D)
```

The first syntax sets the parameter values and returns nothing. The second syntax returns the previously set (or default, if not set) parameter values.

All the `deriv_init_*(D, ...)` functions work the same way.

## Remarks

Remarks are presented under the following headings:

- First example*
- Notation and formulas*
- Type d evaluators*
- Example of a type d evaluator*
- Type v evaluators*
- User-defined arguments*
- Example of a type v evaluator*
- Type t evaluators*
- Example of a type t evaluator*
- Functions*
  - `deriv_init()`
  - `deriv_init_evaluator()` and `deriv_init_evaluortype()`
  - `deriv_init_argument()` and `deriv_init_narguments()`
  - `deriv_init_weights()`
  - `deriv_init_params()`
- Advanced init functions*
  - `deriv_init_h()`, `... _scale()`, `... _bounds()`, and `... _search()`
  - `deriv_init_verbose()`
- `deriv()`
- `_deriv()`
- `deriv_result_value()`
- `deriv_result_values()` and `_deriv_result_values()`
- `deriv_result_gradient()` and `_deriv_result_gradient()`
- `deriv_result_scores()` and `_deriv_result_scores()`
- `deriv_result_Jacobian()` and `_deriv_result_Jacobian()`
- `deriv_result_Hessian()` and `_deriv_result_Hessian()`
- `deriv_result_h()`, `... _scale()`, and `... _delta()`
- `deriv_result_errorcode()`, `... _errortext()`, and `... _returncode()`
- `deriv_query()`

## First example

The derivative functions may be used interactively.

Below we use the functions to compute  $f'(x)$  at  $x = 0$ , where the function is

```
f(x) = exp(-x^2 + x - 3)
: void myeval(x, y)
> {
>     y = exp(-x^2 + x - 3)
> }
: D = deriv_init()
: deriv_init_evaluator(D, &myeval())
: deriv_init_params(D, 0)
: dydx = deriv(D, 1)
: dydx
.0497870683
: exp(-3)
.0497870684
```

The derivative, given the above function, is  $f'(x) = (-2 \times x + 1) \times \exp(-x^2 + x - 3)$ , so  $f'(0) = \exp(-3)$ .

## Notation and formulas

We wrote the above in the way that mathematicians think, that is, differentiate  $y = f(x)$ . Statisticians, on the other hand, think differentiate  $s = f(b)$ . To avoid favoritism, we will write  $v = f(p)$  and write the general problem with the following notation:

Differentiate  $v = f(p)$  with respect to  $p$ , where

$v$ : a scalar  
 $p$ :  $1 \times np$

The gradient vector is  $g = f'(p) = df/dp$ , where

$g$ :  $1 \times np$

and the Hessian matrix is  $H = f''(p) = d^2f/dpdp'$ , where

$H$ :  $np \times np$

`deriv()` can also work with vector-valued functions. Here is the notation for vector-valued functions:

Differentiate  $v = f(p)$  with respect to  $p$ , where

$v$ :  $1 \times nv$ , a vector  
 $p$ :  $1 \times np$

The Jacobian matrix is  $J = f'(p) = df/dp$ , where

$J$ :  $nv \times np$

and where

$J[i, j] = dv[i]/dp[j]$

Second-order derivatives are not computed by `deriv()` when used with vector-valued functions.

`deriv()` uses the following formula for computing the numerical derivative of  $f()$  at  $p$

$$f'(p) = \frac{f(p+d) - f(p-d)}{2d}$$

where we refer to  $d$  as the delta used for computing numerical derivatives. To search for an optimal delta, we decompose  $d$  into two parts.

$$d = h \times scale$$

By default,  $h$  is a fixed value that depends on the parameter value.

$$h = (\text{abs}(p)+1\text{e-}3)*1\text{e-}3$$

`deriv()` searches for a value of  $scale$  that will result in an optimal numerical derivative, that is, one where  $d$  is as small as possible subject to the constraint that  $f(x+d) - f(x-d)$  will be calculated to at least half the accuracy of a double-precision number. This is accomplished by searching for  $scale$  such that  $f(x+d)$  and  $f(x-d)$  fall between  $v0$  and  $v1$ , where

$$v0 = (\text{abs}(f(x))+1\text{e-}8)*1\text{e-}8$$

$$v1 = (\text{abs}(f(x))+1\text{e-}7)*1\text{e-}7$$

Use `deriv_init_h()` to change the default  $h$  values. Use `deriv_init_scale()` to change the default initial  $scale$  values. Use `deriv_init_bounds()` to change the default bounds ( $1\text{e-}8$ ,  $1\text{e-}7$ ) used for determining the optimal  $scale$ .

## Type d evaluators

You must write an evaluator function to calculate  $f()$  before you can use the derivative functions. The example we showed above was of what is called a type d evaluator. Let's stay with that.

The evaluator function we wrote was

```
void myeval(x, y)
{
    y = exp(-x^2 + x - 3)
}
```

All type d evaluators open the same way,

```
void evaluator(x, y)
```

although what you name the arguments is up to you. We named the arguments the way that mathematicians think, although we could just as well have named them the way that statisticians think:

```
void evaluator(b, s)
```

To avoid favoritism, we will write them as

```
void evaluator(p, v)
```

That is, we will think in terms of computing the derivative of  $v = f(p)$  with respect to the elements of  $p$ .

Here is the full definition of a type d evaluator:

---

```
void evaluator(real rowvector p, v)
```

where  $v$  is the value to be returned:

$v$ :     *real scalar*

*evaluator()* is to fill in  $v$  given the values in  $p$ .

*evaluator()* may return  $v = .$  if  $f()$  cannot be evaluated at  $p$ .

---

## Example of a type d evaluator

We wish to compute the gradient of the following function at  $p_1 = 1$  and  $p_2 = 2$ :

$$v = \exp(-p_1^2 - p_2^2 - p_1 p_2 + p_1 - p_2 - 3)$$

Our numerical solution to the problem is

```
: void eval_d(p, v)
> {
>     v = exp(-p[1]^2 - p[2]^2 - p[1]*p[2] + p[1] - p[2] - 3)
> }
: D = deriv_init()
: deriv_init_evaluator(D, &eval_d())
: deriv_init_params(D, (1,2))
: grad = deriv(D, 1)
: grad
      1          2
1  [ -.0000501051  -.0001002102 ]
: (-2*1 - 2 + 1)*exp(-1^2 - 2^2 - 1*2 + 1 - 2 - 3)
  -.0000501051
: (-2*2 - 1 - 1)*exp(-1^2 - 2^2 - 1*2 + 1 - 2 - 3)
  -.0001002102
```

For this problem, the elements of the gradient function are given by the following formulas, and we see that *deriv()* computed values that are in agreement with the analytical results (to the number of significant digits being displayed).

$$\frac{dv}{dp_1} = (-2p_1 - p_2 + 1) \exp(-p_1^2 - p_2^2 - p_1 p_2 + p_1 - p_2 - 3)$$

$$\frac{dv}{dp_2} = (-2p_2 - p_1 - 1) \exp(-p_1^2 - p_2^2 - p_1 p_2 + p_1 - p_2 - 3)$$

## Type v evaluators

In some statistical applications, you will find type v evaluators more convenient to code than type d evaluators.

In statistical applications, one tends to think of a dataset of values arranged in matrix  $X$ , the rows of which are observations. The function  $h(p, X[i, :])$  can be calculated for each row separately, and it is the sum of those resulting values that forms the function  $f(p)$  from which we would like to compute derivatives.

Type v evaluators are for such cases.

In a type d evaluator, you return scalar  $v = f(p)$ .

In a type v evaluator, you return a column vector,  $v$ , such that  $\text{colsum}(v) = f(p)$ .

The code outline for type v evaluators is the same as those for d evaluators. All that differs is that  $v$ , which is a *real scalar* in the d case, is now a *real colvector* in the v case.

## User-defined arguments

The type v evaluators arise in statistical applications and, in such applications, there are data; that is, just knowing  $p$  is not sufficient to calculate  $v$ ,  $g$ , and  $H$ . Actually, that same problem can also arise when coding type d evaluators.

You can pass extra arguments to evaluators. The first line of all evaluators, regardless of type, is

```
void evaluator(p, v)
```

If you code

```
deriv_init_argument(D, 1, X)
```

the first line becomes

```
void evaluator(p, X, v)
```

If you code

```
deriv_init_argument(D, 1, X)
deriv_init_argument(D, 2, Y)
```

the first line becomes

```
void evaluator(p, X, Y, v)
```

and so on, up to nine extra arguments. That is, you can specify extra arguments to be passed to your function.

### Example of a type v evaluator

You have the following data:

: x	
	1
1	.35
2	.29
3	.3
4	.3
5	.65
6	.56
7	.37
8	.16
9	.26
10	.19

You believe that the data are the result of a beta distribution process with fixed parameters alpha and beta, and you wish to compute the gradient vector and Hessian matrix associated with the log likelihood at some values of those parameters alpha and beta (*a* and *b* in what follows). The formula for the density of the beta distribution is

$$\text{density}(x) = \frac{\Gamma(a + b)}{\Gamma(a)\Gamma(b)} x^{a-1} (1 - x)^{b-1}$$

In our type v solution to this problem, we compute the gradient and Hessian at *a* = 0.5 and *b* = 2.

```
: void lnbetaden_v(p, x, lnf)
> {
>     a = p[1]
>     b = p[2]
>     lnf = lngamma(a+b) :- lngamma(a) :- lngamma(b) :+
>           (a-1)*log(x) :+ (b-1)*log(1:-x)
> }

: D = deriv_init()
: deriv_init_evaluator(D, &lnbetaden_v())
: deriv_init_evaluatortype(D, "v")
: deriv_init_params(D, (0.5, 2))
: deriv_init_argument(D, 1, x)
: deriv(D, 2)
[symmetric]
```

	1	2
1	-116.4988089	
2	8.724410052	-1.715062542

```
: deriv_result_gradient(D)
```

	1	2
1	15.12578465	-1.701917722



Note the following:

1. Rather than calling the returned value `v`, we called it `lnf`. You can name the arguments as you please.
2. We arranged for an extra argument to be passed by coding `deriv_init_argument(D, 1, x)`. The extra argument is the vector `x`, which we listed previously for you. In our function, we received the argument as `x`, but we could have used a different name just as we used `lnf` rather than `v`.
3. We set the evaluator type to `"v"`.

## Type t evaluators

Type `t` evaluators are for when you need to compute the Jacobian matrix from a vector-valued function.

Type `t` evaluators are different from type `v` evaluators in that the resulting vector of values should not be summed. One example is when the function `f()` performs a nonlinear transformation from the domain of  $p$  to the domain of  $v$ .

## Example of a type t evaluator

Let's compute the Jacobian matrix for the following transformation:

$$v_1 = p_1 + p_2$$

$$v_2 = p_1 - p_2$$

Here is our numerical solution, evaluating the Jacobian at  $p = (0, 0)$ :

```
: void eval_t1(p, v)
> {
>     v = J(1,2,.)
>     v[1] = p[1] + p[2]
>     v[2] = p[1] - p[2]
> }
: D = deriv_init()
: deriv_init_evaluator(D, &eval_t1())
: deriv_init_evaluatortype(D, "t")
: deriv_init_params(D, (0,0))
: deriv(D, 1)
[symmetric]
      1      2
```

1	1
2	1 -1

Now let's compute the Jacobian matrix for a less trivial transformation:

$$v_1 = p_1^2$$

$$v_2 = p_1 p_2$$

Here is our numerical solution, evaluating the Jacobian at  $p = (1, 2)$ :

```

: void eval_t2(p, v)
> {
>     v = J(1,2,.)
>     v[1] = p[1]^2
>     v[2] = p[1] * p[2]
> }
: D = deriv_init()
: deriv_init_evaluator(D, &eval_t2())
: deriv_init_evaluatortype(D, "t")
: deriv_init_params(D, (1,2))
: deriv(D, 1)

```

	1	2
1	1.999999998	0
2	2	1

## Functions

### deriv\_init()

*transmorphic* deriv\_init()

deriv\_init() is used to begin a derivative problem. Store the returned result in a variable name of your choosing; we have used  $D$  in this documentation. You pass  $D$  as the first argument to the other deriv\*() functions.

deriv\_init() sets all deriv\_init\_\*() values to their defaults. You may use the query form of deriv\_init\_\*() to determine an individual default, or you can use deriv\_query() to see them all.

The query form of deriv\_init\_\*() can be used before or after calling deriv().

### deriv\_init\_evaluator() and deriv\_init\_evaluatortype()

*void* deriv\_init\_evaluator( $D$ , *pointer(function) scalar fptr*)

*void* deriv\_init\_evaluatortype( $D$ , *evaluatortype*)

*pointer(function) scalar* deriv\_init\_evaluator( $D$ )

*string scalar* deriv\_init\_evaluatortype( $D$ )

deriv\_init\_evaluator( $D$ ,  $fptr$ ) specifies the function to be called to evaluate  $f(p)$ . Use of this function is required. If your function is named myfcn(), you code deriv\_init\_evaluator( $D$ , &myfcn()).

deriv\_init\_evaluatortype( $D$ , *evaluatortype*) specifies the capabilities of the function that has been set using deriv\_init\_evaluator(). Alternatives for *evaluatortype* are "d", "v", and "t". The default is "d" if you do not invoke this function.

deriv\_init\_evaluator( $D$ ) returns a pointer to the function that has been set.

deriv\_init\_evaluatortype( $D$ ) returns the evaluator type currently set.

**deriv\_init\_argument() and deriv\_init\_narguments()**

*void*                `deriv_init_argument(D, real scalar k, X)`

*void*                `deriv_init_narguments(D, real scalar K)`

*pointer scalar* `deriv_init_argument(D, real scalar k)`

*real scalar*        `deriv_init_narguments(D)`

`deriv_init_argument(D, k, X)` sets the *k*th extra argument of the evaluator function to be *X*. *X* can be anything, including a view matrix or even a pointer to a function. No copy of *X* is made; it is a pointer to *X* that is stored, so any changes you make to *X* between setting it and *X* being used will be reflected in what is passed to the evaluator function.

`deriv_init_narguments(D, K)` sets the number of extra arguments to be passed to the evaluator function. This function is useless and included only for completeness. The number of extra arguments is automatically set when you use `deriv_init_argument()`.

`deriv_init_argument(D, k)` returns a pointer to the object that was previously set.

`deriv_init_narguments(D)` returns the number of extra arguments that were passed to the evaluator function.

**deriv\_init\_weights()**

*void*                `deriv_init_weights(D, real colvector weights)`

*pointer scalar* `deriv_init_weights(D)`

`deriv_init_weights(D, weights)` sets the weights used with type *v* evaluators to produce the function value. By default, `deriv()` with a type *v* evaluator uses `colsum(v)` to compute the function value. With weights, `deriv()` uses `cross(weights, v)`. *weights* must be row conformable with the column vector returned by the evaluator.

`deriv_init_weights(D)` returns a pointer to the weight vector that was previously set.

**deriv\_init\_params()**

*void*                `deriv_init_params(D, real rowvector params)`

*real rowvector* `deriv_init_params(D)`

`deriv_init_params(D, params)` sets the parameter values at which the derivatives will be computed. Use of this function is required.

`deriv_init_params(D)` returns the parameter values at which the derivatives were computed.

**Advanced init functions**

The rest of the `deriv_init_*`() functions provide finer control of the numerical derivative taker.

**deriv\_init\_h(), ...\_scale(), ...\_bounds(), ...\_search()**

```
void          deriv_init_h(D, real rowvector h)
void          deriv_init_scale(D, real rowvector s)
void          deriv_init_bounds(D, real rowvector minmax)
void          deriv_init_search(D, search)
real rowvector deriv_init_h(D)
real rowvector deriv_init_scale(D)
real rowvector deriv_init_bounds(D)
string scalar deriv_init_search(D)
```

`deriv_init_h(D, h)` sets the *h* values used to compute numerical derivatives.

`deriv_init_scale(D, s)` sets the starting scale values used to compute numerical derivatives.

`deriv_init_bounds(D, minmax)` sets the minimum and maximum values used to search for optimal scale values. The default is *minmax* = (1e-8, 1e-7).

`deriv_init_search(D, "interpolate")` causes `deriv()` to use linear and quadratic interpolation to search for an optimal delta for computing the numerical derivatives. This is the default search method.

`deriv_init_search(D, "bracket")` causes `deriv()` to use a bracketed quadratic formula to search for an optimal delta for computing the numerical derivatives.

`deriv_init_search(D, "off")` prevents `deriv()` from searching for an optimal delta.

`deriv_init_h(D)` returns the user-specified *h* values.

`deriv_init_scale(D)` returns the user-specified starting scale values.

`deriv_init_bounds(D)` returns the user-specified search bounds.

`deriv_init_search(D)` returns the currently set search method.

**deriv\_init\_verbose()**

```
void          deriv_init_verbose(D, verbose)
string scalar deriv_init_verbose(D)
```

`deriv_init_verbose(D, verbose)` sets whether error messages that arise during the execution of `deriv()` or `_deriv()` are to be displayed. Setting *verbose* to "on" means that they are displayed; "off" means that they are not displayed. The default is "on". Setting *verbose* to "off" is of interest only to users of `_deriv()`.

`deriv_init_verbose(D)` returns the current value of *verbose*.

**deriv()**

(*varies*) `deriv(D, todo)`

`deriv(D, todo)` invokes the derivative process. If something goes wrong, `deriv()` aborts with error.

`deriv(D, 0)` returns the function value without computing derivatives.

`deriv(D, 1)` returns the gradient vector; the Hessian matrix is not computed.

`deriv(D, 2)` returns the Hessian matrix; the gradient vector is also computed.

Before you can invoke `deriv()`, you must have defined your evaluator function, `evaluator()`, and you must have set the parameter values at which `deriv()` is to compute derivatives:

```
D = deriv_init()
deriv_init_evaluator(D, &evaluator())
deriv_init_params(D, (...))
```

The above assumes that your evaluator function is type `d`. If your evaluator function type is `v` (that is, it returns a column vector of values instead of a scalar value), you will also have coded

```
deriv_init_evaluortype(D, "v")
```

and you may have coded other `deriv_init_*`() functions as well.

Once `deriv()` completes, you may use the `deriv_result_*`() functions. You may also continue to use the `deriv_init_*`() functions to access initial settings, and you may use them to change settings and recompute derivatives (that is, invoke `deriv()` again) if you wish.

**deriv()**

*real scalar* `_deriv(D, todo)`

`_deriv(D)` performs the same actions as `deriv(D)` except that, rather than returning the requested derivatives, `_deriv()` returns a real scalar and, rather than aborting if numerical issues arise, `_deriv()` returns a nonzero value. `_deriv()` returns 0 if all went well. The returned value is called an error code.

`deriv()` returns the requested result. It can work that way because the numerical derivative calculation must have gone well. Had it not, `deriv()` would have aborted execution.

`_deriv()` returns an error code. If it is 0, the numerical derivative calculation went well, and you can obtain the gradient vector by using `deriv_result_gradient()`. If things did not go well, you can use the error code to diagnose what went wrong and take the appropriate action.

Thus `_deriv(D)` is an alternative to `deriv(D)`. Both functions do the same thing. The difference is what happens when there are numerical difficulties.

`deriv()` and `_deriv()` work around most numerical difficulties. For instance, the evaluator function you write is allowed to return `v` equal to missing if it cannot calculate the  $f()$  at  $p + d$ . If that happens while computing the derivative, `deriv()` and `_deriv()` will search for a better  $d$  for taking the derivative. `deriv()`, however, cannot tolerate that happening at  $p$  (the parameter values you set using `deriv_init_params()`) because the function value must exist at the point when you want

`deriv()` to compute the numerical derivative. `deriv()` issues an error message and aborts, meaning that execution is stopped. There can be advantages in that. The calling program need not include complicated code for such instances, figuring that stopping is good enough because a human will know to address the problem.

`_deriv()`, however, does not stop execution. Rather than aborting, `_deriv()` returns a nonzero value to the caller, identifying what went wrong. The only exception is that `_deriv()` will return a zero value to the caller even when the evaluator function returns `v` equal to missing at `p`, allowing programmers to handle this special case without having to turn `deriv_init_verbose()` off.

Programmers implementing advanced systems will want to use `_deriv()` instead of `deriv()`. Everybody else should use `deriv()`.

Programmers using `_deriv()` will also be interested in the functions `deriv_init_verbose()`, `deriv_result_errorcode()`, `deriv_result_errortext()`, and `deriv_result_returncode()`.

The error codes returned by `_deriv()` are listed below, under the heading `deriv_result_errorcode()`, `..._errortext()`, and `..._returncode()`.

### **`deriv_result_value()`**

*real scalar* `deriv_result_value(D)`

`deriv_result_value(D)` returns the value of `f()` evaluated at `p`.

### **`deriv_result_values()` and `_deriv_result_values()`**

*real matrix* `deriv_result_values(D)`

*void* `_deriv_result_values(D, v)`

`deriv_result_values(D)` returns the vector values returned by the evaluator. For type `v` evaluators, this is the column vector that sums to the value of `f()` evaluated at `p`. For type `t` evaluators, this is the rowvector returned by the evaluator.

`_deriv_result_values(D, v)` uses `swap()` (see [M-5] `swap()`) to interchange `v` with the vector values stored in `D`. This destroys the vector values stored in `D`.

These functions should be called only with type `v` evaluators.

### **`deriv_result_gradient()` and `_deriv_result_gradient()`**

*real rowvector* `deriv_result_gradient(D)`

*void* `_deriv_result_gradient(D, g)`

`deriv_result_gradient(D)` returns the gradient vector evaluated at `p`.

`_deriv_result_gradient(D, g)` uses `swap()` (see [M-5] `swap()`) to interchange `g` with the gradient vector stored in `D`. This destroys the gradient vector stored in `D`.

**deriv\_result\_scores() and \_deriv\_result\_scores()**

```

real matrix deriv_result_scores(D)
void         _deriv_result_scores(D, S)

```

`deriv_result_scores(D)` returns the matrix of the scores evaluated at *p*. The matrix of scores can be summed over the columns to produce the gradient vector.

`_deriv_result_scores(D, S)` uses `swap()` (see [M-5] [swap\(\)](#)) to interchange *S* with the scores matrix stored in *D*. This destroys the scores matrix stored in *D*.

These functions should be called only with type *v* evaluators.

**deriv\_result\_Jacobian() and \_deriv\_result\_Jacobian()**

```

real matrix deriv_result_Jacobian(D)
void         _deriv_result_Jacobian(D, J)

```

`deriv_result_Jacobian(D)` returns the Jacobian matrix evaluated at *p*.

`_deriv_result_Jacobian(D, J)` uses `swap()` (see [M-5] [swap\(\)](#)) to interchange *J* with the Jacobian matrix stored in *D*. This destroys the Jacobian matrix stored in *D*.

These functions should be called only with type *t* evaluators.

**deriv\_result\_Hessian() and \_deriv\_result\_Hessian()**

```

real matrix deriv_result_Hessian(D)
void         _deriv_result_Hessian(D, H)

```

`deriv_result_Hessian(D)` returns the Hessian matrix evaluated at *p*.

`_deriv_result_Hessian(D, H)` uses `swap()` (see [M-5] [swap\(\)](#)) to interchange *H* with the Hessian matrix stored in *D*. This destroys the Hessian matrix stored in *D*.

These functions should not be called with type *t* evaluators.

**deriv\_result\_h(), ...\_scale(), and ...\_delta()**

```

real rowvector deriv_result_h(D)
real rowvector deriv_result_scale(D)
real rowvector deriv_result_delta(D)

```

`deriv_result_h(D)` returns the vector of *h* values that was used to compute the numerical derivatives.

`deriv_result_scale(D)` returns the vector of scale values that was used to compute the numerical derivatives.

`deriv_result_delta(D)` returns the vector of delta values used to compute the numerical derivatives.

**deriv\_result\_errorcode(), ...\_errortext(), and ...\_returncode()**

```
real scalar    deriv_result_errorcode(D)

string scalar  deriv_result_errortext(D)

real scalar    deriv_result_returncode(D)
```

These functions are for use after `_deriv()`.

`deriv_result_errorcode(D)` returns the same error code as `_deriv()`. The value will be zero if there were no errors. The error codes are listed in the table directly below.

`deriv_result_errortext(D)` returns a string containing the error message corresponding to the error code. If the error code is zero, the string will be "".

`deriv_result_returncode(D)` returns the Stata return code corresponding to the error code. The mapping is listed in the table directly below.

In advanced code, these functions might be used as

```
(void) _deriv(D, todo)
... if (ec = deriv_result_code(D)) {
    errprintf("{p}\n")
    errprintf("%s\n", deriv_result_errortext(D))
    errprintf("{p_end}\n")
    exit(deriv_result_returncode(D))
    /*NOTREACHED*/
}
```

The error codes and their corresponding Stata return codes are

Error code	Return code	Error text
1	198	invalid todo argument
2	111	evaluator function required
3	459	parameter values required
4	459	parameter values not feasible
5	459	could not calculate numerical derivatives—discontinuous region with missing values encountered
6	459	could not calculate numerical derivatives—flat or discontinuous region encountered
16	111	<i>function()</i> not found
17	459	Hessian calculations not allowed with type <code>tt</code> evaluators

NOTE: Error 4 can occur only when evaluating *f()* at the parameter values.  
This error occurs only with `deriv()`.

**deriv\_query()**

```
void deriv_query(D)
```

`deriv_query(D)` displays a report on the current `deriv_init_*`() values and some of the `deriv_result_*`() values. `deriv_query(D)` may be used before or after `deriv()`, and it is



useful when using `deriv()` interactively or when debugging a program that calls `deriv()` or `_deriv()`.

## Conformability

All functions have  $1 \times 1$  inputs and have  $1 \times 1$  or *void* outputs, except the following:

`deriv_init_params(D, params):`

*D*: *transmorphic*  
*params*:  $1 \times np$   
*result*: *void*

`deriv_init_params(D):`

*D*: *transmorphic*  
*result*:  $1 \times np$

`deriv_init_argument(D, k, X):`

*D*: *transmorphic*  
*k*:  $1 \times 1$   
*X*: *anything*  
*result*: *void*

`deriv_init_weights(D, params):`

*D*: *transmorphic*  
*params*:  $N \times 1$   
*result*: *void*

`deriv_init_h(D, h):`

*D*: *transmorphic*  
*h*:  $1 \times np$   
*result*: *void*

`deriv_init_h(D):`

*D*: *transmorphic*  
*result*:  $1 \times np$

`deriv_init_scale(D, scale):`

*D*: *transmorphic*  
*scale*:  $1 \times np$  (type *d* and *v* evaluator)  
 $nv \times np$  (type *t* evaluator)  
*void*: *void*

`deriv_init_bounds(D, minmax):`

*D*: *transmorphic*  
*minmax*:  $1 \times 2$   
*result*: *void*

`deriv_init_bounds(D):`

*D*: *transmorphic*  
*result*:  $1 \times w$

```
deriv(D, 0):
    D:    transmorphic
    result: 1 × 1
           1 × nv (type t evaluator)

deriv(D, 1):
    D:    transmorphic
    result: 1 × np
           nv × np (type t evaluator)

deriv(D, 2):
    D:    transmorphic
    result: np × np

deriv_result_values(D):
    D:    transmorphic
    result: N × 1
           1 × nv (type t evaluator)
           N × 1 (type v evaluator)

_deriv_result_values(D, v):
    D:    transmorphic
    v:    N × 1
           1 × nv (type t evaluator)
           N × 1 (type v evaluator)
    result: void

deriv_result_gradient(D):
    D:    transmorphic
    result: 1 × np

_deriv_result_gradient(D, g):
    D:    transmorphic
    g:    1 × np
    result: void

deriv_result_scores(D):
    D:    transmorphic
    result: N × np

_deriv_result_scores(D, S):
    D:    transmorphic
    S:    N × np
    result: void

deriv_result_Jacobian(D):
    D:    transmorphic
    result: nv × np

_deriv_result_Jacobian(D, J):
    D:    transmorphic
    J:    nv × np
    result: void
```

```

deriv_result_Hessian(D):
    D:      transmorphic
    result:  np × np

_deriv_result_Hessian(D, H):
    D:      transmorphic
    H:      np × np
    result:  void

deriv_result_h(D):
    D:      transmorphic
    result:  1 × np

deriv_result_scale(D):
    D:      transmorphic
    result:  1 × np (type d and v evaluator)
            nv × np (type t evaluator)

deriv_result_delta(D):
    D:      transmorphic
    result:  1 × np (type d and v evaluator)
            nv × np (type t evaluator)

```

## Diagnostics

All functions abort with error when used incorrectly.

`deriv()` aborts with error if it runs into numerical difficulties. `_deriv()` does not; it instead returns a nonzero error code.

## Methods and formulas

See sections 1.3.4 and 1.3.5 of [Gould, Pitblado, and Poi \(2010\)](#) for an overview of the methods and formulas `deriv()` uses to compute numerical derivatives.

Carl Gustav Jacob Jacobi (1804–1851) was born in Potsdam, Prussia (now Germany). A prodigy, at secondary school he read advanced mathematics texts such as Euler’s *Introductio in analysin infinitorum*. Even at the University of Berlin, Jacobi studied on his own using the works of Lagrange and other leading mathematicians. His career included university posts in Berlin and Königsberg. Jacobi made major discoveries in analysis (especially elliptic functions and differential equations), algebra, geometry, mechanics, and astronomy. Expanding the theory of the determinant, which was previously defined and presented by Cauchy, Jacobi invented the functional determinant of the Jacobian matrix. Thus the Jacobian was born. It was described in his memoir, “De determinantibus functionalibus”, which was published in 1841. Jacobi was also an inspiring teacher who made pioneering use of seminars. In the last years of his life, Jacobi suffered from much ill-health: he died in Berlin after catching influenza followed by smallpox.

## References

Gould, W. W., J. S. Pitblado, and B. P. Poi. 2010. *Maximum Likelihood Estimation with Stata*. 4th ed. College Station, TX: Stata Press.

James, I. 2002. *Remarkable Mathematicians: From Euler to von Neumann*. Cambridge: Cambridge University Press.

## Also see

[M-4] [mathematical](#) — Important mathematical functions

# Title

[M-5] **designmatrix()** — Design matrices

## Syntax

*real matrix*    **designmatrix**(*real colvector v*)

## Description

**designmatrix**(*v*) returns a  $\text{rows}(v) \times \text{colmax}(v)$  matrix with ones in the indicated columns and zero everywhere else.

## Remarks

**designmatrix**((1\2\3)) is equal to  $I(3)$ , the  $3 \times 3$  identity matrix.

## Conformability

**designmatrix**(*v*):

<i>v</i> :	$r \times 1$
<i>result</i> :	$r \times \text{colmax}(v)$ ( $0 \times 0$ if $r = 0$ )

## Diagnostics

**designmatrix**(*v*) aborts with error if any element of *v* is  $<1$ .

## Also see

[\[M-4\] standard](#) — Functions to create standard matrices

# Title

[M-5] `det()` — Determinant of matrix

## Syntax

*numeric scalar*   `det`(*numeric matrix A*)

*numeric scalar*   `dettriangular`(*numeric matrix A*)

## Description

`det(A)` returns the determinant of  $A$ .

`dettriangular(A)` returns the determinant of  $A$ , treating  $A$  as if it were triangular (even if it is not).

## Remarks

Calculation of the determinant is made by obtaining the LU decomposition of  $A$  and then calculating the determinant of  $U$ :

$$\begin{aligned}\det(A) &= \det(PLU) \\ &= \det(P) \times \det(L) \times \det(U) \\ &= \pm 1 \times 1 \times \det(U) \\ &= \pm \det(U)\end{aligned}$$

Since  $U$  is (upper) triangular,  $\det(U)$  is simply the product of its diagonal elements. See [M-5] `lud()`.

## Conformability

`det(A)`, `dettriangular(A)`:  
          *A*:         $n \times n$   
          *result*:     $1 \times 1$

## Diagnostics

`det(A)` and `dettriangular(A)` return 1 if  $A$  is  $0 \times 0$ .

`det(A)` aborts with error if  $A$  is not square and returns missing if  $A$  contains missing values.

`dettriangular(A)` aborts with error if  $A$  is not square and returns missing if any element on the diagonal of  $A$  is missing.

Both `det(A)` and `dettriangular(A)` will return missing value if the determinant exceeds 8.99e+307.

---

## Also see

[M-5] `lud()` — LU decomposition

[M-4] `matrix` — Matrix functions

# Title

[M-5] **\_diag()** — Replace diagonal of a matrix

## Syntax

*void \_diag(numeric matrix Z, numeric vector v)*

## Description

`_diag(Z, v)` replaces the diagonal of the matrix  $Z$  with  $v$ .  $Z$  need not be square.

1. If  $v$  is a vector, the vector replaces the principal diagonal.
2. If  $v$  is  $1 \times 1$ , each element of the principal diagonal is replaced with  $v$ .
3. If  $v$  is a void vector ( $1 \times 0$  or  $0 \times 1$ ),  $Z$  is left unchanged.

## Conformability

`_diag(Z, v)`:

*input:*

$Z$ :  $n \times m, n \leq m$

$v$ :  $1 \times 1, 1 \times n$ , or  $n \times 1$

or

$Z$ :  $n \times m, n > m$

$v$ :  $1 \times 1, 1 \times m$ , or  $m \times 1$

*output:*

$Z$ :  $n \times m$

## Diagnostics

`_diag(Z, v)` aborts with error if  $Z$  or  $v$  is a view.

## Also see

[M-5] **diag()** — Create diagonal matrix

[M-4] **manipulation** — Matrix manipulation



Syntax

*numeric matrix*    `diag(numeric matrix Z)`

*numeric matrix*    `diag(numeric vector z)`

Description

`diag()` creates diagonal matrices.

`diag(Z)`, *Z* a matrix, extracts the principal diagonal of *Z* to create a new matrix. *Z* must be square.

`diag(z)`, *z* a vector, creates a new matrix with the elements of *z* on its diagonal.

Remarks

Do not confuse `diag()` with its functional inverse, `diagonal()`; see [M-5] [diagonal\(\)](#). `diag()` creates a matrix from a vector (or matrix); `diagonal()` extracts the diagonal of a matrix into a vector.

Use of `diag()` should be avoided because it wastes memory. The [colon operators](#) will allow you to use vectors directly:

Desired calculation	Equivalent
<code>diag(v)*X</code> ,	
<i>v</i> is a column	<code>v:*X</code>
<i>v</i> is a row	<code>v':*X</code>
<i>v</i> is a matrix	<code>diagonal(v):*X</code>
<code>X*diag(v)</code>	
<i>v</i> is a column	<code>X:*v'</code>
<i>v</i> is a row	<code>X:*v</code>
<i>v</i> is a matrix	<code>X:*diagonal(v)'</code>

In the above table, it is assumed that *v* is real. If *v* might be complex, the transpose operators that appear must be changed to `transposeonly()` calls, because we do not want the conjugate. For instance, `v':*X` would become `transposeonly(v):*X`.

## Conformability

`diag(Z)`:

<i>Z</i> :	$m \times n$
<i>result</i> :	$\min(m, n) \times \min(m, n)$

`diag(z)`:

<i>z</i> :	$1 \times n$ or $n \times 1$
<i>result</i> :	$n \times n$

## Diagnostics

None.

## Also see

- [M-5] [\\_diag\(\)](#) — Replace diagonal of a matrix
- [M-5] [diagonal\(\)](#) — Extract diagonal into column vector
- [M-5] [isdiagonal\(\)](#) — Whether matrix is diagonal
- [M-4] [manipulation](#) — Matrix manipulation

# Title

[M-5] `diag0cnt()` — Count zeros on diagonal

## Syntax

*real scalar*   `diag0cnt(real matrix X)`

## Description

`diag0cnt(X)` returns the number of principal diagonal elements of  $X$  that are 0.

## Remarks

`diag0cnt()` is often used after `invsym()` (see [M-5] [invsym\(\)](#)) to count the number of columns dropped because of collinearity.

## Conformability

`diag0cnt(X)`:  
           $X$ :      $r \times c$   
           $result$ :    $1 \times 1$

## Diagnostics

`diag0cnt(X)` returns 0 if  $X$  is void.

## Also see

[M-5] [invsym\(\)](#) — Symmetric real matrix inversion

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] **diagonal()** — Extract diagonal into column vector

## Syntax

*numeric colvector*   **diagonal**(*numeric matrix A*)

## Description

**diagonal**(*A*) extracts the diagonal of *A* and returns it in a column vector.

## Remarks

**diagonal**() may be used with nonsquare matrices.

Do not confuse **diagonal**() with its functional inverse, **diag**(); see [M-5] **diag()**. **diagonal**() extracts the diagonal of a matrix into a vector; **diag**() creates a diagonal matrix from a vector.

## Conformability

**diagonal**(*A*):  
    *A*:       $r \times c$   
    *result*:    $\min(r, c) \times 1$

## Diagnostics

None.

## Also see

[M-5] **diag**() — Create diagonal matrix

[M-5] **isdiagonal**() — Whether matrix is diagonal

[M-5] **blockdiag**() — Block-diagonal matrix

[M-4] **manipulation** — Matrix manipulation

## Syntax

*string colvector*    `dir(dirname, filetype, pattern)`

*string colvector*    `dir(dirname, filetype, pattern, prefix)`

where

*dirname:*    *string scalar* containing directory name

*filetype:*    *string scalar* containing "files", "dirs", or "other"

*pattern:*    *string scalar* containing match pattern

*prefix:*    *real scalar*    containing 0 or 1

## Description

`dir(dirname, filetype, pattern)` returns a column vector containing the names of the files in *dir* that match *pattern*.

`dir(dirname, filetype, pattern, prefix)` does the same thing but allows you to specify whether you want a simple list of files (*prefix* = 0) or a list of filenames prefixed with *dirname* (*prefix* ≠ 0). `dir(dirname, filetype, pattern)` is equivalent to `dir(dirname, filetype, pattern, 0)`.

*pattern* is interpreted by [M-5] `strmatch()`.

## Remarks

Examples:

`dir(".", "dirs", "*")`

returns a list of all directories in the current directory.

`dir(".", "files", "*")`

returns a list of all regular files in the current directory.

`dir(".", "files", "*.sthlp")`

returns a list of all \*.sthlp files found in the current directory.

## Conformability

`dir(dirname, filetype, pattern, prefix):`

*dirname:*     $1 \times 1$

*filetype:*     $1 \times 1$

*pattern:*     $1 \times 1$

*prefix:*     $1 \times 1$  (optional)

*result:*     $k \times 1$ ,     $k$  number of files matching pattern

## Diagnostics

`dir(dirname, filetype, pattern, prefix)` returns `J(0,1,"")` if

1. no files matching *pattern* are found,
2. directory *dirname* does not exist, or
3. *filetype* is misspecified (is not equal to `"files"`, `"dirs"`, or `"others"`).

*dirname* may be specified with or without the directory separator on the end.

*dirname* = `"` is interpreted the same as *dirname* = `"."`; the current directory is searched.

## Also see

[\[M-4\] io](#) — I/O functions

# Title

[M-5] **direxists()** — Whether directory exists

## Syntax

*real scalar*    **direxists**(*string scalar dirname*)

## Description

**direxists**(*dirname*) returns 1 if *dirname* contains a valid path to a directory and returns 0 otherwise.

## Conformability

**direxists**(*dirname*):

<i>dirname</i> :	1 × 1
<i>result</i> :	1 × 1

## Diagnostics

None.

## Also see

[\[M-4\] io](#) — I/O functions

# Title

[M-5] **direxternal()** — Obtain list of existing external globals

## Syntax

*string colvector*    **direxternal**(*string scalar pattern*)

## Description

**direxternal**(*pattern*) returns a column vector containing the names matching *pattern* of the existing external globals. **direxternal**() returns J(0,1,"") if there are no such globals.

*pattern* is interpreted by [M-5] **strmatch**().

## Remarks

See [M-5] **findexternal**() for the definition of a global.

A list of all globals can be obtained by **direxternal**("\*").

## Conformability

**direxternal**(*pattern*):

<i>pattern</i> :	$1 \times 1$
<i>result</i> :	$n \times 1$

## Diagnostics

**direxternal**(*pattern*) returns J(0,1,"") when there are no globals matching *pattern*.

## Also see

[M-5] **findexternal**() — Find, create, and remove external globals

[M-4] **programming** — Programming functions



# Title

[M-5] **display()** — Display text interpreting SMCL

## Syntax

```
void display(string colvector s)
```

```
void display(string colvector s, real scalar asis)
```

## Description

`display(s)` displays the string or strings contained in *s*.

`display(s, asis)` does the same thing but allows you to control how SMCL codes are treated. `display(s, 0)` is equivalent to `display(s)`; any SMCL codes are honored.

`display(s, asis)`, *asis*  $\neq 0$ , displays the contents of *s* exactly as they are. For instance, when *asis*  $\neq 0$ , "{it}" is just the string of characters {, i, t, and } and those characters are displayed; {it} is not given the SMCL interpretation of enter italic mode.

## Remarks

When *s* is a scalar, the differences between coding

```
: display(s)
```

and coding

```
: s
```

are

1. `display(s)` will not indent *s*; *s* by itself causes *s* to be indented by two spaces.
2. `display(s)` will honor any SMCL codes contained in *s*; *s* by itself is equivalent to `display(s, 1)`. For example,

```
: s = "this is an {it:example}"
: display(s)
this is an example
: s
  this is an {it:example}
```

3. When *s* is a vector, `display(s)` simply displays the lines, whereas *s* by itself adorns the lines with row and column numbers:

```

: s = ("this is line 1" \ "this is line 2")
: display(s)
this is line 1
this is line 2
: s

1   this is line 1
2   this is line 2

```

Another alternative to `display()` is `printf()`; see [M-5] `printf()`. When *s* is a scalar, `display()` and `printf()` do the same thing:

```

: display("this is an {it:example}")
this is an example

: printf("%s\n", "this is an {it:example}")
this is an example

```

`printf()`, however, will not allow *s* to be nonscalar; it has other capabilities.

## Conformability

```

display(s, asis)
      s:       $k \times 1$ 
      asis:    $1 \times 1$    (optional)
      result: void

```

## Diagnostics

None.

## Also see

- [M-5] `displayas()` — Set display level
- [M-5] `displayflush()` — Flush terminal-output buffer
- [M-5] `printf()` — Format output
- [M-4] `io` — I/O functions

## Syntax

```
void displayas(string scalar level)
```

where *level* may be

<i>level</i>	Minimum abbreviation
"result"	"res"
"text"	"txt"
"error"	"err"
"input"	"inp"

## Description

`displayas(level)` sets whether and how subsequent output is to be displayed.

## Remarks

If this function is never invoked, then the output level is `result`. Say that Mata was invoked in such a way that all output except error messages is being suppressed (for example, `quietly` was coded in front of the `mata` command or in front of the `ado-file` that called your Mata function). If output is being suppressed, then Mata output is being suppressed, including any output created by your program. Say that you reach a point in your program where you wish to output an error message. You coded

```
printf("{err:you made a mistake}\n")
```

Even though you coded the `SMCL` directive `{err:}`, the error message will still be suppressed. `SMCL` determines how something is rendered, not whether it is rendered. What you need to code is

```
displayas("err")
printf("{err:you made a mistake}\n")
```

Actually, you could code

```
displayas("err")
printf("you made a mistake\n")
```

because, in addition to setting the output level (telling Stata that all subsequent output is of the specified level), it also sets the current `SMCL` rendering to what is appropriate for that kind of output. Hence, if you coded

```
displayas("err")
printf("{res:you made a mistake}\n")
```

the text you made a mistake would appear in the style of results despite any quietly attempting to suppress output. Coding the above is considered bad style.

## Conformability

```
displayas(level):
    level:      1 × 1
    result:     void
```

## Diagnostics

`displayas(level)` aborts with error if *level* contains an inappropriate string.

## Also see

- [M-5] [printf\(\)](#) — Format output
- [M-5] [display\(\)](#) — Display text interpreting SMCL
- [M-4] [io](#) — I/O functions

# Title

[M-5] **displayflush()** — Flush terminal-output buffer

## Syntax

```
void displayflush()
```

## Description

To achieve better performance, Stata buffers terminal output, so, within a program, output may not appear when a [display\(\)](#) or [printf\(\)](#) command is executed. The output might be held in a buffer and displayed later.

`displayflush()` forces Stata to display all pending output at the terminal. `displayflush()` is rarely used.

## Remarks

See [\[M-5\] printf\(\)](#) for an [example](#) of the use of `displayflush()`.

Use of `displayflush()` slows execution. Use `displayflush()` only when it is important that output be displayed at the terminal now, such as when providing messages indicating what your program is doing.

## Diagnostics

None.

## Also see

[\[M-5\] printf\(\)](#) — Format output

[\[M-5\] display\(\)](#) — Display text interpreting SMCL

[\[M-4\] io](#) — I/O functions

# Title

[M-5] **Dmatrix()** — Duplication matrix

## Syntax

*real matrix* **Dmatrix**(*real scalar n*)

## Description

**Dmatrix**( $n$ ) returns the  $n^2 \times n(n+1)/2$  duplication matrix  $D$  for which  $D \cdot \text{vech}(X) = \text{vec}(X)$ , where  $X$  is an arbitrary  $n \times n$  symmetric matrix.

## Remarks

Duplication matrices are frequently used in computing derivatives of functions of symmetric matrices. Section 9.5 of [Lütkepohl \(1996\)](#) lists many useful properties of duplication matrices.

## Conformability

**Dmatrix**( $n$ ):  
     $n$ :  $1 \times 1$   
     $result$ :  $n^2 \times n(n+1)/2$

## Diagnostics

**Dmatrix**( $n$ ) aborts with error if  $n$  is less than 0 or is missing.  $n$  is interpreted as [trunc\( \$n\$ \)](#).

## Reference

Lütkepohl, H. 1996. *Handbook of Matrices*. New York: Wiley.

## Also see

[M-5] [Kmatrix\(\)](#) — Commutation matrix

[M-5] [Lmatrix\(\)](#) — Elimination matrix

[M-5] [vec\(\)](#) — Stack matrix columns

[M-4] [standard](#) — Functions to create standard matrices

# Title

[M-5] **dsign()** — FORTRAN-like DSIGN() function

## Syntax

*real scalar*   **dsign**(*real scalar a*, *real scalar b*)

## Description

**dsign**(*a*, *b*) returns *a* with the sign of *b*, defined as  $|a|$  if  $b \geq 0$  and  $-|a|$  otherwise.

This function is useful when translating FORTRAN programs.

The in-line construction

$(b \geq 0 ? \text{abs}(a) : -\text{abs}(a))$

is clearer. Also, differentiate carefully between what **dsign**() returns (equivalent to the above construction) and **signum**(*b*)\***abs**(*a*), which is almost equivalent but returns 0 when *b* is 0 rather than **abs**(*a*). (Message: **dsign**() is not one of our favorite functions.)

## Conformability

**dsign**(*a*, *b*):

<i>a</i> :	$1 \times 1$
<i>b</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

**dsign**(., *b*) returns . for all *b*.

**dsign**(*a*, .) returns **abs**(*a*) for all *a*.

## Also see

[M-5] **sign()** — Sign and complex quadrant functions

[M-4] **scalar** — Scalar mathematical functions

# Title

[M-5] **e()** — Unit vectors

## Syntax

*real rowvector*    **e**(*real scalar i*, *real scalar n*)

## Description

**e**(*i*, *n*) returns a  $1 \times n$  unit vector, a vector with all elements equal to zero except for the *i*th, which is set to one.

## Conformability

**e**(*i*, *n*):

<i>i</i> :	$1 \times 1$
<i>n</i> :	$1 \times 1$
<i>result</i> :	$1 \times n$

## Diagnostics

**e**(*i*, *n*) aborts with error if  $n < 1$  or if  $i < 1$  or if  $i > n$ . Arguments *i* and *n* are interpreted as **trunc**(*i*) and **trunc**(*n*).

## Also see

[\[M-4\] standard](#) — Functions to create standard matrices



# Title

[M-5] `editmissing()` — Edit matrix for missing values

## Syntax

*numeric matrix*    `editmissing(numeric matrix A, numeric scalar v)`

*void*                `_editmissing(numeric matrix a, numeric scalar v)`

## Description

`editmissing(A, v)` returns *A* with any missing values changed to *v*.  
`_editmissing(A, v)` replaces all missing values in *A* with *v*.

## Remarks

`editmissing()` and `_editmissing()` are very fast.  
If you want to change nonmissing values to other values, including missing, see [M-5] `editvalue()`.

## Conformability

`editmissing(A, v):`

*A:*         $r \times c$   
*v:*         $1 \times 1$   
*result:*     $r \times c$

`_editmissing(A, v):`

*input:*

*A:*         $r \times c$   
*v:*         $1 \times 1$

  
*output:*

*A:*         $r \times c$

## Diagnostics

`editmissing(A, v)` and `_editmissing(A, v)` change all missing elements to *v*, including not only `.` but also `.a`, `.b`, ..., `.z`.

## Also see

- [M-5] `editvalue()` — Edit (change) values in matrix
- [M-4] `manipulation` — Matrix manipulation

# Title

**[M-5] edittoint()** — Edit matrix for roundoff error (integers)

## Syntax

```
numeric matrix  edittoint(numeric matrix Z, real scalar amt)
void            _edittoint(numeric matrix Z, real scalar amt)

numeric matrix  edittointtol(numeric matrix Z, real scalar tol)
void            _edittointtol(numeric matrix Z, real scalar tol)
```

## Description

These edit functions set elements of matrices to integers that are close to integers.

`edittoint(Z, amt)` and `_edittoint(Z, amt)` set

$$Z_{ij} = \text{round}(Z_{ij}) \quad \text{if } |Z_{ij} - \text{round}(Z_{ij})| \leq |tol|$$

for  $Z$  real and set

$$\text{Re}(Z_{ij}) = \text{round}(\text{Re}(Z_{ij})) \quad \text{if } |\text{Re}(Z_{ij}) - \text{round}(\text{Re}(Z_{ij}))| \leq |tol|$$

$$\text{Im}(Z_{ij}) = \text{round}(\text{Im}(Z_{ij})) \quad \text{if } |\text{Im}(Z_{ij}) - \text{round}(\text{Im}(Z_{ij}))| \leq |tol|$$

for  $Z$  complex, where in both cases

$$tol = \text{abs}(amt) * \text{epsilon}(\text{sum}(\text{abs}(Z)) / (\text{rows}(Z) * \text{cols}(Z)))$$

`edittoint()` leaves  $Z$  unchanged and returns the edited matrix. `_edittoint()` edits  $Z$  in place.

`edittointtol(Z, tol)` and `_edittointtol(Z, tol)` do the same thing, except that  $tol$  is specified directly.

## Remarks

These functions mirror the `edittozero()` functions documented in [\[M-5\] edittozero\(\)](#), except that, rather than solely resetting to zero values close to zero, they reset to integer values close to integers.

See [\[M-5\] edittozero\(\)](#). Whereas use of the functions documented there is recommended, use of the functions documented here generally is not. Although zeros commonly arise in real problems so that there is reason to suspect small numbers would be zero but for roundoff error, integers arise more rarely.

If you have reason to believe that integer values are likely, then by all means use these functions.

## Conformability

`edittoint(Z, amt):`

*Z:*         $r \times c$   
*amt:*      $1 \times 1$   
*result:*    $r \times c$

`_edittoint(Z, amt):`

*input:*

*Z:*         $r \times c$   
*amt:*      $1 \times 1$

*output:*

*Z:*         $r \times c$

`edittointtol(Z, tol):`

*Z:*         $r \times c$   
*tol:*       $1 \times 1$   
*result:*    $r \times c$

`_edittointtol(Z, tol):`

*input:*

*Z:*         $r \times c$   
*tol:*       $1 \times 1$

*output:*

*Z:*         $r \times c$

## Diagnostics

None.

## Also see

[M-5] [edittozero\(\)](#) — Edit matrix for roundoff error (zeros)

[M-4] [manipulation](#) — Matrix manipulation

## Syntax

```
numeric matrix  edittozero(numeric matrix Z, real scalar amt)
void            _edittozero(numeric matrix Z, real scalar amt)

numeric matrix  edittozerotol(numeric matrix Z, real scalar tol)
void            _edittozerotol(numeric matrix Z, real scalar tol)
```

## Description

These edit functions set elements of matrices to zero that are close to zero. `edittozero(Z, amt)` and `_edittozero(Z, amt)` set

$$Z_{ij} = 0 \quad \text{if } |Z_{ij}| \leq |tol|$$

for  $Z$  real and set

$$\begin{aligned} \operatorname{Re}(Z_{ij}) &= 0 & \text{if } |\operatorname{Re}(Z_{ij})| &\leq |tol| \\ \operatorname{Im}(Z_{ij}) &= 0 & \text{if } |\operatorname{Im}(Z_{ij})| &\leq |tol| \end{aligned}$$

for  $Z$  complex, where in both cases

$$tol = \operatorname{abs}(amt) * \operatorname{epsilon}(\operatorname{sum}(\operatorname{abs}(Z)) / (\operatorname{rows}(Z) * \operatorname{cols}(Z)))$$

`edittozero()` leaves  $Z$  unchanged and returns the edited matrix. `_edittozero()` edits  $Z$  in place.

`edittozerotol(Z, tol)` and `_edittozerotol(Z, tol)` do the same thing, except that  $tol$  is specified directly.

## Remarks

Remarks are presented under the following headings:

[Background](#)  
[Treatment of complex values](#)  
[Recommendations](#)

## Background

Numerical roundoff error leads to, among other things, numbers that should be zero being small but not zero, and so it is sometimes desirable to reset those small numbers to zero.

The problem is in identifying those small numbers. Is  $1e-14$  small? Is 10,000? The answer is that, given some matrix,  $1e-14$  might not be small because most of the values in the matrix are around  $1e-14$ , and the small values are  $1e-28$ , and given some other matrix, 10,000 might indeed be small because most of the elements are around  $1e+18$ .

`edittozero()` makes an attempt to determine what is small. `edittozerotol()` leaves that determination to you. In `edittozerotol(Z, tol)`, you specify *tol* and elements for which  $|Z_{ij}| \leq tol$  are set to zero.

Using `edittozero(Z, amt)`, however, you specify *amt* and then *tol* is calculated for you based on the size of the elements in *Z* and *amt*, using the formula

$$tol = amt * \text{epsilon}(\text{average value of } |Z_{ij}|)$$

`epsilon()` refers to machine precision, and `epsilon(x)` is the function that returns machine precision in units of *x*:

$$\text{epsilon}(x) = |x| * \text{epsilon}(1)$$

where `epsilon(1)` is approximately  $2.22e-16$  on most computers; see [M-5] `epsilon()`.

## Treatment of complex values

The formula

$$tol = amt * \text{epsilon}(\text{average value of } |Z_{ij}|)$$

is used for both real and complex values. For complex,  $|Z_{ij}|$  refers to the modulus (length) of the complex element.

However, rather than applying the reset rule

$$Z_{ij} = 0 \quad \text{if } |Z_{ij}| \leq |tol|$$

as is done when *Z* is real, the reset rules are

$$\begin{aligned} \text{Re}(Z_{ij}) &= 0 & \text{if } |\text{Re}(Z_{ij})| &\leq |tol| \\ \text{Im}(Z_{ij}) &= 0 & \text{if } |\text{Im}(Z_{ij})| &\leq |tol| \end{aligned}$$

The first rule, applied even for complex, may seem more appealing, but the use of the second has the advantage of mapping numbers close to being purely real or purely imaginary to purely real or purely imaginary results.

## Recommendations

1. Minimal editing is performed by `edittozero(Z, 1)`. Values less than  $2.22e-16$  times the average would be set to zero.
2. It is often reasonable to code `edittozero(Z, 1000)`, which sets to zero values less than  $2.22e-13$  times the average.
3. For a given calculation, the amount of roundoff error that arises with complex matrices (matrices with nonzero imaginary part) is greater than the amount that arises with real matrices (matrices with zero imaginary part even if stored as complex). That is because, in addition to all the usual sources of roundoff error, multiplication of complex values involves the addition operator, which introduces additional roundoff error. Hence, whatever is the appropriate value of *amt* or *tol* with real matrices, it is larger for complex matrices.

# Conformability

```
edittozero(Z, amt):
    Z:       $r \times c$ 
    amt:     $1 \times 1$ 
    result:  $r \times c$ 
```

```
_edittozero(Z, amt):
    input:
        Z:       $r \times c$ 
        amt:     $1 \times 1$ 
    output:
        Z:       $r \times c$ 
```

```
edittozerotol(Z, tol):
    Z:       $r \times c$ 
    tol:     $1 \times 1$ 
    result:  $r \times c$ 
```

```
_edittozerotol(Z, tol):
    input:
        Z:       $r \times c$ 
        tol:     $1 \times 1$ 
    output:
        Z:       $r \times c$ 
```

# Diagnostics

`edittozero(Z, amt)` and `_edittozero(Z, amt)` leave scalars unchanged because they base their calculation of the likely roundoff error on the average value of  $|Z_{ij}|$ .

# Also see

[M-5] [edittoint\(\)](#) — Edit matrix for roundoff error (integers)

[M-4] [manipulation](#) — Matrix manipulation

# Title

[M-5] **editvalue()** — Edit (change) values in matrix

## Syntax

*matrix*    `editvalue(matrix A, scalar from, scalar to)`

*void*    `_editvalue(matrix A, scalar from, scalar to)`

where *A*, *from*, and *to* may be real, complex, or string.

## Description

`editvalue(A, from, to)` returns *A* with all elements equal to *from* changed to *to*.

`_editvalue(A, from, to)` does the same thing but modifies *A* itself.

## Remarks

`editvalue()` and `_editvalue()` are fast.

If you wish to change missing values to nonmissing values, it is better to use [\[M-5\] editmissing\(\)](#). `editvalue(A, ., 1)` would change all . missing values to 1 but leave .a, .b, ..., .z unchanged. `editmissing(A, 1)` would change all missing values to 1.

## Conformability

`editvalue(A, from, to):`

<i>A</i> :	$r \times c$
<i>from</i> :	$1 \times 1$
<i>to</i> :	$1 \times 1$
<i>result</i> :	$r \times c$

`_editvalue(A, from, to):`

*input*:

<i>A</i> :	$r \times c$
<i>from</i> :	$1 \times 1$
<i>to</i> :	$1 \times 1$

*output*:

<i>A</i> :	$r \times c$
------------	--------------

## Diagnostics

`editvalue(A, from, to)` returns a matrix of the same type as *A*.

`editvalue(A, from, to)` and `_editvalue(A, from, to)` abort with error if *from* and *to* are incompatible with *A*. That is, if *A* is real, *to* and *from* must be real. If *A* is complex, *to* and *from* must each be either real or complex. If *A* is string, *to* and *from* must be string.

`_editvalue(A, from, to)` aborts with error if *A* is a view.

## Also see

[M-5] [editmissing\(\)](#) — Edit matrix for missing values

[M-4] [manipulation](#) — Matrix manipulation



## Syntax

```

void                eigensystem(A, X, L [, rcond [, nobalance]])
void                lefteigensystem(A, X, L [, rcond [, nobalance]])
complex rowvector   eigenvalues(A          [, rcond [, nobalance]])

void                symeigensystem(A, X, L)
real rowvector      symeigenvalues(A)

void                _eigensystem(A, X, L [, rcond [, nobalance]])
void                _lefteigensystem(A, X, L [, rcond [, nobalance]])
complex rowvector   _eigenvalues(A          [, rcond [, nobalance]])
void                _symeigensystem(A, X, L)
real rowvector      _symeigenvalues(A)
    
```

where inputs are

*A*:    *numeric matrix*  
*rcond*: *real scalar* (whether *rcond* desired)  
*nobalance*: *real scalar* (whether to suppress balancing)

and outputs are

*X*:    *numeric matrix* of eigenvectors  
*L*:    *numeric vector* of eigenvalues  
*rcond*: *real vector* of reciprocal condition numbers

The columns of *X* will contain the eigenvectors except when using `_lefteigensystem()`, in which case the rows of *X* contain the eigenvectors.

The following routines are used in implementing the above routines:

```

real scalar _eigen_la(real scalar todo, numeric matrix A, X, L, real scalar rcond,
                      real scalar nobalance)

real scalar _symeigen_la(real scalar todo, numeric matrix A, X, L)
    
```

## Description

These routines calculate eigenvectors and eigenvalues of square matrix *A*.

`eigensystem(A, X, L, rcond, nobalance)` calculates eigenvectors and eigenvalues of a general, real or complex, square matrix  $A$ . Eigenvectors are returned in  $X$  and eigenvalues in  $L$ . The remaining arguments are optional:

1. If *rcond* is not specified, then reciprocal condition numbers are not returned in *rcond*.

If *rcond* is specified and contains a value other than 0 or missing—*rcond*=1 is suggested—in *rcond* will be placed a vector of the reciprocals of the condition numbers for the eigenvalues. Each element of the new *rcond* measures the accuracy to which the corresponding eigenvalue has been calculated; large numbers (numbers close to 1) are better and small numbers (numbers close to 0) indicate inaccuracy; see [Eigenvalue condition](#) below.

2. If *nobalance* is not specified, balancing is performed to obtain more accurate results.

If *nobalance* is specified and is not zero nor missing, balancing is not used. Results are calculated more quickly, but perhaps a little less accurately; see [Balancing](#) below.

`lefteigensystem(A, X, L, rcond, nobalance)` mirrors `eigensystem()`, the difference being that `lefteigensystem()` solves for left eigenvectors solving  $XA = \text{diag}(L)*X$  instead of right eigenvectors solving  $AX = X*\text{diag}(L)$ .

`eigenvalues(A, rcond, nobalance)` returns the eigenvalues of square matrix  $A$ ; the eigenvectors are not calculated. Arguments *rcond* and *nobalance* are optional.

`syeigensystem(A, X, L)` and `syeigenvalues(A)` mirror `eigensystem()` and `eigenvalues()`, the difference being that  $A$  is assumed to be symmetric (Hermitian). The eigenvalues returned are real. (Arguments *rcond* and *nobalance* are not allowed; *rcond* because symmetric matrices are inherently well conditioned; *nobalance* because it is unnecessary.)

The underscore routines mirror the routines of the same name without underscores, the difference being that  $A$  is damaged during the calculation and so the underscore routines use less memory.

`_eigen_la()` and `_syeigen_la()` are the interfaces into the [M-1] **LAPACK** routines used to implement the above functions. Their direct use is not recommended.

## Remarks

Remarks are presented under the following headings:

[Eigenvalues and eigenvectors](#)  
[Left eigenvectors](#)  
[Symmetric eigensystems](#)  
[Normalization and order](#)  
[Eigenvalue condition](#)  
[Balancing](#)  
[eigensystem\(\) and eigenvalues\(\)](#)  
[lefteigensystem\(\)](#)  
[syeigensystem\(\) and syeigenvalues\(\)](#)

## Eigenvalues and eigenvectors

A scalar  $\lambda$  is said to be an eigenvalue of square matrix  $A$ :  $n \times n$  if there is a nonzero column vector  $x$ :  $n \times 1$  (called the eigenvector) such that

$$Ax = \lambda x \quad (1)$$

(1) can also be written

$$(A - \lambda I)x = 0$$

where  $I$  is the  $n \times n$  identity matrix. A nontrivial solution to this system of  $n$  linear homogeneous equations exists if and only if

$$\det(A - \lambda I) = 0 \quad (2)$$

This  $n$ th degree polynomial in  $\lambda$  is called the characteristic polynomial or characteristic equation of  $A$ , and the eigenvalues  $\lambda$  are its roots, also known as the characteristic roots.

There are, in fact,  $n$  solutions  $(\lambda_i, x_i)$  that satisfy (1)—although some can be repeated—and we can compactly write the full set of solutions as

$$AX = X * \text{diag}(L) \quad (3)$$

where

$$X = (x_1, x_2, \dots) \quad (X : n \times n)$$

$$L = (\lambda_1, \lambda_2, \dots) \quad (L : 1 \times n)$$

```
: A = (1, 2 \ 9, 4)
: X = .
: L = .
: eigensystem(A, X, L)
: X
```

	1	2
1	-.316227766	-.554700196
2	-.948683298	.832050294

```
: L
```

	1	2
1	7	-2

The first eigenvalue is 7, and the corresponding eigenvector is  $(-.316 \setminus -.949)$ . The second eigenvalue is  $-2$ , and the corresponding eigenvector is  $(-.555 \setminus .832)$ .

In general, eigenvalues and vectors can be complex even if  $A$  is real.

## Left eigenvectors

What we have defined above is properly known as the right-eigensystem problem:

$$Ax = \lambda x \quad (1)$$

In the above,  $x$  is a column vector. The left-eigensystem problem is to find the row vector  $x$  satisfying

$$xA = \lambda x \tag{1'}$$

The eigenvalue  $\lambda$  is the same in (1) and (1'), but  $x$  can differ.

The  $n$  solutions  $(\lambda_i, x_i)$  that satisfy (1') can be compactly written

$$XA = \text{diag}(L) * X \tag{3'}$$

where

$$X = \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_n \end{bmatrix}$$

$n \times n$

$$L = \begin{bmatrix} \lambda_1 \\ \lambda_2 \\ \vdots \\ \lambda_n \end{bmatrix}$$

$n \times 1$

For instance,

```
: A = (1, 2 \ 9, 4)
: X = .
: L = .
: lefteigensystem(A, X, L)
: X
      1      2
1  - .832050294  - .554700196
2  - .948683298   .316227766

: L
      1
1      7
2     -2
```

The first eigenvalue is 7, and the corresponding eigenvector is  $(-.832, -.555)$ . The second eigenvalue is  $-2$ , and the corresponding eigenvector is  $(-.949, .316)$ .

The eigenvalues are the same as in the previous example; the eigenvectors are different.

Symmetric eigensystems

Below we use the term symmetric to encompass [Hermitian matrices](#), even when we do not emphasize the fact.

Eigensystems of symmetric matrices are conceptually no different from general eigensystems, but symmetry introduces certain simplifications:

1. The eigenvalues associated with symmetric matrices are real, whereas those associated with general matrices may be real or complex.
2. The eigenvectors associated with symmetric matrices—which may be real or complex—are orthogonal.

3. The left and right eigenvectors of symmetric matrices are transposes of each other.
4. The eigenvectors and eigenvalues of symmetric matrices are more easily, and more accurately, computed.

For item 3, let us begin with the right-eigensystem problem:

$$AX = X * \text{diag}(L)$$

Taking the transpose of both sides results in

$$X'A = \text{diag}(L) * X'$$

because  $A = A'$  if  $A$  is symmetric (Hermitian).

`symeigensystem(A, X, L)` calculates right eigenvectors. To obtain the left eigenvectors, you simply transpose  $X$ .

## Normalization and order

If  $x$  is a solution to

$$Ax = \lambda x$$

then so is  $cx$ ,  $c: 1 \times 1$ ,  $c \neq 0$ .

The eigenvectors returned by the above routines are scaled to have length (norm) 1.

The eigenvalues are combined and returned in a vector ( $L$ ) and the eigenvectors in a matrix ( $X$ ). The eigenvalues are ordered from largest to smallest in absolute value (or, if the eigenvalues are complex, in length). The eigenvectors are ordered to correspond to the eigenvalues.

## Eigenvalue condition

Optional argument *rcond* may be specified as a value other than 0 or missing—*rcond* = 1 is suggested—and then *rcond* will be filled in with a vector containing the reciprocals of the condition numbers for the eigenvalues. Each element of *rcond* measures the accuracy with which the corresponding eigenvalue has been calculated; large numbers (numbers close to 1) are better and small numbers (numbers close to 0) indicate inaccuracy.

The reciprocal condition number is calculated as  $\text{abs}(y*x)$ , where  $y: 1 \times n$  is the left eigenvector and  $x: n \times 1$  is the corresponding right eigenvector. Since  $y$  and  $x$  each have norm 1,  $\text{abs}(y*x) = \text{abs}(\cos(\theta))$ , where  $\theta$  is the angle made by the vectors. Thus  $0 \leq \text{abs}(y*x) \leq 1$ . For symmetric matrices,  $y*x$  will equal 1. It can be proved that  $\text{abs}(y*x)$  is the reciprocal of the condition number for a simple eigenvalue, and so it turns out that the sensitivity of the eigenvalue to a perturbation is a function of how far the matrix is from symmetric on this scale.

Requesting that *rcond* be calculated increases the amount of computation considerably.

## Balancing

By default, balancing is performed for general matrices. Optional argument *nobalance* allows you to suppress balancing.

Balancing is related to row-and-column equilibration; see [M-5] `_equilrc()`. Here, however, a diagonal matrix  $D$  is found such that  $DAD^{-1}$  is better balanced, the eigenvectors and eigenvalues for  $DAD^{-1}$  are extracted, and then the eigenvectors and eigenvalues are adjusted by  $D$  so that they reflect those for the original  $A$  matrix.

There is no gain from these machinations when  $A$  is symmetric, so the symmetric routines do not have a *nobalance* argument.

## eigensystem() and eigenvalues()

1. Use  $L = \text{eigenvalues}(A)$  and  $\text{eigensystem}(A, X, L)$  for general matrices  $A$ .
2. Use  $L = \text{eigenvalues}(A)$  when you do not need the eigenvectors; it will save both time and memory.
3. The eigenvalues returned by  $L = \text{eigenvalues}(A)$  and by  $\text{eigensystem}(A, X, L)$  are of storage type complex even if the eigenvalues are real (that is, even if  $\text{Im}(L) == 0$ ). If the eigenvalues are known to be real, you can save computer memory by subsequently coding

```
L = Re(L)
```

If you wish to test whether the eigenvalues are real, examine `mrldifre(L)`; see [M-5] `reldif()`.

4. The eigenvectors returned by  $\text{eigensystem}(A, X, L)$  are of storage type complex even if the eigenvectors are real (that is, even if  $\text{Im}(X) == 0$ ). If the eigenvectors are known to be real, you can save computer memory by subsequently coding

```
X = Re(X)
```

If you wish to test whether the eigenvectors are real, examine `mrldifre(X)`; see [M-5] `reldif()`.

5. If you are using  $\text{eigensystem}(A, X, L)$  interactively (outside a program),  $X$  and  $L$  must be predefined. Type

```
: eigensystem(A, X=., L=.)
```

## lefteigensystem()

What was just said about `eigensystem()` applies equally well to `lefteigensystem()`.

If you need only the eigenvalues, use  $L = \text{eigenvalues}(A)$ . The eigenvalues are the same for both left and right systems.

**symeigensystem() and symeigenvalues()**

1. Use  $L = \text{symeigenvalues}(A)$  and  $\text{symeigensystem}(A, X, L)$  for symmetric or Hermitian matrices  $A$ .
2. Use  $L = \text{symeigenvalues}(A)$  when you do not need the eigenvectors; it will save both time and memory.
3. The eigenvalues returned by  $L = \text{symeigenvalues}(A)$  and by  $\text{symeigensystem}(A, X, L)$  are of storage type real. Eigenvalues of symmetric and Hermitian matrices are always real.
4. The eigenvectors returned by  $\text{symeigensystem}(A, X, L)$  are of storage type real when  $A$  is of storage type real and of storage type complex when  $A$  is of storage type complex.
5. If you are using  $\text{symeigensystem}(A, X, L)$  interactively (outside a program),  $X$  and  $L$  must be predefined. Type

```
: symeigensystem(A, X=., L=.)
```

**Conformability**

$\text{eigensystem}(A, X, L, rcond, nobalance)$ :

*input:*

$A$ :	$n \times n$	
$rcond$ :	$1 \times 1$	(optional, specify as 1 to obtain $rcond$ )
$nobalance$ :	$1 \times 1$	(optional, specify as 1 to prevent balancing)

*output:*

$X$ :	$n \times n$	(columns contain eigenvectors)
$L$ :	$1 \times n$	
$rcond$ :	$1 \times n$	(optional)
$result$ :	$void$	

$\text{lefteigensystem}(A, X, L, rcond, nobalance)$ :

*input:*

$A$ :	$n \times n$	
$rcond$ :	$1 \times 1$	(optional, specify as 1 to obtain $rcond$ )
$nobalance$ :	$1 \times 1$	(optional, specify as 1 to prevent balancing)

*output:*

$X$ :	$n \times n$	(rows contain eigenvectors)
$L$ :	$n \times 1$	
$rcond$ :	$n \times 1$	(optional)
$result$ :	$void$	

$\text{eigenvalues}(A, rcond, nobalance)$ :

*input:*

$A$ :	$n \times n$	
$rcond$ :	$1 \times 1$	(optional, specify as 1 to obtain $rcond$ )
$nobalance$ :	$1 \times 1$	(optional, specify as 1 to prevent balancing)

*output:*

$rcond$ :	$1 \times n$	(optional)
$result$ :	$1 \times n$	(contains eigenvalues)

`symeigensystem(A, X, L):`

*input:*

$A: \quad n \times n$

*output:*

$X: \quad n \times n$  (columns contain eigenvectors)

$L: \quad 1 \times n$

*result:* `void`

`symeigenvalues(A):`

*input:*

$A: \quad n \times n$

*output:*

*result:*  $1 \times n$  (contains eigenvalues)

`_eigensystem(A, X, L, rcond, nobalance):`

*input:*

$A: \quad n \times n$

$rcond: \quad 1 \times 1$  (optional, specify as 1 to obtain *rcond*)

$nobalance: \quad 1 \times 1$  (optional, specify as 1 to prevent balancing)

*output:*

$A: \quad 0 \times 0$

$X: \quad n \times n$  (columns contain eigenvectors)

$L: \quad 1 \times n$

$rcond: \quad 1 \times n$  (optional)

*result:* `void`

`_lefteigensystem(A, X, L, rcond, nobalance):`

*input:*

$A: \quad n \times n$

$rcond: \quad 1 \times 1$  (optional, specify as 1 to obtain *rcond*)

$nobalance: \quad 1 \times 1$  (optional, specify as 1 to prevent balancing)

*output:*

$A: \quad 0 \times 0$

$X: \quad n \times n$  (rows contain eigenvectors)

$L: \quad n \times 1$

$rcond: \quad n \times 1$  (optional)

*result:* `void`

`_eigenvalues(A, rcond, nobalance):`

*input:*

$A: \quad n \times n$

$rcond: \quad 1 \times 1$  (optional, specify as 1 to obtain *rcond*)

$nobalance: \quad 1 \times 1$  (optional, specify as 1 to prevent balancing)

*output:*

$A: \quad 0 \times 0$

$rcond: \quad 1 \times n$  (optional)

*result:*  $1 \times n$  (contains eigenvalues)



`_symeigensystem(A, X, L):`

*input:*

$A: n \times n$

*output:*

$A: 0 \times 0$

$X: n \times n$  (columns contain eigenvectors)

$L: 1 \times n$

*result:* void

`_symeigenvalues(A):`

*input:*

$A: n \times n$

*output:*

$A: 0 \times 0$

*result:*  $1 \times n$  (contains eigenvalues)

`_eigen_la(todo, A, X, L, rcond, nobalance):`

*input:*

$todo: 1 \times 1$

$A: n \times n$

$rcond: 1 \times 1$

$nobalance: 1 \times 1$

*output:*

$A: 0 \times 0$

$X: n \times n$

$L: 1 \times n$  or  $n \times 1$

$rcond: 1 \times n$  or  $n \times 1$  (optional)

*result:*  $1 \times 1$  (return code)

`_symeigen_la(todo, A, X, L):`

*input:*

$todo: 1 \times 1$

$A: n \times n$

*output:*

$A: 0 \times 0$

$X: n \times n$

$L: 1 \times n$

*result:*  $1 \times 1$  (return code)

## Diagnostics

All functions return missing-value results if  $A$  has missing values.

`symeigensystem()`, `symeigenvalues()`, `_symeigensystem()`, and `_symeigenvalues()` use the lower triangle of  $A$  without checking for symmetry. When  $A$  is complex, only the real part of the diagonal is used.

## References

Gould, W. W. 2011a. Understanding matrices intuitively, part 1. The Stata Blog: Not Elsewhere Classified. <http://blog.stata.com/2011/03/03/understanding-matrices-intuitively-part-1/>

——. 2011b. Understanding matrices intuitively, part 2, eigenvalues and eigenvectors. The Stata Blog: Not Elsewhere Classified. <http://blog.stata.com/2011/03/09/understanding-matrices-intuitively-part-2/>

## Also see

[M-5] **matexpsym()** — Exponentiation and logarithms of symmetric matrices

[M-5] **matpowersym()** — Powers of a symmetric matrix

[M-4] **matrix** — Matrix functions

## Syntax

```

void      eigensystemselectr(A, range, X, L)
void lefteigensystemselectr(A, range, X, L)
void      eigensystemselecti(A, index, X, L)
void lefteigensystemselecti(A, index, X, L)
void      eigensystemselectf(A, f, X, L)
void lefteigensystemselectf(A, f, X, L)
void symeigensystemselectr(A, range, X, L)
void symeigensystemselecti(A, index, X, L)
    
```

where inputs are

*A*: numeric matrix  
*range*: real vector (range of eigenvalues to be selected)  
*index*: real vector (indices of eigenvalues to be selected)  
*f*: pointer scalar (points to a function used to select eigenvalue)

and outputs are

*X*: numeric matrix of eigenvectors  
*L*: numeric vector of eigenvalues

The following routines are used in implementing the above routines:

```

void _eigenselecti_la(numeric matrix A, XL, XR, L,
                     string scalar side, real vector index)
void _eigenselectr_la(numeric matrix A, XL, XR, L,
                     string scalar side, real vector range)
void _eigenselectf_la(numeric matrix A, XL, XR, L,
                     string scalar side, pointer scalar f)
real scalar _eigenselect_la(numeric matrix A, XL, XR, L, select,
                          string scalar side, real scalar nroflop)
real scalar _symeigenselect_la(numeric matrix A, X, L, ifail,
                              real scalar type, lower, upper, abstol)
    
```

## Description

`eigensystemselectr(A, range, X, L)` computes selected right eigenvectors of a square, numeric matrix *A* along with their corresponding eigenvalues. Only the eigenvectors corresponding to selected eigenvalues are computed. Eigenvalues that lie in a [range](#) are selected. The selected eigenvectors are returned in *X*, and their corresponding eigenvalues are returned in *L*.

*range* is a vector of length 2. All eigenvalues with absolute value in the half-open interval (*range*[1], *range*[2]] are selected.

`lefteigensystemselectr(A, range, X, L)` mirrors `eigensystemselectr()`, the difference being that it computes selected left eigenvectors instead of selected right eigenvectors.

`eigensystemselecti(A, index, X, L)` computes selected right eigenvectors of a square, numeric matrix, *A*, along with their corresponding eigenvalues. Only the eigenvectors corresponding to selected eigenvalues are computed. Eigenvalues are selected by an [index](#). The selected eigenvectors are returned in *X*, and the selected eigenvalues are returned in *L*.

*index* is a vector of length 2. The eigenvalues are sorted by their absolute values, in descending order. The eigenvalues whose rank is *index*[1] through *index*[2], inclusive, are selected.

`lefteigensystemselecti(A, index, X, L)` mirrors `eigensystemselecti()`, the difference being that it computes selected left eigenvectors instead of selected right eigenvectors.

`eigensystemselectf(A, f, X, L)` computes selected right eigenvectors of a square, numeric matrix, *A*, along with their corresponding eigenvalues. Only the eigenvectors corresponding to selected eigenvalues are computed. Eigenvalues are selected by a user-written function described [below](#). The selected eigenvectors are returned in *X*, and the selected eigenvalues are returned in *L*.

`lefteigensystemselectf(A, f, X, L)` mirrors `eigensystemselectf()`, the difference being that it computes selected left eigenvectors instead of selected right eigenvectors.

`symeigensystemselectr(A, range, X, L)` computes selected eigenvectors of a symmetric (Hermitian) matrix, *A*, along with their corresponding eigenvalues. Only the eigenvectors corresponding to selected eigenvalues are computed. Eigenvalues that lie in a [range](#) are selected. The selected eigenvectors are returned in *X*, and their corresponding eigenvalues are returned in *L*.

`symeigensystemselecti(A, index, X, L)` computes selected eigenvectors of a symmetric (Hermitian) matrix, *A*, along with their corresponding eigenvalues. Only the eigenvectors corresponding to selected eigenvalues are computed. Eigenvalues are selected by an [index](#). The selected eigenvectors are returned in *X*, and the selected eigenvalues are returned in *L*.

`_eigensystemselectr_la()`, `_eigensystemselecti_la()`, `_eigensystemselectf_la()`, `_symeigensystemselectr_la()`, and `_symeigensystemselecti_la()` are the interfaces into the [\[M-1\] LAPACK](#) routines used to implement the above functions. Their direct use is not recommended.

## Remarks

[Introduction](#)  
[Range selection](#)  
[Index selection](#)  
[Criterion selection](#)  
[Other functions](#)

## Introduction

These routines compute subsets of the available eigenvectors. This computation can be much faster than computing all the eigenvectors. (See [\[M-5\] eigensystem\(\)](#) for routines to compute all the eigenvectors and an introduction to the eigensystem problem.)

There are three methods for selecting which eigenvectors to compute; all of them are based on the corresponding eigenvalues. First, we can select only those eigenvectors whose eigenvalues have absolute values that fall in a half-open interval. Second, we can select only those eigenvectors whose eigenvalues have certain indices, after sorting the eigenvalues by their absolute values in descending order. Third, we can select only those eigenvectors whose eigenvalues meet a criterion encoded in a function.

Below we illustrate each of these methods. For comparison purposes, we begin by computing all the eigenvectors of the matrix

```
: A
```

	1	2	3	4
1	.31	.69	.13	.56
2	.31	.5	.72	.42
3	.68	.37	.71	.8
4	.09	.16	.83	.9

We perform the computation with `eigensystem()`:

```
: eigensystem(A, X=., L=.)
```

The absolute values of the eigenvalues are

```
: abs(L)
```

	1	2	3	4
1	2.10742167	.4658398402	.4005757984	.4005757984

The corresponding eigenvectors are

```
: X
```

	1	2	3
1	.385302069	-.394945842	.672770333
2	.477773165	-.597299386	-.292386384 - .171958335i
3	.604617181	-.192938403	-.102481414 + .519705293i
4	.50765459	.670839771	-.08043663 - .381122722i

	4
1	.672770333
2	-.292386384 + .171958335i
3	-.102481414 - .519705293i
4	-.08043663 + .381122722i

## Range selection

In applications, an eigenvalue whose absolute value is greater than 1 frequently corresponds to an explosive solution, whereas an eigenvalue whose absolute value is less than 1 corresponds to a stable solution. We frequently want to use only the eigenvectors from the stable solutions, which we can do using `eigensystemselectr()`. We begin by specifying

```
: range = (-1, .999999999)
```

which starts from  $-1$  to include 0 and stops at .999999999 to exclude 1. (The half-open interval in range is open on the left and closed on the right.)

Using this range in eigensystemselectr() requests each eigenvector for which the absolute value of the corresponding eigenvalue falls in the interval  $(-1, .999999999]$ . For the example at hand, we have

```
: eigensystemselectr(A, range, X=., L=.)
: X
      1                2                3
1  -.442004357    .201218963 - .875384534i    .201218963 + .875384534i
2  -.668468693    .136296114 + .431873675i    .136296114 - .431873675i
3  -.215927364   -.706872994 - .022093594i   -.706872994 + .022093594i
4   .750771548    .471845361 + .218651289i    .471845361 - .218651289i

: L
      1                2                3
1  .46583984   -.076630755 + .393177692i   -.076630755 - .393177692i

: abs(L)
      1                2                3
1  .4658398402   .4005757984   .4005757984
```

The above output illustrates that eigensystemselectr() has not included the results for the eigenvalue whose absolute value is greater than 1, as desired.

Index selection

In many statistical applications, an eigenvalue measures the importance of an eigenvector factor. In these applications, we want only to compute several of the largest eigenvectors. Here we use eigensystemselecti() to compute the eigenvectors corresponding to the two largest eigenvalues:

```
: index = (1, 2)
: eigensystemselecti(A, index, X=., L=.)
: L
      1                2
1  2.10742167    .46583984

: X
      1                2
1  .385302069   -.442004357
2  .477773165   -.668468693
3  .604617181   -.215927364
4  .50765459    .750771548
```

## Criterion selection

In some applications, we want to compute only those eigenvectors whose corresponding eigenvalues satisfy a more complicated criterion. We can use `eigensystemselectf()` to solve these problems.

We must pass `eigensystemselectf()` a [pointer to a function](#) that implements our criterion. The function must accept a complex scalar argument so that it can receive an eigenvalue, and it must return the real value 0 to indicate rejection and a nonzero real value to indicate selection.

In the example below, we consider the common criterion of whether the eigenvalue is real. We want only to compute the eigenvectors corresponding to real eigenvalues. After deciding that anything smaller than  $1e-15$  is zero, we define our function to be

```
: real scalar onlyreal(complex scalar ev)
> {
>     return( (abs(Im(ev))<1e-15) )
> }
```

We compute only the eigenvectors corresponding to the real eigenvalues by typing

```
: eigensystemselectf(A, &onlyreal(), X=., L=.)
```

The eigenvalues that satisfy this criterion and their corresponding eigenvectors are

```
: L
      1      2
1  [ 2.10742167  .46583984 ]

: X
      1      2
1  [ .385302069  -.442004357 ]
2  [ .477773165  -.668468693 ]
3  [ .604617181  -.215927364 ]
4  [ .50765459   .750771548 ]
```

## Other functions

`lefteigensystemselectr()` and `symeigensystemselectr()` use a *range* like `eigensystemselectr()`.

`lefteigensystemselecti()` and `symeigensystemselecti()` use an *index* like `eigensystemselecti()`.

`lefteigensystemselectf()` uses a pointer to a function like `eigensystemselectf()`.

## Conformability

`eigensystemselectr(A, range, X, L):`

*input:*

$A:$       $n \times n$   
 $range:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $n \times m$   
 $L:$       $1 \times m$

`lefteigensystemselectr(A, range, X, L):`

*input:*

$A:$       $n \times n$   
 $range:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $m \times n$   
 $L:$       $1 \times m$

`eigensystemselecti(A, index, X, L):`

*input:*

$A:$       $n \times n$   
 $index:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $n \times m$   
 $L:$       $1 \times m$

`lefteigensystemselecti(A, index, X, L):`

*input:*

$A:$       $n \times n$   
 $index:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $m \times n$   
 $L:$       $1 \times m$

`eigensystemselectf(A, f, X, L):`

*input:*

$A:$       $n \times n$   
 $f:$       $1 \times 1$

*output:*

$X:$       $n \times m$   
 $L:$       $1 \times m$



lefteigensystemselectf( $A, f, X, L$ ):

*input:*

$A:$   $n \times n$

$f:$   $1 \times 1$

*output:*

$X:$   $m \times n$

$L:$   $1 \times m$

symeigensystemselectr( $A, range, X, L$ ):

*input:*

$A:$   $n \times n$

$range:$   $1 \times 2$  or  $2 \times 1$

*output:*

$X:$   $n \times m$

$L:$   $1 \times m$

symeigensystemselecti( $A, index, X, L$ ):

*input:*

$A:$   $n \times n$

$index:$   $1 \times 2$  or  $2 \times 1$

*output:*

$X:$   $n \times m$

$L:$   $1 \times m$

## Diagnostics

All functions return missing-value results if  $A$  has missing values.

`symeigensystemselectr()` and `symeigensystemselecti()` use the lower triangle of  $A$  without checking for symmetry. When  $A$  is complex, only the real part of the diagonal is used.

If the  $i$ th eigenvector failed to converge, `symeigensystemselectr()` and `symeigensystemselecti()` insert a vector of missing values into the  $i$ th column of the returned eigenvector matrix.

## Also see

[M-1] **LAPACK** — The LAPACK linear-algebra routines

[M-5] **eigensystem()** — Eigenvectors and eigenvalues

[M-5] **matexpsym()** — Exponentiation and logarithms of symmetric matrices

[M-5] **matpowersym()** — Powers of a symmetric matrix

[M-4] **matrix** — Matrix functions

[M-5] `eltype()` — Element type and organizational type of object

Syntax

```
string scalar  eltype(X)
string scalar  orgtype(X)
```

Description

`eltype()` returns the current *eltype* of the argument.  
`orgtype()` returns the current *orgtype* of the argument.  
See [M-6] [Glossary](#) for a definition of *eltype* and *orgtype*.

Remarks

If *X* is a matrix (syntax 1), returned is

eltype( <i>X</i> )	orgtype( <i>X</i> )
real	scalar
complex	rowvector
string	colvector
pointer	matrix
struct	
class	

The returned value reflects the current contents of *X*. That is, *X* might be declared a *transmorphic matrix*, but at any instant, it contains something, and if that something were 5, returned would be "real" and "scalar".

For `orgtype()`, returned is "scalar" if the object is currently  $1 \times 1$ ; "rowvector" if it is  $1 \times k$ ,  $k \neq 1$ ; "colvector" if it is  $k \times 1$ ,  $k \neq 1$ ; and "matrix" otherwise (it is  $r \times c$ ,  $r \neq 1$ ,  $c \neq 1$ ).

*X* can be a function (syntax 2). Returned is

eltype(*(&func()))	orgtype(*(&func()))
transmorphic	matrix
numeric	vector
real	rowvector
complex	colvector
string	scalar
pointer	void
struct	
structdef	
class	
classdef	

These types are obtained from the declaration of the function.

Aside: `struct` and `structdef` have to do with structures; see [M-2] [struct](#). `structdef` indicates that the function not only returns a structure but is the routine that defines the structure as well. `class` and `classdef` have to do with Mata classes; see [M-2] [class](#). `classdef` indicates the function not only returns a class but is the routine that defines the class as well.

## Conformability

`eltype(X)`, `orgtype(X)`:

<i>X</i> :	$r \times c$
<i>result</i> :	$1 \times 1$

## Diagnostics

None.

## Also see

[M-5] [isreal\(\)](#) — Storage type of matrix

[M-5] [isview\(\)](#) — Whether matrix is view

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] **epsilon()** — Unit roundoff error (machine precision)

## Syntax

*real scalar*    **epsilon**(*real scalar x*)

## Description

**epsilon**(*x*) returns the unit roundoff error in quantities of size **abs**(*x*).

## Remarks

On all computers on which Stata and Mata are currently implemented—which are computers following IEEE standards—**epsilon**(1) is 1.0X–34, or about 2.22045e–16. This is the smallest amount by which a real number can differ from 1.

**epsilon**(*x*) is **abs**(*x*)\***epsilon**(1). This is an approximation of the smallest amount by which a real number can differ from *x*. The approximation is exact at integer powers of 2.

## Conformability

**epsilon**(*x*):

<i>x</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

**epsilon**(*x*) returns . if *x* is missing.

## Also see

[M-5] **mindouble()** — Minimum and maximum nonmissing value

[M-5] **edittozero()** — Edit matrix for roundoff error (zeros)

[M-4] **utility** — Matrix utility functions

## Syntax

```

void          _equilrc(numeric matrix A, r, c)
void          _equilr(numeric matrix A, r)
void          _equilc(numeric matrix A, c)

real scalar   _perhapsequilrc(numeric matrix A, r, c)
real scalar   _perhapsequilr(numeric matrix A, r)
real scalar   _perhapsequilc(numeric matrix A, c)

real colvector rowscalefactors(numeric matrix A)
real rowvector colscalefactors(numeric matrix A)

```

The types of *r* and *c* are irrelevant because they are overwritten.

## Description

`_equilrc(A, r, c)` performs row and column equilibration (balancing) on matrix *A*, returning the equilibrated matrix in *A*, the row-scaling factors in *r*, and the column-scaling factors in *c*.

`_equilr(A, r)` performs row equilibration on matrix *A*, returning the row-equilibrated matrix in *A* and the row-scaling factors in *r*.

`_equilc(A, c)` performs column equilibration on matrix *A*, returning the column-equilibrated matrix in *A* and the column-scaling factors in *c*.

`_perhapsequilrc(A, r, c)` performs row and/or column equilibration on matrix *A*—as is necessary and which decision is made by `_perhapsequilrc()`—returning the equilibrated matrix in *A*, the row-scaling factors in *r*, the column-scaling factors in *c*, and returning 0 (no equilibration performed), 1 (row equilibration performed), 2 (column equilibration performed), or 3 (row and column equilibration performed).

`_perhapsequilr(A, r)` performs row equilibration on matrix *A*—if necessary and which decision is made by `_perhapsequilr()`—returning the equilibrated matrix in *A*, the row-scaling factors in *r*, and returning 0 (no equilibration performed) or 1 (row equilibration performed).

`_perhapsequilc(A, c)` performs column equilibration on matrix *A*—if necessary and which decision is made by `_perhapsequilc()`—returning the equilibrated matrix in *A*, the column-scaling factors in *c*, and returning 0 (no equilibration performed) or 1 (column equilibration performed).

`rowscalefactors(A)` returns the row-scaling factors of *A*.

`colscalefactors(A)` returns the column-scaling factors of *A*.

# Remarks

Remarks are presented under the following headings:

- [Introduction](#)
- [Is equilibration necessary?](#)
- [The `\_equil\*\(\)` family of functions](#)
- [The `\_perhapsequil\*\(\)` family of functions](#)
- [`rowscalefactors\(\)` and `colscalefactors\(\)`](#)

# Introduction

Equilibration (also known as balancing) takes a matrix with poorly scaled rows and columns, such as

	1	2
1	1.00000e+10	5.00000e+10
2	2.00000e-10	8.00000e-10

and produces a related matrix, such as

	1	2
1	.2	1
2	.25	1

that will yield improved accuracy in, for instance, the solution to linear systems. The improved matrix above has been row equilibrated. All we did was find the maximum of each row of the original and then divide the row by its maximum. If we were to take the result and repeat the process on the columns—divide each column by the column’s maximum—we would obtain

	1	2
1	.8	1
2	1	1

which is the row-and-column equilibrated form of the original matrix.

In terms of matrix notation, equilibration can be thought about in terms of multiplication by diagonal matrices. The row-equilibrated form of  $A$  is  $RA$ , where  $R$  contains the reciprocals of the row maximums on its diagonal. The column-equilibrated form of  $A$  is  $AC$ , where  $C$  contains the reciprocals of the column maximums on its diagonal. The row-and-column equilibrated form of  $A$  is  $RAC$ , where  $R$  contains the reciprocals of the row maximums of  $A$  on its diagonal, and  $C$  contains the reciprocals of the column maximums of  $RA$  on its diagonal.

Say we wished to find the solution  $x$  to

$$Ax = b$$

We could compute the solution by solving for  $y$  in the equilibrated system

$$(RAC)y = Rb$$

and then setting

$$x = Cy$$

Thus routines that perform equilibration need to return to you, in some fashion,  $R$  and  $C$ . The routines here do that by returning  $r$  and  $c$ , the reciprocals of the maximums in vector form. You could obtain  $R$  and  $C$  from them by coding

$$R = \text{diag}(r)$$

$$C = \text{diag}(c)$$

but that is not in general necessary, and it is wasteful of memory. In code, you will need to multiply by  $R$  and  $C$ , and you can do that using the `:*` operator with  $r$  and  $c$ :

$$RA \leftrightarrow r:*A$$

$$AC \leftrightarrow A:*c$$

$$RAC \leftrightarrow r:*A:*c$$

## Is equilibration necessary?

Equilibration is not a panacea. Equilibration can reduce the condition number of some matrices and thereby improve the accuracy of the solution to linear systems, but equilibration is not guaranteed to reduce the condition number, and counterexamples exist in which equilibration actually decreases the accuracy of the solution. That said, you have to look long and hard to find such examples.

Equilibration is not especially computationally expensive, but neither is it cheap, especially when you consider the extra computational costs of using the equilibrated matrices. In statistical contexts, equilibration may buy you little because matrices are already nearly equilibrated. Data analysts know variables should be on roughly the same scale, and observations are assumed to be draws from an underlying distribution. The computational cost of equilibration is probably better spent somewhere else. For instance, consider obtaining regression estimates from  $X'X$  and  $X'y$ . The gain from equilibrating  $X'X$  and  $X'y$ , or even from equilibrating the original  $X$  matrix, is nowhere near that from the gain to be had in removal of the means before  $X'X$  and  $X'y$  are formed.

In the example in the previous section, we showed you a matrix that assuredly benefited from equilibration. Even so, after row equilibration, column equilibration was unnecessary. It is often the case that solely row or column equilibration is sufficient, and in those cases, although the extra equilibration will do no numerical harm, it will burn computer cycles. And, as we have already argued, some matrices do not need equilibration at all.

Thus programmers who want to use equilibration and obtain the best speed possible examine the matrix and on that basis perform (1) no equilibration, (2) row equilibration, (3) column equilibration, or (4) both. They then write four branches in their subsequent code to handle each case efficiently.

In terms of determining whether equilibration is necessary, measures of the row and column condition can be obtained from  $\min(r)/\max(r)$  and  $\min(c)/\max(c)$ , where  $r$  and  $c$  are the scaling factors (reciprocals of the row and column maximums). If those measures are near 1 (LAPACK uses  $\geq .1$ ), then equilibration can be skipped.

There is also the issue of the overall scale of the matrix. In theory, the overall scale should not matter, but many packages set tolerances in absolute rather than relative terms, and so overall scale does matter. In most of Mata's other functions, relative scales are used, but provisions are made so that you can specify tolerances in relative or absolute terms. In any case, LAPACK uses the rule that equilibration is necessary if the matrix is too small (its maximum value is less than `epsilon(100)`, approximately  $2.22045\text{e-}14$ ) or too large (greater than  $1/\text{epsilon}(100)$ , approximately  $4.504\text{e+}13$ ).

To summarize,

1. In statistical applications, we believe that equilibration burns too much computer time and adds too much code complexity for the extra accuracy it delivers. This is a judgment call, and we would probably recommend the use of equilibration were it computationally free.
2. If you are going to use equilibration, there is nothing numerically wrong with simply equilibrating matrices in all cases, including those in which equilibration is not necessary. The advantages of this is that you will gain the precision to be had from equilibration while still keeping your code reasonably simple.
3. If you wish to minimize execution time, then you want to perform the minimum amount of equilibration possible and write code to deal efficiently with each case: (1) no equilibration, (2) row equilibration, (3) column equilibration, and (4) row and column equilibration. The defaults used by LAPACK and incorporated in Mata's `_perhapsequil*()` routines are
  - a. Perform row equilibration if  $\min(r)/\max(r) < .1$ , or if  $\min(\text{abs}(A)) < \text{epsilon}(100)$ , or if  $\min(\text{abs}(A)) > 1/\text{epsilon}(100)$ .
  - b. After performing row equilibration, perform column equilibration if  $\min(c)/\max(c) < .1$ , where  $c$  is calculated on  $r*A$ , the row-equilibrated  $A$ , if row equilibration was performed.

## The `_equil*()` family of functions

The `_equil*()` family of functions performs equilibration as follows:

`_equilrc(A, r, c)` performs row equilibration followed by column equilibration; it returns in  $r$  and in  $c$  the row- and column-scaling factors, and it modifies  $A$  to be the fully equilibrated matrix  $r:A:c$ .

`_equilr(A, r)` performs row equilibration only; it returns in  $r$  the row-scaling factors, and it modifies  $A$  to be the row-equilibrated matrix  $r:A$ .

`_equilc(A, c)` performs column equilibration only; it returns in  $c$  the row-scaling factors, and it modifies  $A$  to be the row-equilibrated matrix  $A:c$ .

Here is code to solve  $Ax = b$  using the fully equilibrated form, which damages  $A$  in the process:

```
_equilrc(A, r, c)
x = c:*lusolve(A, r:*b)
```

## The `_perhapsequil*()` family of functions

The `_perhapsequil*()` family of functions mirrors `_equil*()`, except that these functions apply the rules mentioned above for whether equilibration is necessary.

Here is code to solve  $Ax = b$ , which may damage  $A$  in the process:

```
result = _perhapsequilrc(A, r, c)
if (result==0)      x = lusolve(A, b)
else if (result==1) x = lusolve(A, r:*b)
else if (result==2) x = c:*lusolve(A, b)
else if (result==3) x = c:*lusolve(A, r:*b)
```



As a matter of fact, the `_perhapsequilrc()` family returns a vector of 1s when equilibration is not performed, so you could code

```
(void) _perhapsequilrc(A, r, c)
x = c:*lusolve(A, r:*b)
```

but that would be inefficient.

## `rowscalefactors()` and `colscalefactors()`

`rowscalefactors(A)` and `colscalefactors(A)` return the scale factors (reciprocals of row and column maximums) to perform row and column equilibration. These functions are used by the other functions above and are provided for those who wish to write their own equilibration routines.

## Conformability

`_equilrc(A, r, c):`

*input:*

$A: \quad m \times n$

*output:*

$A: \quad m \times n$

$r: \quad m \times 1$

$c: \quad 1 \times n$

`_equilr(A, r):`

*input:*

$A: \quad m \times n$

*output:*

$A: \quad m \times n$

$r: \quad m \times 1$

`_equilc(A, c):`

*input:*

$A: \quad m \times n$

*output:*

$A: \quad m \times n$

$c: \quad 1 \times n$

`_perhapsequilrc(A, r, c):`

*input:*

$A: \quad m \times n$

*output:*

$A: \quad m \times n$  (unmodified if *result* = 0)

$r: \quad m \times 1$

$c: \quad 1 \times n$

*result:*  $1 \times 1$

```
_perhapsequilr(A, r):  
    input:  
        A:       $m \times n$   
  
    output:  
        A:       $m \times n$  (unmodified if result = 0)  
        r:       $m \times 1$   
        result:   $1 \times 1$   
  
_perhapsequilc(A, c):  
    input:  
        A:       $m \times n$   
  
    output:  
        A:       $m \times n$  (unmodified if result = 0)  
        c:       $1 \times n$   
        result:   $1 \times 1$   
  
rowscalefactors(A):  
        A:       $m \times n$   
        result:   $m \times 1$   
  
colscalefactors(A):  
        A:       $m \times n$   
        result:   $1 \times n$ 
```

## Diagnostics

Scale factors used and returned by all functions are calculated by `rowscalefactors(A)` and `colscalefactors(A)`. The functions are defined as `1:/rowmaxabs(A)` and `1:/colmaxabs(A)`, with missing values changed to 1. Thus rows or columns that contain missing or are entirely zero are defined to have scale factors of 1.

Equilibration functions do not equilibrate rows or columns that contain missing or all zeros.

The `_equil*()` functions convert *A* to an array if *A* was a view. The Stata dataset is not changed.

The `_perhapsequil*()` functions convert *A* to an array if *A* was a view and returned is nonzero. The Stata dataset is not changed.

## Also see

[\[M-4\] `matrix`](#) — Matrix functions

## Syntax

```

real scalar    error(real scalar rc)

void           _error(real scalar errnum)
void           _error(string scalar errtxt)
void           _error(real scalar errnum, string scalar errtxt)
    
```

## Description

`error(rc)` displays the standard Stata error message associated with return code *rc* and returns *rc*; see [\[P\] \*\*error\*\*](#) for a listing of return codes. `error()` does not abort execution; standard usage is `exit(error(rc))`.

`_error()` aborts execution and produces a traceback log.

`_error(errnum)` produces a traceback log with standard Mata error message *errnum*; see [\[M-2\] \*\*errors\*\*](#) for a listing of the standard Mata error codes.

`_error(errtxt)` produces a traceback log with error number 3498 and custom text *errtxt*.

`_error(errnum, errtxt)` produces a traceback log with error number *errnum* and custom text *errtxt*.

If *errtxt* is specified, it should contain straight text; SMCL codes are not interpreted.

## Remarks

Remarks are presented under the following headings:

[Use of `\_error\(\)`](#)  
[Use of `error\(\)`](#)

### Use of `_error()`

`_error()` aborts execution and produces a traceback log:

```

: myfunction(A,B)
      mysub(): 3200 conformability error
myfunction(): - function returned error
      <istmt>: - function returned error
r(3200);
    
```

The above output was created because function `mysub()` contained the line

```
_error(3200)
```

and 3200 is the error number associated with the standard message “conformability error”; see [M-2] **errors**. Possibly, the code read

```
if (rows(A)!=rows(B) | cols(A)!=cols(B)) {
    _error(3200)
}
```

Another kind of mistake might produce

```
: myfunction(A,B)
      mysub(): 3498 zeros on diagonal not allowed
myfunction(): - function returned error
<istmt>:      - function returned error
r(3498);
```

and that could be produced by the code

```
if (diag0cnt(A)>0) {
    _error("zeros on diagonal not allowed")
}
```

If we wanted to produce the same text but change the error number to 3300, we could have coded

```
if (diag0cnt(A)>0) {
    _error(3300, "zeros on diagonal not allowed")
}
```

Coding `_error()` is not always necessary. In our conformability-error example, imagine that more of the code read

```
...
if (rows(A)!=rows(B) | cols(A)!=cols(B)) {
    _error(3200)
}
C = A + B
...
```

If we simplified the code to read

```
...
C = A + B
...
```

the conformability error would still be detected because `+` requires p-conformability:

```
: myfunction(A,B)
      +: 3200 conformability error
      mysub(): - conformability error
myfunction(): - function returned error
<istmt>:      - function returned error
r(3200);
```

Sometimes, however, you must detect the error yourself. For instance,

```

...
if (rows(A)!=rows(B) | cols(A)!=cols(B) | rows(A)!=2*cols(A)) {
    _error(3200)
}
C = A + B
...

```

We assume we have some good reason to require that A has twice as many rows as columns. +, however, will not require that, and perhaps no other calculation we will make will require that, either. Or perhaps it will be subsequently detected, but in a way that leads to a confusing error message for the caller.

## Use of error()

error(*rc*) does not cause the program to terminate. Standard usage is

```
exit(error(rc))
```

such as

```
exit(error(503))
```

In any case, error() does not produce a traceback log:

```

: myfunction(A,B)
conformability error
r(503);

```

error() is intended to be used in functions that are subroutines of ado-files:

---

```

----- begin example.ado -----

program example
    version 12
    ...
    mata: myfunction("'mat1'", "'mat2'")
    ...
end
version 12
mata:
void myfunction(string scalar matname1, string scalar matname2)
{
    ...
    A = st_matrix(matname1)
    B = st_matrix(matname2)
    if (rows(A)!=rows(B) | cols(A)!=cols(B)) {
        exit(error(503))
    }
    C = A + B
    ...
}
end

```

---

----- end example.ado -----

This way, when the example command is used incorrectly, the user will see

```

. example ...
conformability error
r(503);

```

rather than the traceback log that would have been produced had we omitted the test and `exit(error(503))`:

```
. example ...
      +: 3200 conformability error
myfunction(): - function returned error
      <istmt>: - function returned error
r(3200);
```

## Conformability

`error(rc)`:

```
rc:      1 × 1
result:  1 × 1
```

`_error(errnum)`:

```
errnum:  1 × 1
result:  void
```

`_error(errtxt)`:

```
errtxt:  1 × 1
result:  void
```

`_error(errnum, errtxt)`:

```
errnum:  1 × 1
errtxt:  1 × 1
result:  void
```

## Diagnostics

`error(rc)` does not abort execution; code `exit(error(rc))` if that is your desire; see [M-5] `exit()`.

The code `error(rc)` returns can differ from `rc` if `rc` is not a standard code or if there is a better code associated with it.

`error(rc)` with `rc = 0` produces no output and returns 0.

`_error(errnum)`, `_error(errtxt)`, and `_error(errnum, errtxt)` always abort with error. `_error()` will abort with error because you called it wrong if you specify an `errnum` less than 1 or greater than 2,147,483,647 or if you specify an `errtxt` longer than 100 characters. If you specify an `errnum` that is not a standard code, the text of the error messages will read “Stata returned error”.

## Also see

[M-2] **errors** — Error codes

[M-5] `exit()` — Terminate execution

[M-4] **programming** — Programming functions

# Title

**[M-5] `errprintf()`** — Format output and display as error message

## Syntax

```
void errprintf(string scalar fmt, r1, r2, ... , rN)
```

## Description

`errprintf()` is a convenience tool for displaying an error message.

`errprintf(...)` is equivalent to `printf(...)` except that it executes `displayas("error")` before the `printf()` is executed; see [\[M-5\] `printf\(\)`](#) and [\[M-5\] `displayas\(\)`](#).

## Remarks

You have written a program. At one point in the code, you have variable `fn` that should contain the name of an existing file:

```
if (!fileexists(fn)) {  
    // you wish to display the error message  
    // file ____ not found  
    exit(601)  
}
```

One solution is

```
if (!fileexists(fn)) {  
    displayas("error")  
    printf("file %s not found\n", fn)  
    exit(601)  
}
```

Equivalent is

```
if (!fileexists(fn)) {  
    errprintf("file %s not found\n", fn)  
    exit(601)  
}
```

It is important that you either `displayas("error")` before using `printf()` or that you use `errprintf()`, to ensure that your error message is displayed (is not suppressed by a [quietly](#)) and that it is displayed in red; see [\[M-5\] `displayas\(\)`](#).

## Conformability

`errprintf(fmt, r1, r2, ..., rN)`

*fmt*:        1 × 1

*r*<sub>1</sub>:        1 × 1

*r*<sub>2</sub>:        1 × 1

...

*r*<sub>*N*</sub>:       1 × 1

*result*:     *void*

## Diagnostics

`errprintf()` aborts with error if a *%fmt* is misspecified, if a numeric *%fmt* corresponds to a string result or a string *%fmt* corresponds to a numeric result, or there are too few or too many *%fmts* in *fmt* relative to the number of *results* specified.

## Also see

[M-5] `printf()` — Format output

[M-5] `displayas()` — Set display level

[M-4] `io` — I/O functions



# Title

[M-5] `exit()` — Terminate execution

## Syntax

```
exit(real scalar rc)
```

```
exit()
```

## Description

`exit(rc)` terminates execution and sets the overall return code to *rc*.

`exit()` with no argument specified is equivalent to `exit(0)`.

## Remarks

Do not confuse `exit()` and `return`. `return` stops execution of the current function and returns to the caller, whereupon execution continues. `exit()` terminates execution. For instance, consider

```
function first()
{
    "begin execution"
    second()
    "this message will never be seen"
}

function second()
{
    "hello from second()"
    exit(0)
}
```

The result of running this would be

```
: first
  begin execution
  hello from second()
```

If we changed the `exit(0)` to be `exit(198)` in `second()`, the result would be

```
: first
  begin execution
  hello from second()
r(198);
```

No error message is presented. If you want to present an error message and exit, you should code `exit(error(198))`; see [M-5] `error()`.

## Conformability

`exit(rc)`:  
*rc*:  $1 \times 1$  (optional)

## Diagnostics

`exit(rc)` and `exit()` do not return.

## Also see

[\[M-5\] `error\(\)`](#) — Issue error message

[\[M-4\] `programming`](#) — Programming functions

# Title

[M-5] **exp()** — Exponentiation and logarithms

## Syntax

*numeric matrix*   **exp**(*numeric matrix Z*)

*numeric matrix*   **ln**(*numeric matrix Z*)

*numeric matrix*   **log**(*numeric matrix Z*)

*numeric matrix*   **log10**(*numeric matrix Z*)

## Description

**exp**(*Z*) returns the elementwise exponentiation of *Z*. **exp**() returns real if *Z* is real and complex if *Z* is complex.

**ln**(*Z*) and **log**(*Z*) return the elementwise natural logarithm of *Z*. The functions are synonyms. **ln**() and **log**() return real if *Z* is real and complex if *Z* is complex.

**ln**(*x*), *x* real, returns the natural logarithm of *x* or returns missing (.) if  $x \leq 0$ .

**ln**(*z*), *z* complex, returns the complex natural logarithm of *z*. **Im**(**ln**(.)) is chosen to be in the interval  $[-\pi, \pi]$ .

**log10**(*Z*) returns the elementwise log base 10 of *Z*. **log10**() returns real if *Z* is real and complex if *Z* is complex. **log10**(*Z*) is defined mathematically and operationally as **ln**(*Z*)/**ln**(10).

## Conformability

**exp**(*Z*), **ln**(*Z*), **log**(*Z*), **log10**(*Z*):

<i>Z</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

**exp**(*Z*) returns missing when  $\text{Re}(Z) > 709$ .

**ln**(*Z*), **log**(*Z*), and **log10**(*Z*) return missing when *Z* is real and  $Z \leq 0$ . In addition, the functions return missing (.) for real arguments when the result would be complex. For instance, **ln**(-1) = ., whereas **ln**(-1+0i) = 3.14159265i.

## Also see

[M-4] **scalar** — Scalar mathematical functions

## Syntax

<i>real matrix</i>	<code>factorial(<i>real matrix R</i>)</code>
<i>real matrix</i>	<code>lnfactorial(<i>real matrix R</i>)</code>
<i>numeric matrix</i>	<code>lngamma(<i>numeric matrix Z</i>)</code>
<i>numeric matrix</i>	<code>gamma(<i>numeric matrix Z</i>)</code>
<i>real matrix</i>	<code>digamma(<i>real matrix R</i>)</code>
<i>real matrix</i>	<code>trigamma(<i>real matrix R</i>)</code>

## Description

`factorial(R)` returns the elementwise factorial of *R*.

`lnfactorial(R)` returns the elementwise  $\ln(\text{factorial}(R))$ , calculated differently. Very large values of *R* may be evaluated.

`lngamma(Z)`, for *Z* real, returns the elementwise real result  $\ln(\text{abs}(\text{gamma}(Z)))$ , but calculated differently. `lngamma(Z)`, for *Z* complex, returns the elementwise  $\ln(\text{gamma}(Z))$ , calculated differently. Thus,  $\text{lngamma}(-2.5) = -.056244$ , whereas  $\text{lngamma}(-2.5+0i) = -.056244 + 3.1416i$ . In both cases, very large values of *Z* may be evaluated.

`gamma(Z)` returns  $\exp(\text{lngamma}(Z))$  for complex arguments and  $\text{Re}(\exp(\text{lngamma}(C(Z))))$  for real arguments. Thus `gamma()` can correctly calculate, say, `gamma(-2.5)` even for real arguments.

`digamma(R)` returns the derivative of `lngamma()` for  $R > 0$ , sometimes called the psi function. `digamma()` requires a real argument.

`trigamma(R)` returns the second derivative of `lngamma()` for  $R > 0$ . `trigamma()` requires a real argument.

## Conformability

All functions return a matrix of the same dimension as input, containing element-by-element calculated results.

## Diagnostics

`factorial()` returns missing for noninteger arguments, negative arguments, and arguments  $> 167$ .

`lnfactorial()` returns missing for noninteger arguments, negative arguments, and arguments  $> 1e+305$ .

lngamma() returns missing for 0, negative integer arguments, negative arguments  $\leq -2,147,483,648$ , and arguments  $> 1e+305$ .

gamma() returns missing for real arguments  $> 171$  and for negative integer arguments.

digamma() returns missing for 0 and negative integer arguments and for arguments  $< -10,000,000$ .

trigamma() returns missing for 0 and negative integer arguments and for arguments  $< -10,000,000$ .

## Also see

[M-4] [scalar](#) — Scalar mathematical functions

[M-4] [statistical](#) — Statistical functions

# Title

[M-5] **favorspeed()** — Whether speed or space is to be favored

## Syntax

*real scalar*   favorspeed()

## Description

`favorspeed()` returns 1 if the user has `mata set matafavor speed` and 0 if the user has `mata set matafavor space` or has not set `matafavor` at all; see [\[M-3\] mata set](#).

## Remarks

Sometimes in programming you can choose between writing code that runs faster but consumes more memory or writing code that conserves memory at the cost of execution speed. `favorspeed()` tells you the user's preference:

```
if (favorspeed()) {  
    /* code structured for speed over memory */  
}  
else {  
    /* code structured for memory over speed */  
}
```

## Conformability

`favorspeed()`:  
*result:*      1 × 1

## Diagnostics

None.

## Also see

[\[M-3\] mata set](#) — Set and display Mata system parameters

[\[M-4\] programming](#) — Programming functions

# Title

**[M-5] ferrortext()** — Text and return code of file error code

## Syntax

```
string scalar ferrortext(real scalar ec)  
real scalar   freturncode(real scalar ec)
```

## Description

`ferrortext(ec)` returns the text associated with a file error code returned by, for instance, `_fopen()`, `_fwrite()`, `fstatus()`, or any other file-processing functions that return an error code. See [\[M-5\] fopen\(\)](#).

`freturncode(ec)` returns the Stata return code associated with the file error code.

## Remarks

Most file-processing functions abort with error if something goes wrong. You attempt to read a nonexisting file, or attempt to write a read-only file, and the file functions you use to do that, usually documented in [\[M-5\] fopen\(\)](#), abort with error. Abort with error means not only that the file function you called stops when the error arises but also that the calling function is aborted. The user is presented with a traceback log documenting what went wrong.

Sometimes you will want to write code to handle such errors for itself. For instance, you are writing a subroutine for a command to be used in Stata and, if the file does not exist, you want the subroutine to present an informative error message and exit without a traceback log but with a nonzero return code. Or in another application, if the file does not exist, that is not an error at all; your code will create the file.

Most file-processing functions have a corresponding underscore function that, rather than aborting, returns an error code when things go wrong. `fopen()` opens a file or aborts with error. `_fopen()` opens a file or returns an error code. The error code is sufficient for your program to take the appropriate action. One uses the underscore functions when the calling program will deal with any errors that might arise.

Let's take the example of simply avoiding traceback logs. If you code

```
fh = fopen(filename, "r")
```

and the file does not exist, execution aborts and you see a traceback log. If you code

```
if ((fh = _fopen(filename, "r"))<0) {  
    errprintf("%s\n", ferrortext(fh))  
    exit(freturncode(fh))  
}
```

execution still stops if the file does not exist, but this time, it stops because you coded `exit()`. You still see an error message, but this time, you see the message because you coded `errprintf()`. No traceback log is produced because you did not insert code to produce one. You could have coded `_exit()` if you wanted one.

The file error codes and the messages associated with them are

Negative code	Meaning
0	all is well
−1	end of file ( <i>this code is usually not an error</i> )
−601	file not found
−602	file already exists
−603	file could not be opened
−608	file is read-only
−610	file not Stata format
−612	unexpected end of file
−630	web files not supported in this version of Stata
−631	host not found
−632	web file not allowed in this context
−633	may not write to web file
−639	file transmission error—checksums do not match
−660	proxy host not found
−662	proxy server refused request to send
−663	remote connection to proxy failed
−665	could not set socket nonblocking
−667	wrong version of winsock.dll
−668	could not find valid winsock.dll or astsys0.lib
−669	invalid URL
−670	invalid network port number
−671	unknown network protocol
−672	server refused to send file
−673	authorization required by server
−674	unexpected response from server
−675	server reported error
−676	server refused request to send
−677	remote connection failed—see <code>r(677)</code> for troubleshooting
−678	could not open local network socket
−679	unexpected web error
−680	could not find valid odbcc32.dll
−681	too many open files
−682	could not connect to ODBC data source name
−683	could not fetch variable in ODBC table
−684	could not find valid dlxabi32.dll
−691	I/O error
−699	insufficient disk space



---

−3601	invalid file handle
−3602	invalid filename
−3611	too many open files
−3621	attempt to write read-only file
−3622	attempt to read write-only file
−3623	attempt to seek append-only file
−3698	file seek error

---

File error codes are usually negative, but neither `ferrortext(ec)` nor `freturncode(ec)` cares whether *ec* is of the positive or negative variety.

## Conformability

`ferrortext(ec)`, `freturncode(ec)`:

*ec*:  $1 \times 1$   
*result*:  $1 \times 1$

## Diagnostics

`ferrortext(ec)` and `freturncode(ec)` treat *ec* = −1 (end of file) the same as *ec* = 612 (unexpected end of file). Code −1 usually is not an error; it just denotes end of file. It is assumed that you will not call `ferrortext()` and `freturncode()` in such cases. If you do call the functions here, it is assumed that you mean that the end of file was unexpected.

## Also see

[M-5] `fopen()` — File I/O

[M-4] `io` — I/O functions

## Syntax

```

complex vector  fft(numeric vector h)
numeric vector invfft(numeric vector H)

void            _fft(complex vector h)
void            _invfft(complex vector H)

numeric vector convolve(numeric vector r, numeric vector s)
numeric vector deconvolve(numeric vector r, numeric vector sm)

numeric vector Corr(numeric vector g, numeric vector h, real scalar k)

real vector    ftperiodogram(numeric vector H)

numeric vector ftpad(numeric vector h)
numeric vector ftwrap(numeric vector r, real scalar n)
numeric vector ftunwrap(numeric vector H)
numeric vector ftretime(numeric vector r, numeric vector s)
real vector    ftfreqs(numeric vector H, real scalar delta)

```

## Description

$H = \text{fft}(h)$  and  $h = \text{invfft}(H)$  calculate the Fourier transform and inverse Fourier transform. The length of  $h$  ( $H$ ) must be a power of 2.

`_fft(h)` and `_invfft(H)` do the same thing, but they perform the calculation in place, replacing the contents of  $h$  and  $H$ .

`convolve(r, s)` returns the convolution of the signal  $s$  with the response function  $r$ . `deconvolve(r, sm)` deconvolves the smeared signal  $sm$  with the response function  $r$  and is thus the inverse of `convolve()`.

`Corr(g, h, k)` returns a  $2k + 1$  element vector containing the correlations of  $g$  and  $h$  for lags and leads as large as  $k$ .

`ftperiodogram(H)` returns a real vector containing the one-sided periodogram of  $H$ .

`ftpad(h)` returns  $h$  padded with 0s to have a length that is a power of 2.

`ftwrap(r, n)` converts the symmetrically stored response function  $r$  into wraparound format of length  $n$ ,  $n \geq \text{rows}(r) * \text{cols}(r)$  and  $\text{rows}(r) * \text{cols}(r)$  odd.

`ftunwrap(H)` unwraps frequency-wraparound order such as returned by `fft()`. You may find this useful when graphing or listing results, but it is otherwise unnecessary.

`ftretime(r, s)` retimes the signal *s* to be on the same time scale as `convolve(r, s)`. This is useful in graphing data and listing results but is otherwise not required.

`ftfreqs(H, delta)` returns a vector containing the frequencies associated with the elements of *H*; *delta* is the sampling interval and is often specified as 1.

## Remarks

Remarks are presented under the following headings:

*Definitions, notation, and conventions*  
*Fourier transform*  
*Convolution and deconvolution*  
*Correlation*  
*Utility routines*  
*Warnings*

## Definitions, notation, and conventions

A signal *h* is a row or column vector containing real or complex elements. The length of the signal is defined as the number of elements of the vector. It is occasionally necessary to pad a signal to a given length. This is done by forming a new vector equal to the original and with zeros added to the end.

The Fourier transform of a signal *h*, typically denoted by capital letter *H* of *h*, is stored in frequency-wraparound order. That is, if there are *n* elements in *H*:

$H[1]$	frequency 0
$H[2]$	frequency 1
$H[3]$	frequency 2
$\vdots$	
$H[n/2]$	frequency $n/2-1$
$H[n/2 + 1]$	frequency $n/2$ ( $-n/2$ , aliased)
$H[n/2 + 2]$	frequency $-(n/2-1)$
$\vdots$	
$H[n-1]$	frequency $-2$
$H[n]$	frequency $-1$

All routines expect and use this order, but see `ftunwrap()` below.

A response function *r* is a row or column vector containing  $m = 2k + 1$  real or complex elements. *m* is called the duration of the response function. Response functions are generally stored symmetrically, although the response function itself need not be symmetric. The response vector contains

$r[1]$	response at lag $-k$
$r[2]$	response at lag $-k + 1$
$\vdots$	
$r[k]$	response at lag $-1$
$r[k + 1]$	contemporaneous response
$r[k + 2]$	response at lead 1
$r[k + 3]$	response at lead 2
$\vdots$	
$r[2k + 1]$	response at lead $k$

Response functions always have odd lengths. Response vectors are never padded.

You may occasionally find it convenient to store a response vector in “wraparound” order (similar to frequency-wraparound order), although none of the routines here require this. In wraparound order:

<code>wrap[1]</code>	contemporaneous response
<code>wrap[2]</code>	response at lead 1
<code>wrap[3]</code>	response at lead 2
$\vdots$	
<code>wrap[k + 1]</code>	response at lead $k$
<code>wrap[k + 2]</code>	response at lag $-k$
<code>wrap[k + 3]</code>	response at lag $-k + 1$
$\vdots$	
<code>wrap[2k + 1]</code>	response at lag $-1$

Response vectors stored in wraparound order may be internally padded (as opposed to merely padded) to a given length by the insertion of zeros between `wrap[k + 1]` and `wrap[k + 2]`.

## Fourier transform

`fft(h)` returns the discrete Fourier transform of  $h$ .  $h$  may be either real or complex, but its length must be a power of 2, so one typically codes `fft(ftpadd(h))`; see `ftpadd()`, below. The returned result is p-conformable with  $h$ . The calculation is performed by `_fft()`.

`invfft(H)` returns the discrete inverse Fourier transform of  $H$ .  $H$  may be either real or complex, but its length must be a power of 2. The returned result is p-conformable with  $H$ . The calculation is performed by `_invfft()`.

`invfft(H)` may return a real or complex. This should be viewed as a feature, but if you wish to ensure the complex interpretation, code `C(invfft(H))`.

`_fft(h)` is the built-in procedure that performs the fast Fourier transform in place.  $h$  must be complex, and its length must be a power of 2.

`_invfft(H)` is the built-in procedure that performs the inverse fast Fourier transform in place.  $H$  must be complex, and its length must be a power of 2.

## Convolution and deconvolution

`convolve(r, s)` returns the convolution of the signal  $s$  with the response function  $r$ . Calculation is performed by taking the `fft()` of the elements, multiplying, and using `invfft()` to transform the results back. Nevertheless, it is not necessary that the length of  $s$  be a power of 2. `convolve()` handles all paddings necessary, including paddings to  $s$  required to prevent the result from being contaminated by erroneous wrapping around of  $s$ . Although one thinks of the convolution operator as being commutative, `convolve()` is not commutative since required zero-padding of the response and signal differ.

If  $n$  is the length of the signal and  $2k + 1$  is the length of the response function, the returned result has length  $n + 2k$ . The first  $k$  elements are the convoluted signal before the true signal begins, and the last  $k$  elements are the convoluted signal after the true signal ends. See `ftretime()`, below. In any case, you may be interested only in the elements `convolve()[k+1:n-k]`, the part contemporaneous with  $s$ .

The returned vector is a row vector if  $s$  is a row vector and a column vector otherwise. The result is guaranteed to be real if both  $r$  and  $s$  are real; the result may be complex or real, otherwise.

It is not required that the response function be shorter than the signal, although this will typically be the case.

`deconvolve(r, sm)` deconvolves the smeared signal  $sm$  with the response function  $r$  and is thus the inverse of `convolve()`. In particular,

$$\text{deconvolve}(r, \text{convolve}(r, s)) = s \quad (\text{up to roundoff error})$$

Everything said about `convolve()` applies equally to `deconvolve()`.

## Correlation

Here we refer to correlation in the signal-processing sense, not the statistical sense.

`Corr(g, h, k)` returns a  $2k + 1$  element vector containing the correlations of  $g$  and  $h$  for lags and leads as large as  $k$ . For instance, `Corr(g, h, 2)` returns a five-element vector, the first element of which contains the correlation for lag 2, the second element lag 1, the third (middle) element the contemporaneous correlation, the fourth element lead 1, and the fifth element lead 2.  $k$  must be greater than or equal to 1. The returned vector is a row or column vector depending on whether  $g$  is a row or column vector.  $g$  and  $h$  must have the same number of elements but need not be p-conformable.

The result is obtained by padding with zeros to avoid contamination, taking the Fourier transform, multiplying  $G \times \text{conj}(H)$ , and rearranging the inverse transformed result. Nevertheless, it is not required that the number of elements of  $g$  and  $h$  be powers of 2 because the program pads internally.

## Utility routines

`ftpad(h)` returns  $h$  padded with 0s to have a length that is a power of 2. For instance,

```
: h = (1,2,3,4,5)
: ftpad(h)
```

	1	2	3	4	5	6	7	8
1	1	2	3	4	5	0	0	0

If  $h$  is a row vector, a row vector is returned. If  $h$  is a column vector, a column vector is returned.

`ftwrap(r, n)` converts the symmetrically stored response function  $r$  into wraparound format of length  $n$ ,  $n \geq \text{rows}(r) \times \text{cols}(r)$  and  $\text{rows}(r) \times \text{cols}(r)$  odd. A symmetrically stored response function is a vector of odd length, for example:

(.1, .5, 1, .2, .4)

The middle element of the vector represents the response function at lag 0. Elements before the middle represent lags while elements after the middle represent leads. Here .1 is the response for lag 2 and .5 for lag 1, 1 the contemporaneous response, .2 the response for lead 1, and .4 the response for lead 2. The wraparound format of a response function records the response at times 0, 1, and so on in the first positions, has some number of zeros, and then records the most negative time value of the response function, and so on.

For instance,

```
: r
      1      2      3      4      5
1  [ .1  .5  1  .2  .4 ]

: ftwrap(r, 5)
      1      2      3      4      5
1  [ 1  .2  .4  .1  .5 ]

: ftwrap(r, 6)
      1      2      3      4      5      6
1  [ 1  .2  .4  0  .1  .5 ]

: ftwrap(r, 8)
      1      2      3      4      5      6      7      8
1  [ 1  .2  .4  0  0  0  .1  .5 ]
```

`ftunwrap(H)` unwraps frequency-wraparound order such as returned by `fft()`. You may find this useful when graphing or listing results, but it is otherwise unnecessary. Frequency-unwrapped order is defined as

```
unwrap[1]      frequency  $-(n/2) + 1$ 
unwrap[2]      frequency  $-(n/2) + 2$ 
:
unwrap[n/2 - 1] frequency  $-1$ 
unwrap[n/2]     frequency  $0$ 
unwrap[n/2 + 1] frequency  $1$ 
:
unwrap[n - 1]   frequency  $n/2 - 1$ 
unwrap[n]       frequency  $n/2$ 
```

Here we assume that  $n$  is even, as will usually be true. The aliased (highest) frequency is assigned the positive sign.

Also see `ftperiodogram()`, below.

`ftretime(r, s)` retimes the signal  $s$  to be on the same time scale as `convolve(r, s)`. This is useful in graphing and listing results but is otherwise not required. `ftretime()` uses only the length of  $r$ , and not its contents, to perform the retiming. If the response vector is of length  $2k + 1$ , a vector containing  $k$  zeros,  $s$ , and  $k$  more zeros is returned. Thus the result of `ftretime(r, s)` is p-conformable with `convolve(r, s)`.

`ftfreqs(H, delta)` returns a p-conformable-with- $H$  vector containing the frequencies associated with the elements of  $H$ .  $delta$  is the sampling interval and is often specified as 1.

`ftperiodogram(H)` returns a real vector of length  $n/2$  containing the one-sided periodogram of  $H$  (length  $n$ ), calculated as

$$|H(f)|^2 + |H(-f)|^2$$

excluding frequency 0. Thus `ftperiodogram(H)[1]` corresponds to frequency 1 ( $-1$ ), `ftperiodogram(H)[2]` to frequency 2 ( $-2$ ), and so on.

## Warnings

`invfft(H)` will cast the result down to real if possible. Code `C(invfft(H))` if you want to be assured of the result being stored as complex.

`convolve(r, s)` is not the same as `convolve(s, r)`.

`convolve(r, s)` will cast the result down to real if possible. Code `C(convolve(r, s))` if you want to be assured of the result being stored as complex.

For `convolve(r, s)`, the response function  $r$  must have odd length.

## Conformability

`fft(h)`:

$h$ :	$1 \times n$	or	$n \times 1$ , $n$ a power of 2
result:	$1 \times n$	or	$n \times 1$

`invfft(H)`:

$H$ :	$1 \times n$	or	$n \times 1$ , $n$ a power of 2
result:	$1 \times n$	or	$n \times 1$

`_fft(h)`:

$h$ :	$1 \times n$	or	$n \times 1$ , $n$ a power of 2
result:	void		

`_invfft(H)`:

$H$ :	$1 \times n$	or	$n \times 1$ , $n$ a power of 2
result:	void		

`convolve(r, s)`:

$r$ :	$1 \times n$	or	$n \times 1$ , $n > 0$ , $n$ odd
$s$ :	$1 \times 2k + 1$	or	$2k + 1 \times 1$ , i.e., $s$ of odd length
result:	$1 \times 2k + n$	or	$2k + n \times 1$

`deconvolve(r, sm):`

<i>r</i> :	$1 \times n$	or	$n \times 1, \quad n > 0, n \text{ odd}$
<i>sm</i> :	$1 \times 2k + n$	or	$2k + n \times 1$
<i>result</i> :	$1 \times 2k + 1$	or	$2k + 1 \times 1$

`Corr(g, h, k):`

<i>g</i> :	$1 \times n$	or	$n \times 1, \quad n > 0$
<i>h</i> :	$1 \times n$	or	$n \times 1$
<i>k</i> :	$1 \times 1$	or	$1 \times 1, \quad k > 0$
<i>result</i> :	$1 \times 2k + 1$	or	$2k + 1 \times 1$

`ftperiodogram(H):`

<i>H</i> :	$1 \times n$	or	$n \times 1, \quad n \text{ even}$
<i>result</i> :	$n/2 \times 1$	or	$1 \times n/2$

`ftpad(h):`

<i>h</i> :	$1 \times n$	or	$n \times 1$
<i>result</i> :	$1 \times N$	or	$N \times 1, \quad N = n \text{ rounded up to power of } 2$

`ftwrap(r, n):`

<i>r</i> :	$1 \times m$	or	$m \times 1, \quad m > 0, m \text{ odd}$
<i>n</i> :	$1 \times 1$	or	$1 \times 1, \quad n \geq m$
<i>result</i> :	$1 \times n$	or	$n \times 1$

`ftunwrap(H):`

<i>H</i> :	$1 \times n$	or	$n \times 1$
<i>result</i> :	$1 \times n$	or	$n \times 1$

`ftretime(r, s):`

<i>r</i> :	$1 \times n$	or	$n \times 1, \quad n > 0, n \text{ odd}$
<i>s</i> :	$1 \times 2k + 1$	or	$2k + 1 \times 1, \quad \text{i.e., } s \text{ of odd length}$
<i>result</i> :	$1 \times 2k + n$	or	$2k + n \times 1$

`ftfreqs(H, delta):`

<i>H</i> :	$1 \times n$	or	$n \times 1, \quad n \text{ even}$
<i>delta</i> :	$1 \times 1$		
<i>result</i> :	$1 \times n$	or	$n \times 1$

## Diagnostics

All functions abort with error if the conformability requirements are not met. This is always true, of course, but pay particular attention to the requirements outlined under [Conformability](#) directly above.

`fft(h)`, `_fft(h)`, `invfft(H)`, `_invfft(H)`, `convolve(r, s)`, `deconvolve(r, sm)`, and `Corr(g, h, k)` return missing results if any argument contains missing values.

`ftwrap(r, n)` aborts with error if *n* contains missing value.

## Also see

[M-4] [mathematical](#) — Important mathematical functions



# Title

[M-5] fileexists() — Whether file exists

## Syntax

*real scalar*    fileexists(*string scalar fn*)

## Description

fileexists(*fn*) returns 1 if file *fn* exists and is readable and returns 0 otherwise.

## Conformability

fileexists(*fn*):  
          *fn*:        1 × 1  
          *result*:    1 × 1

## Diagnostics

None.

## Also see

[\[M-4\] io](#) — I/O functions

# Title

[M-5] `_fillmissing()` — Fill matrix with missing values

## Syntax

```
void _fillmissing(transmorphic matrix A)
```

## Description

`_fillmissing(transmorphic matrix A)` changes the contents of *A* to missing values.

## Remarks

The definition of missing depends on the storage type of *A*:

Storage type	Contents
real	.
complex	C(.)
string	" "
pointer	NULL

## Conformability

`_fillmissing(A)`:

*input:*

*A:*      $r \times c$

*output:*

*A:*      $r \times c$

## Diagnostics

None.

## Also see

[M-4] [manipulation](#) — Matrix manipulation

## Syntax

*pointer()*   *scalar*   **findexternal**(*string scalar name*)

*pointer()*   *scalar*   **crexternal**(*string scalar name*)

*void*                      **rmexternal**(*string scalar name*)

*string*            *scalar*   **nameexternal**(*pointer()* *scalar p*)

## Description

**findexternal**(*name*) returns a pointer (see [M-2] [pointers](#)) to the external global matrix, vector, or scalar whose name is specified by *name*, or to the external global function if the contents of *name* end in (). **findexternal**() returns NULL if the external global is not found.

**crexternal**(*name*) creates a new external global  $0 \times 0$  real matrix with the specified name and returns a pointer to it; it returns NULL if an external global of that name already exists.

**rmexternal**(*name*) removes (deletes) the specified external global or does nothing if no such external global exists.

**nameexternal**(*p*) returns the name of *\*p*.

## Remarks

Remarks are presented under the following headings:

*Definition of a global*

*Use of globals*

Also see [Linking to external globals](#) in [M-2] [declarations](#).

## Definition of a global

When you use Mata interactively, any variables you create are known, equivalently, as externals, globals, or external globals.

```
: myvar = x
```

Such variables can be used by subsequent functions that you run, and there are two ways that can happen:

```
function example1(...)
{
    external real myvar
    ... myvar ...
}
```

and

```
function example2(...)
{
    pointer(real) p

    p = findexternal("myvar")
    ... *p ...
}
```

Using the first method, you must know the name of the global at the time you write the source code, and when you run your program, if the global does not exist, it will refuse to run (abort with `myvar` not found). With the second method, the name of the global can be specified at run time and what is to happen when the global is not found is up to you.

In the second example, although we declared `p` as a pointer to a real, `myvar` will not be required to contain a real. After `p = findexternal("myvar")`, if `p!=NULL`, `p` will point to whatever `myvar` contains, whether it be real, complex, string, or another pointer. (You can diagnose the contents of `*p` using `eltype(*p)` and `orgtype(*p)`; see [M-5] [eltype\(\)](#).)

## Use of globals

Globals are useful when a function must remember something from one call to the next:

```
function example3(real scalar x)
{
    pointer() scalar p

    if ( (p = findexternal("myprivatevar")) == NULL) {
        printf("you haven't called me previously")
        p = crexternal("myprivatevar")
    }
    else {
        printf("last time, you said \"%g\", *p)
    }
    *p = x
}

: example3(2)
you haven't called me previously
: example3(31)
last time, you said 2
: example3(9)
last time, you said 31
```

Note our use of the name `myprivatevar`. It actually is not a private variable; it is global, and you would see the variable listed if you described the contents of Mata's memory. Because global variables are so exposed, it is best that you give them long and unlikely names.

In general, programs do not need global variables. The exception is when a program must remember something from one invocation to the next, and especially if that something must be remembered from one invocation of Mata to the next.

When you do need globals, you probably will have more than one thing you will need to recall. There are two ways to proceed. One way is simply to create separate global variables for each thing you need to remember. The other way is to create one global pointer vector and store everything in that. In the following example, we remember one scalar and one matrix:

```
function example4()
{
    pointer(pointer() vector) scalar    p
    scalar                                     s
    real matrix                               X
    pointer() scalar                         ps, pX

    if ( (p = findexternal("mycollection")) == NULL) {
        ... calculate scalar s and X from nothing ...
        ... and save them:
        p = crexternal("mycollection")
        *p = (&s, &X)
    }
    else {
        ps = (*p)[1]
        pX = (*p)[2]
        ... calculate using *ps and *pX ...
    }
}
```

In the above example, even though `crexternal()` created a  $0 \times 0$  real global, we morphed it into a  $1 \times 2$  pointer vector:

```
p = crexternal("mycollection")    *p is  $0 \times 0$  real
*p = (&s, &X)                     *p is  $1 \times 2$  vector
```

just as we could with any nonpointer object.

In the else part of our program, where we use the previous values, we do not use variables `s` and `X`, but `ps` and `pX`. Actually, we did not really need them, we could just as well have used `*((*p)[1])` and `*((*p)[2])`, but the code is more easily understood by introducing `*ps` and `*pX`.

Actually, we could have used the variables `s` and `X` by changing the else part of our program to read

```
else {
    s = (*p)[1]
    X = (*p)[2]
    ... calculate using s and X ...
    *p = (&s, &X)          ← remember to put them back
}
```

Doing that is inefficient because `s` and `X` contain copies of the global values. Obviously, the amount of inefficiency depends on the sizes of the elements being copied. For `s`, there is really no inefficiency at all because `s` is just a scalar. For `X`, the amount of inefficiency depends on the dimensions of `X`. Making a copy of a small `X` matrix would introduce just a little inefficiency.

The best balance between efficiency and readability is achieved by introducing a subroutine:

```
function example5()
{
    pointer(pointer() vector) scalar    p
    scalar                                     s
    real matrix                                X

    if ( (p = findexternal("mycollection")) == NULL) {
        example5_sub(1, s=., X=J(0,0,.))
        p = crexternal("mycollection")
        *p = (&s, &X)
    }
    else {
        example5_sub(0, (*p)[1], (*p)[2])
    }
}

function example5_sub(scalar firstcall, scalar x, matrix X)
{
    ...
}
```

The last two lines in the not-found case

```
p = crexternal("mycollection")
*p = (&s, &X)
```

could also be coded

```
*crexternal("mycollection") = (&s, &X)
```

## Conformability

`findexternal(name)`, `crexternal(name)`:

<i>name</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

`rmexternal(name)`:

<i>name</i> :	$1 \times 1$
<i>result</i> :	<i>void</i>

`nameexternal(p)`:

<i>p</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

`findexternal(name)`, `crexternal(name)`, and `rmexternal(name)` abort with error if *name* contains an invalid name.

`findexternal(name)` returns NULL if *name* does not exist.

`crexternal(name)` returns `NULL` if *name* already exists.

`nameexternal(p)` returns `""` if *p* = `NULL`. Also, `nameexternal()` may be used not just with pointers to globals but pointers to locals as well. For example, you can code `nameexternal(&myx)`, where `myx` is declared in the same program or a calling program. `nameexternal()` will usually return the expected local name, such as “`myx`”. In such cases, however, it is also possible that `""` will be returned. That can occur because, in the compilation/optimization process, the identity of local variables can be lost.

## Also see

[M-5] [`valofexternal\(\)`](#) — Obtain value of external global

[M-4] [programming](#) — Programming functions

# Title

[M-5] **findfile()** — Find file

## Syntax

*string scalar*    **findfile**(*string scalar fn*, *string scalar dirlist*)

*string scalar*    **findfile**(*string scalar fn*)

## Description

**findfile**(*fn*, *dirlist*) looks for file *fn* along the semicolon-separated list of directories *dirlist* and returns the fully qualified path and filename if *fn* is found. **findfile**() returns "" if the file is not found.

**findfile**(*fn*) is equivalent to **findfile**(*fn*, c("adopath")). **findfile**() with one argument looks along the official Stata ado-path for file *fn*.

## Remarks

For instance,

```
findfile("kappa.ado")
```

might return C:\Program Files\Stata12\ado\updates\k\kappa.ado.

## Conformability

```
findfile(fn, dirlist):  
   fn:           1 × 1  
   dirlist:       1 × 1  (optional)  
   result:       1 × 1
```

## Diagnostics

**findfile**(*fn*, *dirlist*) and **findfile**(*fn*) return "" if the file is not found. If the file is found, the returned fully qualified path and filename is guaranteed to exist and be readable at the instant **findfile**() concluded.

## Also see

[M-4] **io** — I/O functions



# Title

[M-5] floatround() — Round to float precision

## Syntax

*real matrix* floatround(*real matrix x*)

## Description

floatround(*x*) returns *x* rounded to IEEE 4-byte real (float) precision. floatround() is the element-by-element equivalent of Stata's [float\(\)](#) function. The Mata function could not be named float() because the word float is reserved in Mata.

## Remarks

```
: printf(" %21x\n", .1)
+1.999999999999999aX-004
: printf(" %21x\n", floatround(.1))
+1.99999a0000000X-004
```

## Conformability

```
floatround(x):
      x:       $r \times c$ 
      result:  $r \times c$ 
```

## Diagnostics

floatround(*x*) returns missing (.) if  $x < -1.fffffeX+7e$  (approximately  $-1.70141173319e+38$ ) or  $x > 1.fffffeX+7e$  (approximately  $1.70141173319e+38$ ).

In contrast with most functions, floatround(*x*) returns the same kind of missing value as *x* if *x* contains missing; . if *x* == ., .a if *x* == .a, .b if *x* == .b, ..., and .z if *x* == .z.

## Also see

[\[M-4\] utility](#) — Matrix utility functions

# Title

[M-5] `fmtwidth()` — Width of %fmt

## Syntax

*real matrix* `fmtwidth(string matrix f)`

## Description

`fmtwidth(f)` returns the width of the *%fmt* contained in *f*.

## Remarks

`fmtwidth("%9.2f")` returns 9.

`fmtwidth("%20s")` returns 20.

`fmtwidth("%tc")` returns 18.

`fmtwidth("%tcDay_Mon_DD_hh:mm:ss!C!D!T_CCYY")` returns 28.

`fmtwidth("not a format")` returns . (missing).

## Conformability

<code>fmtwidth(f):</code>	
<i>f</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

`fmtwidth(f)` returns . (missing) when *f* does not contain a valid Stata format.

## Also see

[M-5] `strlen()` — Length of string

[M-4] `string` — String manipulation functions

## Syntax

*real scalar*    `fopen(string scalar fn, mode)`

*real scalar*    `fopen(string scalar fn, mode, public)`

*real scalar*    `_fopen(string scalar fn, mode)`

*real scalar*    `_fopen(string scalar fn, mode, public)`

where

*mode*:    *string scalar* containing "r", "w", "rw", or "a"

*public*:    optional *real scalar* containing zero or nonzero

In what follows, *fh* is the value returned by `fopen()` or `_fopen()`:

*void*            `fclose(fh)`

*real scalar*    `_fclose(fh)`

*string scalar*    `fget(fh)`

*string scalar*    `_fget(fh)`

*string scalar*    `fgetnl(fh)`

*string scalar*    `_fgetnl(fh)`

*string scalar*    `fread(fh, real scalar len)`

*string scalar*    `_fread(fh, real scalar len)`

*void*            `fput(fh, string scalar s)`

*real scalar*    `_fput(fh, string scalar s)`

*void*            `fwrite(fh, string scalar s)`

*real scalar*    `_fwrite(fh, string scalar s)`

*matrix*        `fgetmatrix(fh)`

*matrix*        `_fgetmatrix(fh)`

*void*            `fputmatrix(fh, transmorphic matrix X)`

*real scalar*    `_fputmatrix(fh, transmorphic matrix X)`

*real scalar*    `fstatus(fh)`

*real scalar*    `ftell(fh)`

*real scalar*    `_ftell(fh)`

*void*            `fseek(fh, real scalar offset, real scalar whence)`

*real scalar*    `_fseek(fh, real scalar offset, real scalar whence)`

(*whence* is coded `-1`, `0`, or `1`, meaning from start of file, from current position, or from end of file; *offset* is signed: positive values mean after *whence* and negative values mean before)

*void*            `ftruncate(fh)`

*real scalar*    `_ftruncate(fh)`

## Description

These functions read and write files. First, open the file and get back a file handle (*fh*). The file handle, which is nothing more than an integer, is how you refer to the file in the calls to other file I/O functions. When you are finished, close the file.

Most file I/O functions come in two varieties: without and with an underscore in front of the name, such as `fopen()` and `_fopen()`, and `fwrite()` and `_fwrite()`.

In functions without a leading underscore, errors cause execution to be aborted. For instance, you attempt to open a file for read and the file does not exist. Execution stops. Or, having successfully opened a file, you attempt to write into it and the disk is full. Execution stops. When execution stops, the appropriate error message is presented.

In functions with the leading underscore, execution continues and no error message is displayed; it is your responsibility (1) to verify that things went well and (2) to take the appropriate action if they did not. Concerning (1), some underscore functions return a status value; others require that you call `fstatus()` to obtain the status information.

You can mix and match use of underscore and nonunderscore functions, using, say, `_fopen()` to open a file and `fread()` to read it, or `fopen()` to open and `_fwrite()` to write.

## Remarks

Remarks are presented under the following headings:

- Opening and closing files*
- Reading from a file*
- Writing to a file*
- Reading and writing in the same file*
- Reading and writing matrices*
- Repositioning in a file*
- Truncating a file*
- Error codes*

## Opening and closing files

### Functions

`fopen(string scalar fn, string scalar mode)`

`_fopen(string scalar fn, string scalar mode)`

`fopen(string scalar fn, string scalar mode, real scalar public)`

`_fopen(string scalar fn, string scalar mode, real scalar public)`

open a file. The file may be on a local disk, a network disk, or even on the web (such as <http://www.stata.com/index.html>). *fn* specifies the filename, and *mode* specifies how the file is to be opened:

<i>mode</i>	Meaning
"r"	Open for reading; file must exist and be readable. File may be "http://..." file. File will be positioned at the beginning.
"w"	Open for writing; file must not exist and the directory be writable. File may not be "http://..." file. File will be positioned at the beginning.
"rw"	Open for reading and writing; file must either exist and be writable or not exist and directory be writable. File may not be "http://..." file. File will be positioned at the beginning (new file) or at the end (existing file).
"a"	Open for appending; file must either exist and be writable or not exist and directory be writable. File may not be "http://..." file. File will be positioned at the end.

Other values for *mode* cause `fopen()` and `_fopen()` to abort with an invalid-mode error.

Optional third argument *public* specifies whether the file, if it is being created, should be given permissions so that everyone can read it, or if it instead should be given the normal permissions. Not specifying *public*, or specifying *public* as 0, gives the file the normal permissions. Specifying *public* as nonzero makes the file publicly readable. *public* is relevant only when the file is being created, that is, is being opened "w", or being opened "rw" and not previously existing.

`fopen()` returns a file handle; the file is opened or execution is aborted.

`_fopen()` returns a file handle or returns a negative number. If a negative number is returned, the file is not open, and the number indicates the reason. For `_fopen()`, there are a few likely possibilities

Negative value	Meaning
−601	file not found
−602	file already exists
−603	file could not be opened
−608	file is read-only
−691	I/O error

and there are many other possibilities. For instance, perhaps you attempted to open a file on the web (say, <http://www.newurl.org/upinfo.doc>) and the URL was not found, or the server refused to send back the file, etc. See *Error codes* below for a complete list of codes.

After opening the file, you use the other file I/O commands to read and write it, and then you close the file with `fclose()` or `_fclose()`. `fclose()` returns nothing; if the file cannot be closed, execution is aborted. `_fclose` returns 0 if successful, or a negative number otherwise. For `_fclose()`, the likely possibilities are

Negative value	Meaning
−691	filesystem I/O error

## Reading from a file

You may read from a file opened "r" or "rw". The commands to read are

```
fget(fh)
fgetnl(fh)
fread(fh, real scalar len)
```

and, of course,

```
_fget(fh)
_fgetnl(fh)
_fread(fh, real scalar len)
```

All functions, with or without an underscore, require a file handle be specified, and all the functions return a string scalar or they return `J(0,0,"")`, a  $0 \times 0$  string matrix. They return `J(0,0,"")` on end of file and, for the underscore functions, when the read was not successful for other reasons. When using the underscore functions, you use `fstatus()` to obtain the status code; see *Error codes* below. The underscore read functions are rarely used because the only reason a read can fail is I/O error, and there is not much that can be done about that except abort, which is exactly what the nonunderscore functions do.

`fget(fh)` is for reading ASCII files; the next line from the file is returned, without end-of-line characters. (If the line is longer then 32,768 characters, the first 32,768 characters are returned.)

`fgetnl(fh)` is much the same as `fgetc()`, except that the new-line characters are not removed from the returned result. (If the line is longer than 32,768 characters, the first 32,768 characters are returned.)

`fread(fh, len)` is usually used for reading binary files and returns the next *len* characters (bytes) from the file or, if there are fewer than that remaining to be read, however many remain. (*len* may not exceed 2,147,483,647 on 32-bit computers and 9,007,199,254,740,991 [sic] on 64-bit computers; memory shortages for storing the result will arise long before these limits are reached on most computers.)

The following code reads and displays a file:

```
fh = fopen(filename, "r")
while ((line=fgetc(fh))!=J(0,0,"")) {
    printf("%s\n", line)
}
fclose(fh)
```

## Writing to a file

You may write to a file opened "w", "rw", or "a". The functions are

```
fputc(fh, string scalar s)
fwrite(fh, string scalar s)
```

and, of course,

```
_fputc(fh, string scalar s)
_ffwrite(fh, string scalar s)
```

*fh* specifies the file handle, and *s* specifies the string to be written. `fputc()` writes *s* followed by the new-line characters appropriate for your operating system. `fwrite()` writes *s* alone.

`fputc()` and `fwrite()` return nothing; `_fputc()` and `_ffwrite()` return a real scalar equal to 0 if all went well or a negative error code; see [Error codes](#) below.

The following code copies text from one file to another:

```
fh_in = fopen(inputname, "r")
fh_out = fopen(outputname, "w")
while ((line=fgetc(fh_in))!=J(0,0,"")) {
    fputc(fh_out, line)
}
fclose(fh_out)
fclose(fh_in)
```

The following code reads a file (binary or text) and changes every occurrence of "a" to "b":

```
fh_in = fopen(inputname, "r")
fh_out = fopen(outputname, "w")
while ((c=fread(fh_in, 1))!=J(0,0,"")) {
    fwrite(fh_out, (c=="a" ? "b" : c))
}
fclose(fh_out)
fclose(fh_in)
```

## Reading and writing in the same file

You may read and write from a file opened "rw", using any of the read or write functions described above. When reading and writing in the same file, one often uses file repositioning functions, too; see [Repositioning in a file](#) below.

## Reading and writing matrices

Functions

```
fputmatrix(fh, transmorphic matrix X)
```

and

```
_fputmatrix(fh, transmorphic matrix X)
```

will write a matrix to a file. In the usual fashion, `fputmatrix()` returns nothing (it aborts if there is an I/O error) and `_fputmatrix()` returns a scalar equal to 0 if all went well and a negative error code otherwise.

Functions

```
fgetmatrix(fh)
```

and

```
_fgetmatrix(fh)
```

will read a matrix written by `fputmatrix()` or `_fputmatrix()`. Both functions return the matrix read or return `J(0,0,.)` on end of file (both functions) or error (`_fgetmatrix()` only). Because `J(0,0,.)` could be the matrix that was written, distinguishing between that and end of file requires subsequent use of `fstatus()`. `fstatus()` will return 0 if `fgetmatrix()` or `_fgetmatrix()` returned a written matrix, -1 if end of file, or (after `_fgetmatrix()`) a negative error code.

`fputmatrix()` writes matrices in a compact, efficient, and portable format; a matrix written on a Windows computer can be read back on a Mac or Unix computer and vice versa.

The following code creates a file containing three matrices

```
fh = fopen(filename, "w")
fputmatrix(fh, a)
fputmatrix(fh, b)
fputmatrix(fh, c)
fclose(fh)
```



and the following code reads them back:

```
fh = fopen(filename, "r")
a = fgetmatrix(fh)
b = fgetmatrix(fh)
c = fgetmatrix(fh)
fclose(fh)
```

## □ Technical note

You may even write pointer matrices

```
mats = (&a, &b, &c, NULL)
fh = fopen(filename, "w")
fputmatrix(fh, mats)
fclose(fh)
```

and read them back:

```
fh = fopen(filename, "r")
mats = fgetmatrix(fh)
fclose(fh)
```

When writing pointer matrices, `fputmatrix()` writes `NULL` for any pointer-to-function value. It is also recommended that you do not write self-referential matrices (matrices that point to themselves, either directly or indirectly), although the elements of the matrix may be cross linked and even recursive themselves. If you are writing pointer matrix `p`, no element of `p`, `*p`, `**p`, etc., should contain `&p`. That one address cannot be preserved because in the assignment associated with reading back the matrix (the "*result*"= part of *result*=`fgetmatrix(fh)`), a new matrix with a different address is associated with the contents.

□

## Repositioning in a file

The function

```
ftell(fh)
```

returns a real scalar reporting where you are in a file, and function

```
fseek(fh, real scalar offset, real scalar whence)
```

changes where you are in the file to be *offset* bytes from the beginning of the file (*whence* = -1), *offset* bytes from the current position (*whence* = 0), or *offset* bytes from the end of the file (*whence* = 1).

Functions `_ftell()` and `_fseek()` do the same thing as `ftell()` and `fseek()`, the difference being that, rather than aborting on error, the underscore functions return negative error codes. `_ftell()` is pretty well useless as the only error that can arise is I/O error, and what else are you going to do other than abort? `_fseek()`, however, has a use, because it allows you to try out a repositioning and check whether it was successful. With `fseek()`, if the repositioning is not successful, execution is aborted.

Say you open a file for read:

```
fh = fopen(filename, "r")
```

After opening the file in mode `r`, you are positioned at the beginning of the file or, in the jargon of file processing, at position 0. Now say that you read 10 bytes from the file:

```
part1 = fread(fh, 10)
```

Assuming that was successful, you are now at position 10 of the file. Say that you next read a line from the file

```
line = fget(fh)
```

and assume that `fget()` returns "abc". You are now at position 14 or 15. (No, not 13: `fget()` read the line and the new-line characters and returned the line. abc was followed by carriage return and line feed (two characters) if the file was written under Windows and by a carriage return or line feed alone (one character) if the file was written under Mac or Unix).

`ftell(fh)` and `_ftell(fh)` tell you where you are in the file. Coding

```
pos = ftell(fh)
```

would store 14 or 15 in `pos`. Later in your code, after reading more of the file, you could return to this position by coding

```
fseek(fh, pos, -1)
```

You could return to the beginning of the file by coding

```
fseek(fh, 0, -1)
```

and you could move to the end of the file by coding

```
fseek(fh, 0, 1)
```

`ftell(fh)` is equivalent to `_fseek(fh, 0, 0)`.

Repositioning functions cannot be used when the file has been opened "a".

## Truncating a file

Truncation refers to making a longer file shorter. If a file was opened "w" or "rw", you may truncate it at its current position by using

```
ftruncate(fh)
```

or

```
_ftruncate(fh)
```

`ftruncate()` returns nothing; `_ftruncate()` returns 0 on success and otherwise returns a negative error code.

The following code shortens a file to its first 100 bytes:

```
fh = fopen(filename, "rw")
fseek(fh, 100, -1)
ftruncate(fh)
fclose(fh)
```

## Error codes

If you use the underscore I/O functions, if there is an error, they will return a negative code. Those codes are

Negative code	Meaning
0	all is well
−1	end of file
−2	connection timed out
−601	file not found
−602	file already exists
−603	file could not be opened
−608	file is read-only
−610	file format error
−612	unexpected end of file
−630	web files not supported in this version of Stata
−631	host not found
−632	web file not allowed in this context
−633	web file not allowed in this context
−660	proxy host not found
−661	host or file not found
−662	proxy server refused request to send
−663	remote connection to proxy failed
−665	could not set socket nonblocking
−669	invalid URL
−670	invalid network port number
−671	unknown network protocol
−672	server refused to send file
−673	authorization required by server
−674	unexpected response from server
−675	server reported server error
−676	server refused request to send
−677	remote connection failed
−678	could not open local network socket
−679	unexpected web error
−691	I/O error
−699	insufficient disk space
−3601	invalid file handle
−3602	invalid filename
−3603	invalid file mode
−3611	too many open files
−3621	attempt to write read-only file
−3622	attempt to read write-only file
−3623	attempt to seek append-only file
−3698	file seek error

---

Other codes in the −600 to −699 range are possible. The codes in this range correspond to the negative of the corresponding Stata return code; see [\[P\] error](#).

Underscore functions that return a real scalar will return one of these codes if there is an error.

If an underscore function does not return a real scalar, then you obtain the outcome status using `fstatus()`. For instance, the read-string functions return a string scalar or `J(0,0,"")` on end of file. The underscore variants do the same, and they also return `J(0,0,"")` on error, meaning error looks like end of file. You can determine the error code using the function

```
fstatus(fh)
```

`fstatus()` returns 0 (no previous error) or one of the negative codes above.

`fstatus()` may be used after any underscore I/O command to obtain the current error status.

`fstatus()` may also be used after the nonunderscore I/O commands; then `fstatus()` will return `-1` or 0 because all other problems would have stopped execution.

## Conformability

`fopen(fn, mode, public)`, `_fopen(fn, mode, public)`:

```
fn:      1 × 1
mode:    1 × 1
public:  1 × 1   (optional)
result:  1 × 1
```

`fclose(fh)`:

```
fh:      1 × 1
result:  void
```

`_fclose(fh)`:

```
fh:      1 × 1
result:  1 × 1
```

`fget(fh)`, `_fget(fh)`, `fgetnl(fh)`, `_fgetnl(fh)`:

```
fh:      1 × 1
result:  1 × 1   or   0 × 0 if end of file
```

`fread(fh, len)`, `_fread(fh, len)`:

```
fh:      1 × 1
len:     1 × 1
result:  1 × 1   or   0 × 0 if end of file
```

`fput(fh, s)`, `fwrite(fh, s)`:

```
fh:      1 × 1
s:       1 × 1
result:  void
```

`_fput(fh, s)`, `_fwrite(fh, s)`:

```
fh:      1 × 1
s:       1 × 1
result:  1 × 1
```

`fgetmatrix(fh)`, `_fgetmatrix(fh)`:

```
fh:      1 × 1
result:  r × c   or   0 × 0 if end of file
```

```

fputmatrix(fh, X):
    fh:      1 × 1
    X:      r × c
    result:  void

_fputmatrix(fh, X):
    fh:      1 × 1
    X:      r × c
    result:  1 × 1

fstatus(fh):
    fh:      1 × 1
    result:  1 × 1

ftell(fh), _ftell(fh):
    fh:      1 × 1
    result:  1 × 1

fseek(fh, offset, whence):
    fh:      1 × 1
    offset:  1 × 1
    whence:  1 × 1
    result:  void

_fseek(fh, offset, whence):
    fh:      1 × 1
    offset:  1 × 1
    whence:  1 × 1
    result:  1 × 1

ftruncate(fh):
    fh:      1 × 1
    result:  void

_ftruncate(fh):
    fh:      1 × 1
    result:  1 × 1

```

## Diagnostics

`fopen(fn, mode)` aborts with error if *mode* is invalid or if *fn* cannot be opened or if an attempt is made to open too many files simultaneously.

`_fopen(fn, mode)` aborts with error if *mode* is invalid or if an attempt is made to open too many files simultaneously. `_fopen()` returns the appropriate negative error code if *fn* cannot be opened.

All remaining I/O functions—even functions with leading underscore—abort with error if *fh* is not a handle to a currently open file.

Also, the functions that do not begin with an underscore abort with error when a file was opened read-only and a request requiring write access is made, when a file is opened write-only and a request requiring read access is made, etc. See [Error codes](#) above; all problems except code `-1` (end of file) cause the nonunderscore functions to abort with error.

Finally, the following functions will also abort with error for the following specific reasons:

`fseek(fh, offset, whence)` and `_fseek(fh, offset, whence)` abort with error if *offset* is outside the range  $\pm 2,147,483,647$  on 32-bit computers; if *offset* is outside the range  $\pm 9,007,199,254,740,991$  on 64-bit computers; or, on all computers, if *whence* is not `-1`, `0`, or `1`.

## Also see

[M-5] `cat()` — Load file into string matrix `sprintf()` in

[M-5] `printf()` — Format output

[M-5] `bufio()` — Buffered (binary) I/O

[M-4] `io` — I/O functions

# Title

**[M-5] fullsvd()** — Full singular value decomposition

## Syntax

```
void          fullsvd(numeric matrix A, U, s, Vt)
numeric matrix fullsdiag(numeric colvector s, real scalar k)

void          _fullsvd(numeric matrix A, U, s, Vt)
real scalar  _svd_la(numeric matrix A, U, s, Vt)
```

## Description

`fullsvd(A, U, s, Vt)` calculates the singular value decomposition of  $m \times n$  matrix  $A$ , returning the result in  $U$ ,  $s$ , and  $Vt$ . Singular values in  $s$  are sorted from largest to smallest.

`fullsdiag(s, k)` converts column vector  $s$  returned by `fullsvd()` into matrix  $S$ . In all cases, the appropriate call for this function is

$$S = \text{fullsdiag}(s, \text{rows}(A) - \text{cols}(A))$$

`_fullsvd(A, U, s, Vt)` does the same as `fullsvd()`, except that, in the process, it destroys  $A$ . Use of `_fullsvd()` in place of `fullsvd()` conserves memory.

`_svd_la()` is the interface into the [\[M-1\] LAPACK](#) SVD routines and is used in the implementation of the previous functions. There is no reason you should want to use it. `_svd_la()` is similar to `_fullsvd()`. It differs in that it returns a real scalar equal to 1 if the numerical routines fail to converge, and it returns 0 otherwise. The previous SVD routines set  $s$  to contain missing values in this unlikely case.

## Remarks

Remarks are presented under the following headings:

- [Introduction](#)
- [Relationship between the full and thin SVDs](#)
- [The contents of  \$s\$](#)
- [Possibility of convergence problems](#)

Documented here is the full SVD, appropriate in all cases, but of interest mainly when  $A: m \times n$ ,  $m < n$ . There is a thin SVD that conserves memory when  $m \geq n$ ; see [\[M-5\] svd\(\)](#). The relationship between the two is discussed in [Relationship between the full and thin SVDs](#) below.

## Introduction

The SVD is used to compute accurate solutions to linear systems and least-squares problems, to compute the 2-norm, and to determine the numerical rank of a matrix.

The singular value decomposition (SVD) of  $A$ :  $m \times n$  is given by

$$A = USV'$$

where

- $U$ :

$m \times m$  and orthogonal (unitary)
- $S$ :

$m \times n$  and diagonal
- $V$ :

$n \times n$  and orthogonal (unitary)

When  $A$  is complex, the transpose operator  $'$  is understood to mean the conjugate transpose operator.

Diagonal matrix  $S$  contains the singular values and those singular values are real even when  $A$  is complex. It is usual (but not required) that  $S$  is arranged so that the largest singular value appears first, then the next largest, and so on. The SVD routines documented here do this.

The full SVD routines return  $U$  and  $Vt = V'$ .  $S$  is returned as a column vector  $s$ , and  $S$  can be obtained by

$$S = \text{fullsdiag}(s, \text{rows}(A) - \text{cols}(A))$$

so we will write the SVD as

$$A = U * \text{fullsdiag}(s, \text{rows}(A) - \text{cols}(A)) * Vt$$

Function `fullsvd(A, U, s, Vt)` returns the  $U$ ,  $s$ , and  $Vt$  corresponding to  $A$ .

## Relationship between the full and thin SVDs

A popular variant of the SVD is known as the thin SVD and is suitable for use when  $m \geq n$ . Both SVDs have the same formula,

$$A = USV'$$

but  $U$  and  $S$  have reduced dimensions in the thin version:

Matrix	Full SVD	Thin SVD
$U$ :	$m \times m$	$m \times n$
$S$ :	$m \times n$	$n \times n$
$V$ :	$n \times n$	$n \times n$

When  $m = n$ , the two variants are identical.

The thin SVD is of use when  $m > n$ , because then only the first  $m$  diagonal elements of  $S$  are nonzero, and therefore only the first  $m$  columns of  $U$  are relevant in  $A = USV'$ . There are considerable memory savings to be had in calculating the thin SVD when  $m \gg n$ .

As a result, many people call the thin SVD the SVD and ignore the full SVD altogether. If the matrices you deal with have  $m \geq n$ , you will want to do the same. To obtain the thin SVD, see [M-5] `svd()`.

Regardless of the dimension of your matrix, you may wish to obtain only the singular values. In this case, see `svdsv()` documented in [M-5] `svd()`. That function is appropriate in all cases.



## The contents of `s`

Given  $A: m \times n$ , the singular values are returned in  $s: \min(m, n) \times 1$ .

Let's consider the  $m = n$  case first.  $A$  is  $m \times m$  and the  $m$  singular values are returned in  $s$ , an  $m \times 1$  column vector. If  $A$  were  $3 \times 3$ , perhaps we would get back

```
: s
      1
1  13.47
2   5.8
3   2.63
```

1	13.47
2	5.8
3	2.63

If we needed it, we could obtain  $S$  from  $s$  simply by creating a diagonal matrix from  $s$

```
: S = diag(s)
: S
[symmetric]
      1      2      3
1  13.47
2     0   5.8
3     0     0  2.63
```

1	13.47		
2	0	5.8	
3	0	0	2.63

although the official way we are supposed to do this is

```
: S = fullsdiag(s, rows(A)-cols(A))
```

and that will return the same result.

Now let's consider  $m < n$ . Let's pretend that  $A$  is  $3 \times 4$ . The singular values will be returned in  $3 \times 1$  vector  $s$ . For instance,  $s$  might still contain

```
: s
      1
1  13.47
2   5.8
3   2.63
```

1	13.47
2	5.8
3	2.63

The  $S$  matrix here needs to be  $3 \times 4$ , and `fullsdiag()` will form it:

```
: fullsdiag(s, rows(A)-cols(A))
      1      2      3      4
1  13.47     0     0     0
2     0   5.8     0     0
3     0     0  2.63     0
```

1	13.47	0	0	0
2	0	5.8	0	0
3	0	0	2.63	0

The final case is  $m > n$ . We will pretend that  $A$  is  $4 \times 3$ . The  $s$  vector we get back will look the same

: s	
	1
1	13.47
2	5.8
3	2.63

but this time, we need a  $4 \times 3$  rather than a  $3 \times 4$  matrix formed from it.

: fullsdiag(s, rows(A)-cols(A))			
	1	2	3
1	13.47	0	0
2	0	5.8	0
3	0	0	2.63
4	0	0	0

Possibility of convergence problems

See [Possibility of convergence problems](#) in [M-5] `svd()`; what is said there applies equally here.

Conformability

`fullsvd(A, U, s, Vt):`

*input:*

*A:*  $m \times n$

*output:*

*U:*  $m \times m$

*s:*  $\min(m,n) \times 1$

*Vt:*  $n \times n$

*result:* *void*

`fullsdiag(s, k):`

*input:*

*s:*  $r \times 1$

*k:*  $1 \times 1$

*output:*

*result:*  $r + k \times r$ , if  $k \geq 0$

$r \times r - k$ , otherwise

`_fullsvd(A, U, s, Vt):`

*input:*

*A:*  $m \times n$

*output:*

*A:*  $0 \times 0$

*U:*  $m \times m$

*s:*  $\min(m,n) \times 1$

*Vt:*  $n \times n$

*result:* *void*

`_svd_la(A, U, s, Vt):`

*input:*

$A:$       $m \times n$

*output:*

$A:$       $m \times n$ , but contents changed

$U:$       $m \times m$

$s:$       $\min(m, n) \times 1$

$Vt:$      $n \times n$

*result:*     $1 \times 1$

## Diagnostics

`fullsvd(A, U, s, Vt)` and `_fullsvd(A, s, Vt)` return missing results if  $A$  contains missing. In all other cases, the routines should work, but there is the unlikely possibility of convergence problems, in which case missing results will also be returned; see [Possibility of convergence problems](#) in [M-5] `svd()`.

`_fullsvd()` aborts with error if  $A$  is a view.

Direct use of `_svd_la()` is not recommended.

## Also see

[M-5] `svd()` — Singular value decomposition

[M-5] `svsolve()` — Solve  $AX=B$  for  $X$  using singular value decomposition

[M-5] `pinv()` — Moore–Penrose pseudoinverse

[M-5] `norm()` — Matrix and vector norms

[M-5] `rank()` — Rank of matrix

[M-4] `matrix` — Matrix functions

## Syntax

```
void                                geigensystem(A, B, X, w, b)
```

```
void                                leftgeigensystem(A, B, X, w, b)
```

```
void    geigensystemselectr(A, B, range, X, w, b)
```

```
void leftgeigensystemselectr(A, B, range, X, w, b)
```

```
void    geigensystemselecti(A, B, index, X, w, b)
```

```
void leftgeigensystemselecti(A, B, index, X, w, b)
```

```
void    geigensystemselectf(A, B, f, X, w, b)
```

```
void    leftgeigensystemselectf(A, B, f, X, w, b)
```

where inputs are

<i>A</i> :	<i>numeric matrix</i>	
<i>B</i> :	<i>numeric matrix</i>	
<i>range</i> :	<i>real vector</i>	(range of generalized eigenvalues to be selected)
<i>index</i> :	<i>real vector</i>	(indices of generalized eigenvalues to be selected)
<i>f</i> :	<i>pointer scalar</i>	(points to a function used to select generalized eigenvalues)

and outputs are

<i>X</i> :	<i>numeric matrix</i> of generalized eigenvectors
<i>w</i> :	<i>numeric vector</i> (numerators of generalized eigenvalues)
<i>b</i> :	<i>numeric vector</i> (denominators of generalized eigenvalues)

The following routines are used in implementing the above routines:

```
void _geigensystem_la(numeric matrix H, R, XL, XR, w, b,  
                      string scalar side)
```

```
void _geigenselectr_la(numeric matrix H, R, XL, XR, w, b,  
                       range, string scalar side)
```

```
void _geigenselecti_la(numeric matrix H, R, XL, XR, w, b,  
                       index, string scalar side)
```

```
void _geigenselectf_la(numeric matrix H, R, XL, XR, w, b,  
                       pointer scalar f, string scalar side)
```

```
real scalar _geigen_la(numeric matrix H, R, XL, XR, w, select,  
                       string scalar side, string scalar howmany)
```

## Description

`geigensystem(A, B, X, w, b)` computes [generalized eigenvectors](#) of two general, real or complex, square matrices,  $A$  and  $B$ , along with their corresponding [generalized eigenvalues](#).

- $A$  and  $B$  are two general, real or complex, square matrices with the same dimensions.
- $X$  contains generalized eigenvectors.
- $w$  contains numerators of generalized eigenvalues.
- $b$  contains denominators of generalized eigenvalues.

`leftgeigensystem(A, B, X, w, b)` mirrors `geigensystem()`, the difference being that `leftgeigensystem()` computes left, generalized eigenvectors.

`geigensystemselectr(A, B, range, X, w, b)` computes selected generalized eigenvectors of two general, real or complex, square matrices,  $A$  and  $B$ , along with their corresponding generalized eigenvalues. Only the generalized eigenvectors corresponding to selected generalized eigenvalues are computed. Generalized eigenvalues that lie in a [range](#) are selected. The selected generalized eigenvectors are returned in  $X$ , and their corresponding generalized eigenvalues are returned in  $(w, b)$ .

*range* is a vector of length 2. All finite, generalized eigenvalues with absolute value in the half-open interval  $(range[1], range[2]]$  are selected.

`leftgeigensystemselectr(A, B, range, X, w, b)` mirrors `geigensystemselectr()`, the difference being that `leftgeigensystemr()` computes left, generalized eigenvectors.

`geigensystemselecti(A, B, index, X, w, b)` computes selected right, generalized eigenvectors of two general, real or complex, square matrices,  $A$  and  $B$ , along with their corresponding generalized eigenvalues. Only the generalized eigenvectors corresponding to selected generalized eigenvalues are computed. Generalized eigenvalues are selected by an [index](#). The selected generalized eigenvectors are returned in  $X$ , and the selected generalized eigenvalues are returned in  $(w, b)$ .

The finite, generalized eigenvalues are sorted by their absolute values, in descending order, followed by the infinite, generalized eigenvalues. There is no particular order among infinite, generalized eigenvalues.

*index* is a vector of length 2. The generalized eigenvalues in elements  $index[1]$  through  $index[2]$ , inclusive, are selected.

`leftgeigensystemselecti(A, B, index, X, w, b)` mirrors `geigensystemselecti()`, the difference being that `leftgeigensystemi()` computes left, generalized eigenvectors.

`geigensystemselectf(A, B, f, X, w, b)` computes selected generalized eigenvectors of two general, real or complex, square matrices  $A$  and  $B$  along with their corresponding generalized eigenvalues. Only the generalized eigenvectors corresponding to selected generalized eigenvalues are computed. Generalized eigenvalues are selected by a user-written function described [below](#). The selected generalized eigenvectors are returned in  $X$ , and the selected generalized eigenvalues are returned in  $(w, b)$ .

`leftgeigensystemselectf(A, B, f, X, w, b)` mirrors `geigensystemselectf()`, the difference being that `leftgeigensystemselectf()` computes selected left, generalized eigenvectors.

`_geigen_la()`, `_geigensystem_la()`, `_geigenselectr_la()`, `_geigenselecti_la()`, and `_geigenselectf_la()` are the interfaces into the LAPACK routines used to implement the above functions; see [M-1] **LAPACK**. Their direct use is not recommended.

## Remarks

Remarks are presented under the following headings:

*Generalized eigenvalues*  
*Generalized eigenvectors*  
*Criterion selection*  
*Range selection*  
*Index selection*

## Generalized eigenvalues

A scalar,  $l$  (usually denoted by *lambda*), is said to be a generalized eigenvalue of a pair of  $n \times n$  square, numeric matrices  $(\mathbf{A}, \mathbf{B})$  if there is a nonzero column vector  $\mathbf{x}$ :  $n \times 1$  (called the generalized eigenvector) such that

$$\mathbf{A}\mathbf{x} = l\mathbf{B}\mathbf{x} \quad (1)$$

(1) can also be written

$$(\mathbf{A} - l\mathbf{B})\mathbf{x} = 0$$

A nontrivial solution to this system of  $n$  linear homogeneous equations exists if and only if

$$\det(\mathbf{A} - l\mathbf{B}) = 0 \quad (2)$$

In practice, the generalized eigenvalue problem for the matrix pair  $(\mathbf{A}, \mathbf{B})$  is usually formulated as finding a pair of scalars  $(w, b)$  and a nonzero column vector  $\mathbf{x}$  such that

$$w\mathbf{A}\mathbf{x} = b\mathbf{B}\mathbf{x}$$

The scalar  $w/b$  is a finite, generalized eigenvalue if  $b$  is not zero. The pair  $(w, b)$  represents an infinite, generalized eigenvalue if  $b$  is zero or numerically close to zero. This situation may arise if  $\mathbf{B}$  is singular.

The Mata functions that compute generalized eigenvalues return them in two complex vectors,  $\mathbf{w}$  and  $\mathbf{b}$ , of length  $n$ . If  $\mathbf{b}[i]=0$ , the  $i$ th generalized eigenvalue is infinite; otherwise, the  $i$ th generalized eigenvalue is  $\mathbf{w}[i]/\mathbf{b}[i]$ .

## Generalized eigenvectors

A column vector,  $\mathbf{x}$ , is a right, generalized eigenvector or simply a generalized eigenvector of a generalized eigenvalue  $(w, b)$  for a pair of matrices,  $\mathbf{A}$  and  $\mathbf{B}$ , if

$$w\mathbf{A}\mathbf{x} = b\mathbf{B}\mathbf{x}$$

A row vector,  $\mathbf{v}$ , is a left, generalized eigenvector of a generalized eigenvalue  $(w, b)$  for a pair of matrices,  $\mathbf{A}$  and  $\mathbf{B}$ , if

$$w\mathbf{v}\mathbf{A} = b\mathbf{v}\mathbf{B}$$

For instance, let's consider the linear system

$$dx/dt = A1 \times x + A2 \times u$$

$$dy/dt = A3 \times x + A4 \times u$$

where

```
: A1 = (-4, -3 \ 2, 1)
```

```
: A2 = (3 \ 1)
```

```
: A3 = (1, 2)
```

and

```
: A4 = 0
```

The finite solutions of zeros for the transfer function

$$g(s) = A3 \times (sI - A1)^{-1} \times A2 + A4 \quad (3)$$

of this linear time-invariant state-space model is given by the finite, generalized eigenvalues of A and B where

```
: A = (A1, A2 \ A3, A4)
```

and

```
: B = (1, 0, 0 \ 0, 1, 0 \ 0, 0, 0)
```

We obtain generalized eigenvectors in X and generalized eigenvalues in w and b by using

```
: geigensystem(A, B, X=., w=., b=.)
```

```
: X
```

	1	2	3
1	-1	0	2.9790e-16
2	.5	0	9.9301e-17
3	.1	1	1

```
: w
```

	1	2	3
1	-1.97989899	3.16227766	2.23606798

```
: b
```

	1	2	3
1	.7071067812	0	0

The only finite, generalized eigenvalue of A and B is

```
: w[1,1]/b[1,1]
-2.8
```

In this simple example, (3) can be explicitly written out as

$$g(s) = (5s + 14)/(s^2 + 3s + 2)$$

which clearly has the solution of zero at  $-2.8$ .

## Criterion selection

We sometimes want to compute only those generalized eigenvectors whose corresponding generalized eigenvalues satisfy certain criterion. We can use `geigensystemselectf()` to solve these problems.

We must pass `geigensystemselectf()` a [pointer](#) to a function that implements our conditions. The function must accept two numeric scalar arguments so that it can receive the numerator `w` and the denominator `b` of a generalized eigenvalue, and it must return the real value 0 to indicate rejection and a nonzero real value to indicate selection.

In this example, we want to compute only finite, generalized eigenvalues for each of which `b` is not zero. After deciding that anything smaller than  $1e-15$  is zero, we define our function to be

```
: real scalar finiteonly(numeric scalar w, numeric scalar b)
> {
>     return((abs(b)>=1e-15))
> }
```

By using

```
: geigensystemselectf(A, B, &finiteonly(), X=., w=., b=.)
```

we get the only finite, generalized eigenvalue of `A` and `B` in `(w, b)` and its corresponding eigenvector in `X`:

```
: X
      1
1  - .894427191
2   .447213595
3   .089442719

: w
-1.97989899

: b
.7071067812

: w:/b
-2.8
```

## Range selection

We can use `geigensystemselectr()` to compute only those generalized eigenvectors whose generalized eigenvalues have absolute values that fall in a half-open interval.

For instance,

```
: A = (-132, -88, 84, 104 \ -158.4, -79.2, 76.8, 129.6 \
> 129.6, 81.6, -79.2, -100.8 \ 160, 84, -80, -132)
: B = (-60, -50, 40, 50 \ -69, -46.4, 38, 58.2 \ 58.8, 46, -37.6, -48 \
> 70, 50, -40, -60)
: range = (0.99, 2.1)
```



We obtain generalized eigenvectors in `X` and generalized eigenvalues in `w` and `b` by using

```
: geigensystemselectr(A, B, range, X=., w=., b=.)
: X
      1      2
1  .089442719 .02236068
2  .04472136 .067082039
3  .04472136 .067082039
4  .089442719 .02236068

: w
      1      2
1  .02820603 .170176379

: b
      1      2
1  .0141030148 .1701763791
```

The generalized eigenvalues have absolute values in the half-open interval  $(0.99, 2.1]$ .

```
: abs(w:/b)
      1      2
1  2      1
```

## Index selection

`geigensystemselect()` sorts the finite, generalized eigenvalues using their absolute values, in descending order, placing the infinite, generalized eigenvalues after the finite, generalized eigenvalues. There is no particular order among infinite, generalized eigenvalues.

If we want to compute only generalized eigenvalues whose ranks are `index[1]` through `index[2]` in the list of generalized eigenvalues obtained by `geigensystemselect()`, we can use `geigensystemselecti()`.

To compute the first two generalized eigenvalues and generalized eigenvectors in this example, we can specify

```
: index = (1, 2)
: geigensystemselecti(A, B, index, X=., w=., b=.)
```

The results are

```
: X
      1      2
1  .02981424 -.059628479
2  .04472136 -.059628479
3  .089442719 -.02981424
4  .01490712 -.119256959
```

```
: w
      1      2
1  [.012649111 .379473319]

: b
      1      2
1  [.0031622777 .1264911064]

: w:/b
      1      2
1  [ 4      3]
```

Conformability

```
geigensystem(A, B, X, w, b):
  input:
    A:  n × n
    B:  n × n
  output:
    X:  n × n
    w:  1 × n
    b:  1 × n

leftgeigensystem(A, B, X, w, b):
  input:
    A:  n × n
    B:  n × n
  output:
    X:  n × n
    w:  1 × n
    b:  1 × n

geigensystemselectr(A, B, range, X, w, b):
  input:
    A:  n × n
    B:  n × n
    range:  1 × 2 or 2 × 1
  output:
    X:  n × m
    w:  1 × m
    b:  1 × m
```

`leftgeigensystemselectr(A, B, range, X, w, b):`

*input:*

$A:$       $n \times n$   
 $B:$       $n \times n$   
 $range:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $m \times n$   
 $w:$       $1 \times m$   
 $b:$       $1 \times m$

`geigensystemselecti(A, B, index, X, w, b):`

*input:*

$A:$       $n \times n$   
 $B:$       $n \times n$   
 $index:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $n \times m$   
 $w:$       $1 \times m$   
 $b:$       $1 \times m$

`leftgeigensystemselecti(A, B, index, X, w, b):`

*input:*

$A:$       $n \times n$   
 $B:$       $n \times n$   
 $index:$     $1 \times 2$  or  $2 \times 1$

*output:*

$X:$       $m \times n$   
 $w:$       $1 \times m$   
 $b:$       $1 \times m$

`geigensystemselectf(A, B, f, X, w, b):`

*input:*

$A:$       $n \times n$   
 $B:$       $n \times n$   
 $f:$       $1 \times 1$

*output:*

$X:$       $n \times m$   
 $w:$       $1 \times m$   
 $b:$       $1 \times m$

`leftgeigensystemselectf(A, B, f, X, w, b):`

*input:*

$A:$       $n \times n$   
 $B:$       $n \times n$   
 $f:$       $1 \times 1$

*output:*

$X:$       $m \times n$   
 $w:$       $1 \times m$   
 $b:$       $1 \times m$

## Diagnostics

All functions return missing-value results if  $A$  or  $B$  has missing values.

## References

- Gould, W. W. 2011a. Understanding matrices intuitively, part 1. The Stata Blog: Not Elsewhere Classified. <http://blog.stata.com/2011/03/03/understanding-matrices-intuitively-part-1/>
- . 2011b. Understanding matrices intuitively, part 2, eigenvalues and eigenvectors. The Stata Blog: Not Elsewhere Classified. <http://blog.stata.com/2011/03/09/understanding-matrices-intuitively-part-2/>

## Also see

- [M-1] **LAPACK** — The LAPACK linear-algebra routines
- [M-5] **geigensystem()** — Generalized eigenvectors and eigenvalues
- [M-5] **ghessenbergd()** — Generalized Hessenberg decomposition
- [M-5] **gschurd()** — Generalized Schur decomposition
- [M-4] **matrix** — Matrix functions

Syntax

```
void ghessenbergd(numeric matrix A, B, H, R, U, V)
void _ghessenbergd(numeric matrix A, B, U, V)
```

Description

ghessenbergd(A, B, H, R, U, V) computes the generalized Hessenberg decomposition of two general, real or complex, square matrices, A and B, returning the upper Hessenberg form matrix in H, the upper triangular matrix in R, and the orthogonal (unitary) matrices in U and V.

\_ghessenbergd(A, B, U, V) mirrors ghessenbergd(), the difference being that it returns H in A and R in B.

\_ghessenbergd\_la() is the interface into the LAPACK routines used to implement the above function; see [M-1] LAPACK. Its direct use is not recommended.

Remarks

The generalized Hessenberg decomposition of two square, numeric matrices (**A** and **B**) can be written as

$$\begin{aligned} \mathbf{U}' \times \mathbf{A} \times \mathbf{V} &= \mathbf{H} \\ \mathbf{U}' \times \mathbf{B} \times \mathbf{V} &= \mathbf{R} \end{aligned}$$

where **H** is in upper Hessenberg form, **R** is upper triangular, and **U** and **V** are orthogonal matrices if **A** and **B** are real or are unitary matrices otherwise.

In the example below, we define A and B, obtain the generalized Hessenberg decomposition, and list H and Q.

```
: A = (6, 2, 8, -1\ -3, -4, -6, 4\ 0, 8, 4, 1\ -8, -7, -3, 5)
: B = (8, 0, -8, -1\ -6, -2, -6, -1\ -7, -6, 2, -6\ 1, -7, 9, 2)
: ghessenbergd(A, B, H=., R=., U=., V=.)
: H
      1      2      3      4
1  -4.735680169  1.363736029  5.097381347  3.889763589
2   9.304479208 -8.594240253 -7.993282943  4.803411217
3      0  4.553169015  3.236266637 -2.147709419
4      0      0  6.997043028 -3.524816722

: R
      1      2      3      4
1 -12.24744871 -1.089095534 -1.848528639 -5.398470103
2      0 -5.872766311  8.891361089  3.86967647
3      0      0  9.056748937  1.366322731
4      0      0      0  8.357135399
```

## Conformability

`ghessenbergd(A, B, H, R, U, V):`

*input:*

*A:*      $n \times n$

*B:*      $n \times n$

*output:*

*H:*      $n \times n$

*R:*      $n \times n$

*U:*      $n \times n$

*V:*      $n \times n$

`_ghessenbergd(A, B, U, V):`

*input:*

*A:*      $n \times n$

*B:*      $n \times n$

*output:*

*A:*      $n \times n$

*B:*      $n \times n$

*U:*      $n \times n$

*V:*      $n \times n$

## Diagnostics

`_ghessenbergd()` aborts with error if *A* or *B* is a view.

`ghessenbergd()` and `_ghessenbergd()` return missing results if *A* or *B* contains missing values.

## Also see

[M-1] **LAPACK** — The LAPACK linear-algebra routines

[M-5] **gschurd()** — Generalized Schur decomposition

[M-4] **matrix** — Matrix functions

Syntax

```
S = ghk_init(real scalar npts)

(varies)    ghk_init_method(S [ , string scalar method])
(varies)    ghk_init_start(S [ , real scalar start])
(varies)    ghk_init_pivot(S [ , real scalar pivot])
(varies)    ghk_init_antithetics(S [ , real scalar anti])
real scalar ghk_query_npts(S)
real scalar ghk(S, real vector x, V)
real scalar ghk(S, real vector x, V, real rowvector dfdx, dfdv)
```

where *S*, if declared, should be declared

```
transmorphic S
```

and where *method*, optionally specified in `ghk_init_method()`, is

<i>method</i>	Description
"halton"	Halton sequences
"hammersley"	Hammersley's variation of the Halton set
"random"	pseudorandom uniforms

Description

`S = ghk_init(npts)` constructs the transmorphic object *S*. Calls to `ghk_init_method(S, method)`, `ghk_init_start(S, start)`, `ghk_init_pivot(S, pivot)`, and `ghk_init_antithetics(S, anti)` prior to calling `ghk(S, ...)` allow you to modify the simulation algorithm through the object *S*.

`ghk(S, x, V)` returns a real scalar containing the simulated value of the multivariate normal (MVN) distribution with variance–covariance *V* at the point *x*. First, code `S = ghk_init(npts)` and then use `ghk(S, ...)` to obtain the simulated value based on *npts* simulation points.

`ghk(S, x, V, dfdx, dfdv)` does the same thing but also returns the first-order derivatives of the simulated probability with respect to *x* in *dfdx* and the simulated probability derivatives with respect to `vech(V)` in *dfdv*. See `vech()` in [M-5] `vec()` for details of the half-vectorized operator.

The `ghk_query_npts(S)` function returns the number of simulation points, the same value given in the construction of the transmorphic object *S*.

## Remarks

Halton and Hammersley point sets are composed of deterministic sequences on  $[0,1]$  and, for sets of dimension less than 10, generally have better coverage than that of the uniform pseudorandom sequences.

Antithetic draws effectively double the number of points and reduce the variability of the simulated probability. For draw  $u$ , the antithetic draw is  $1 - u$ . To use antithetic draws, call `ghk_init_antithetic(S, 1)` prior to executing `ghk()`.

By default, `ghk()` will pivot the wider intervals of integration (and associated rows/columns of the covariance matrix) to the interior of the multivariate integration. This improves the accuracy of the quadrature estimate. When `ghk()` is used in a likelihood evaluator for [R] `ml` or [M-5] `optimize()`, discontinuities may result in the computation of numerical second-order derivatives using finite differencing (for the Newton–Raphson optimize technique) when few simulation points are used, resulting in a non-positive-definite Hessian. To turn off the interval pivoting, call `ghk_init_pivot(S, 0)` prior to executing `ghk()`.

If you are using `ghk()` in a likelihood evaluator, be sure to use the same sequence with each call to the likelihood evaluator. For a uniform pseudorandom sequence, `ghk_init_method("random")`, set the seed of the uniform random-variate generator—see `rseed()` in [M-5] `runiform()`—to the same value with each call to the likelihood evaluator.

If you are using the Halton or Hammersley point sets, you will want to keep the sequences going with each call to `ghk()` within one likelihood evaluation. This can be done in one expression executed after each call to `ghk()`: `ghk_init_start(S, ghk_init_start(S) + ghk_query_npts(S))`. With each call to the likelihood evaluator, you will need to reset the starting index to 1. This last point assumes that the transmorphic object  $S$  is not re-created on each call to the likelihood evaluator.

Unlike `ghkfast_init()` (see [M-5] `ghkfast()`), the transmorphic object  $S$  created by `ghk_init()` is inexpensive to create, so it is possible to re-create it with each call to your likelihood evaluator instead of storing it as `external` global and reusing the object with each likelihood evaluation. Alternatively, the initialization function for `optimize()`, `optimize_init_arguments()`, can be used.



## Conformability

All initialization functions have  $1 \times 1$  inputs and have  $1 \times 1$  or *void* outputs except

`ghk_init(npts)`:

*input*:

*npts*:  $1 \times 1$

*output*:

*S*: transmorphic

`ghk_query_npts(S)`:

*input*:

*S*: transmorphic

*output*:

*result*:  $1 \times 1$

`ghk(S, x, V)`:

*input*:

*S*: transmorphic

*x*:  $1 \times m$  or  $m \times 1$

*V*:  $m \times m$  (symmetric, positive definite)

*output*:

*result*:  $1 \times 1$

`ghk(S, x, V, dfdx, dfdv)`:

*input*:

*S*: transmorphic

*x*:  $1 \times m$  or  $m \times 1$

*V*:  $m \times m$  (symmetric, positive definite)

*output*:

*result*:  $1 \times 1$

*dfdx*:  $1 \times m$

*dfdv*:  $1 \times m(m+1)/2$

## Diagnostics

The maximum dimension,  $m$ , is 20.

$V$  must be symmetric and positive definite. `ghk()` will return a missing value when  $V$  is not positive definite. When `ghk()` is used in an `m1` (or `optimize()`) likelihood evaluator, return a missing likelihood to `m1` and let `m1` take the appropriate action (that is, step halving).

## Also see

[M-5] [ghkfast\(\)](#) — GHK multivariate normal simulator using pregenerated points

[M-5] [halton\(\)](#) — Generate a Halton or Hammersley set

[M-4] [statistical](#) — Statistical functions

Syntax

$S = \text{ghkfast\_init}(\text{real scalar } n, \text{npts}, \text{dim}, \text{string scalar method})$

(varies)  $\text{ghkfast\_init\_pivot}(S \text{ [ , real scalar pivot ]})$   
(varies)  $\text{ghkfast\_init\_antithetics}(S \text{ [ , real scalar anti ]})$   
real scalar  $\text{ghkfast\_query\_n}(S)$   
real scalar  $\text{ghkfast\_query\_npts}(S)$   
real scalar  $\text{ghkfast\_query\_dim}(S)$   
string scalar  $\text{ghkfast\_query\_method}(S)$   
string scalar  $\text{ghkfast\_query\_rseed}(S)$   
real matrix  $\text{ghkfast\_query\_pointset\_i}(S, i)$   
real colvector  $\text{ghkfast}(S, \text{real matrix } X, V)$   
real colvector  $\text{ghkfast}(S, \text{real matrix } X, V, \text{dfdx}, \text{dfdv})$   
real scalar  $\text{ghkfast\_i}(S, \text{real matrix } X, V, i)$   
real scalar  $\text{ghkfast\_i}(S, \text{real matrix } X, V, i, \text{dfdx}, \text{dfdv})$

where  $S$ , if it is declared, should be declared

transmorphic  $S$

and where *method* specified in `ghkfast_init()` is

<i>method</i>	Description
"halton"	Halton sequences
"hammersley"	Hammersley's variation of the Halton set
"random"	pseudorandom uniforms
"ghalton"	generalized Halton sequences

Description

Please see [M-5] [ghk\(\)](#). The routines documented here do the same thing, but `ghkfast()` can be faster at the expense of using more memory. First, code  $S = \text{ghkfast\_init}(\dots)$  and then use  $\text{ghkfast}(S, \dots)$  to obtain the simulated values. There is a time savings because the simulation points are generated once in `ghkfast_init()`, whereas for `ghk()` the points are generated on each call to `ghk()`. Also, `ghkfast()` can generate simulated probabilities from the generalized Halton sequence; see [M-5] [halton\(\)](#).

`ghkfast_init(n, npts, dim, method)` computes the simulation points to be used by `ghkfast()`. Inputs *n*, *npts*, and *dim* are the number of observations, the number of repetitions for the simulation, and the maximum dimension of the multivariate normal (MVN) distribution, respectively. Input *method* specifies the type of points to generate and can be one of "halton", "hammersley", "random", or "ghalton".

`ghkfast(S, X, V)` returns an  $n \times 1$  real vector containing the simulated values of the MVN distribution with  $\text{dim} \times \text{dim}$  variance–covariance matrix *V* at the points stored in the rows of the  $n \times \text{dim}$  matrix *X*.

`ghkfast(S, X, V, dfdx, dfdv)` does the same thing as `ghkfast(S, X, V)` but also returns the first-order derivatives of the simulated probability with respect to the rows of *X* in *dfdx* and the simulated probability derivatives with respect to `vech(V)` in *dfdv*. See `vech()` in [M-5] `vec()` for details of the half-vectorized operator.

The `ghk_query_n(S)`, `ghk_query_npts(S)`, `ghk_query_dim(S)`, and `ghk_query_method(S)` functions extract the number of observations, number of simulation points, maximum dimension, and method of point-set generation that is specified in the construction of the transmorphic object *S*. Use `ghk_query_rseed(S)` to retrieve the uniform random-variate seed used to generate the "random" or "ghalton" point sets. The `ghkfast_query_pointset_i(S, i)` function will retrieve the *i*th point set used to simulate the MVN probability for the *i*th observation.

The `ghkfast_i(S, X, V, i, ...)` function computes the probability and derivatives for the *i*th observation,  $i = 1, \dots, n$ .

## Remarks

For problems where repetitive calls to the GHK algorithm are required, `ghkfast()` might be a preferred alternative to `ghk()`. Generating the points once at the outset of a program produces a speed increase. For problems with many observations or many simulation points per observation, `ghkfast()` will be faster than `ghk()` at the cost of requiring more memory.

If `ghkfast()` is used within a likelihood evaluator for `ml` or `optimize()`, you will need to store the transmorphic object *S* as an [external](#) global and reuse the object with each likelihood evaluation. Alternatively, the initialization function for `optimize()`, `optimize_init_arguments()`, can be used.

Prior to calling `ghkfast()`, call `ghkfast_init_npivot(S, 1)` to turn off the integration interval pivoting that takes place in `ghkfast()`. By default, `ghkfast()` pivots the wider intervals of integration (and associated rows/columns of the covariance matrix) to the interior of the multivariate integration to improve quadrature accuracy. This option may be useful when `ghkfast()` is used in a likelihood evaluator for [R] `ml` or [M-5] `optimize()` and few simulation points are used for each observation. Here the pivoting may cause discontinuities when computing numerical second-order derivatives using finite differencing (for the Newton–Raphson technique), resulting in a non–positive-definite Hessian.

Also the sequences "halton", "hammersley", and "random", `ghkfast()` will use the generalized Halton sequence, "ghalton". Generalized Halton sequences have the same uniform coverage (low discrepancy) as the Halton sequences with the addition of a pseudorandom uniform component. Therefore, "ghalton" sequences are like "random" sequences in that you should set the random-number seed before using them if you wish to replicate the same point set; see [M-5] `runiform()`.

## Conformability

All initialization functions have  $1 \times 1$  inputs and have  $1 \times 1$  or *void* outputs, and all query functions have the *transmorphic* input and  $1 \times 1$  outputs except

`ghkfast_init(n, npts, dim, method):`

input:

*n*:  $1 \times 1$   
*npts*:  $1 \times 1$   
*dim*:  $1 \times 1$   
*method*:  $1 \times 1$

output:

result: *transmorphic*

`ghkfast_query_pointset_i(S, i):`

input:

*S*: *transmorphic*  
*i*:  $1 \times 1$

output:

result:  $npts \times dim$

`ghkfast(S, X, V):`

input:

*S*: *transmorphic*  
*X*:  $n \times dim$   
*V*:  $dim \times dim$  (symmetric, positive definite)

output:

result:  $n \times 1$

`ghkfast(S, X, V, dfdx, dfdv):`

input:

*S*: *transmorphic*  
*X*:  $n \times dim$   
*V*:  $dim \times dim$  (symmetric, positive definite)

output:

result:  $n \times 1$   
*dfdx*:  $n \times dim$   
*dfdv*:  $n \times dim(dim + 1)/2$

`ghkfast_i(S, X, V, i, dfdx, dfdv):`

input:

*S*: *transmorphic*  
*X*:  $n \times dim$  or  $1 \times dim$   
*V*:  $dim \times dim$  (symmetric, positive definite)  
*i*:  $1 \times 1$  ( $1 \leq i \leq n$ )

output:

result:  $n \times 1$   
*dfdx*:  $1 \times dim$   
*dfdv*:  $1 \times dim(dim + 1)/2$

## Diagnostics

`ghkfast_init(n, npts, dim, method)` aborts with error if the dimension, *dim*, is greater than 20.

`ghkfast(S, X, V, ...)` and `ghkfast_i(S, X, V, i, ...)` require that *V* be symmetric and positive definite. If *V* is not positive definite, then the returned vector (scalar) is filled with missings.

## Also see

[M-5] [ghk\(\)](#) — Geweke–Hajivassiliou–Keane (GHK) multivariate normal simulator

[M-5] [halton\(\)](#) — Generate a Halton or Hammersley set

[M-4] [statistical](#) — Statistical functions

## Syntax

```

void          gschurd(A, B, T, R, U, V, w, b)
void          _gschurd(A, B, U, V, w, b)
void  gschurdgroupby(A, B, f, T, R, U, V, w, b, m)
void  _gschurdgroupby(A, B, f, U, V, w, b, m)
    
```

## Description

`gschurd(A, B, T, R, U, V, w, b)` computes the generalized Schur decomposition of two square, numeric matrices, *A* and *B*, and the [generalized eigenvalues](#). The decomposition is returned in the [Schur-form](#) matrix, *T*; the upper-triangular matrix, *R*; and the orthogonal (unitary) matrices, *U* and *V*. The generalized eigenvalues are returned in the complex vectors *w* and *b*.

`gschurdgroupby(A, B, f, T, R, U, V, w, b, m)` computes the generalized Schur decomposition of two square, numeric matrices, *A* and *B*, and the [generalized eigenvalues](#), and groups the results according to whether a condition on each generalized eigenvalue is satisfied. *f* is a pointer to the function that implements the condition on each generalized eigenvalue, as discussed [below](#). The number of generalized eigenvalues for which the condition is true is returned in *m*.

`_gschurd()` mirrors `gschurd()`, the difference being that it returns *T* in *A* and *R* in *B*.

`_gschurdgroupby()` mirrors `gschurdgroupby()`, the difference being that it returns *T* in *A* and *R* in *B*.

`_gschurd_la()` and `_gschurdgroupby_la()` are the interfaces into the LAPACK routines used to implement the above functions; see [\[M-1\] LAPACK](#). Their direct use is not recommended.

## Remarks

Remarks are presented under the following headings:

[Generalized Schur decomposition](#)  
[Grouping the results](#)

## Generalized Schur decomposition

The generalized Schur decomposition of a pair of square, numeric matrices, **A** and **B**, can be written as

$$\mathbf{U}' \times \mathbf{A} \times \mathbf{V} = \mathbf{T}$$

$$\mathbf{U}' \times \mathbf{B} \times \mathbf{V} = \mathbf{R}$$

where **T** is in Schur form, **R** is upper triangular, and **U** and **V** are orthogonal if **A** and **B** are real and are unitary if **A** or **B** is complex. The complex vectors **w** and **b** contain the generalized eigenvalues.

If **A** and **B** are real, **T** is in real Schur form and **R** is a real upper-triangular matrix. If **A** or **B** is complex, **T** is in complex Schur form and **R** is a complex upper-triangular matrix.

In the example below, we define **A** and **B**, obtain the generalized Schur decomposition, and list **T** and **R**.

```
: A = (6, 2, 8, -1\ -3, -4, -6, 4\ 0, 8, 4, 1\ -8, -7, -3, 5)
: B = (8, 0, -8, -1\ -6, -2, -6, -1\ -7, -6, 2, -6\ 1, -7, 9, 2)
: gschurd(A, B, T=., R=., U=., V=., w=., b=.)
: T
```

	1	2	3	4
1	12.99313938	1.746927947	3.931212285	-10.91622337
2	0	.014016016	6.153566902	1.908835695
3	0	-4.362999645	1.849905717	-2.998194791
4	0	0	0	-5.527285433

```
: R
```

	1	2	3	4
1	4.406836593	6.869534063	-1.840892081	1.740906311
2	0	13.88730687	0	-.6995556735
3	0	0	9.409495218	-4.659386723
4	0	0	0	9.453808732

```
: w
```

	1	2	3	4
1	12.9931394	.409611804+1.83488354i	.024799819-.111092453i	-5.52728543

```
: b
```

	1	2	3	4
1	4.406836593	4.145676341	.2509986829	9.453808732

Generalized eigenvalues can be obtained by typing

```
: w:/b
```

	1	2	3	4
1	2.94840508	.098804579+.442601735i	.098804579-.442601735i	-.584662287

## Grouping the results

`gschurddgroupby()` reorders the generalized Schur decomposition so that a selected group of generalized eigenvalues appears in the leading block of the pair **w** and **b**. It also reorders the generalized Schur form **T**, **R**, and orthogonal (unitary) matrices, **U** and **V**, correspondingly.

We must pass `gschurddgroupby()` a [pointer](#) to a function that implements our criterion. The function must accept two arguments, a complex scalar and a real scalar, so that it can receive a generalized eigenvalue, and it must return the real value 0 to indicate rejection and a nonzero real value to indicate selection.

In the following example, we use `gschurddgroupby()` to put the finite, real, generalized eigenvalues first. One of the arguments to `schurddgroupby()` is a pointer to the function `onlyreal()` which

accepts two arguments, a complex scalar and a real scalar that define a generalized eigenvalue. `onlyreal()` returns 1 if the generalized eigenvalue is finite and real; it returns zero otherwise.

```

: real scalar onlyreal(complex scalar w, real scalar b)
> {
>     if(b==0) return(0)
>     if(Im(w/b)==0) return(1)
>     return(0)
> }
: gschurdgroupby(A, B, &onlyreal(), T=., R=., U=., V=., w=., b=., m=.)

```

We obtain

```

: T
      1      2      3      4
1  12.99313938   8.19798168   6.285710813   5.563547054
2      0  -5.952366071  -1.473533834   2.750066482
3      0      0  -0.2015830885   3.882051743
4      0      0   6.337230739   1.752690714

: R
      1      2      3      4
1  4.406836593   2.267479575  -6.745927817   1.720793701
2      0  10.18086202  -2.253089622   5.74882307
3      0      0  -12.5704981      0
4      0      0      0   9.652818299

: w
      1      2      3      4
1  12.9931394  -5.95236607  .36499234+1.63500766i  .36499234-1.63500766i

: b
      1      2      3      4
1  4.406836593  10.18086202  3.694083258  3.694083258

: w:/b
      1      2      3      4
1  2.94840508  -.584662287  .098804579+.442601735i  .098804579-.442601735i

```

`m` contains the number of real, generalized eigenvalues

```

: m
2

```



## Conformability

`gschurd(A, B, T, R, U, V, w, b):`

*input:*

*A:*      $n \times n$

*B:*      $n \times n$

*output:*

*T:*      $n \times n$

*R:*      $n \times n$

*U:*      $n \times n$

*V:*      $n \times n$

*w:*      $1 \times n$

*b:*      $1 \times n$

`_gschurd(A, B, U, V, w, b):`

*input:*

*A:*      $n \times n$

*B:*      $n \times n$

*output:*

*A:*      $n \times n$

*B:*      $n \times n$

*U:*      $n \times n$

*V:*      $n \times n$

*w:*      $1 \times n$

*b:*      $1 \times n$

`gschurdgroupby(A, B, f, T, R, U, V, w, b, m):`

*input:*

*A:*      $n \times n$

*B:*      $n \times n$

*f:*      $1 \times 1$

*output:*

*T:*      $n \times n$

*R:*      $n \times n$

*U:*      $n \times n$

*V:*      $n \times n$

*w:*      $1 \times n$

*b:*      $1 \times n$

*m:*      $1 \times 1$

`_gschurdgroupby(A, B, f, U, V, w, b, m)`:

*input:*

*A:*      $n \times n$   
*B:*      $n \times n$   
*f:*      $1 \times 1$

*output:*

*A:*      $n \times n$   
*B:*      $n \times n$   
*U:*      $n \times n$   
*V:*      $n \times n$   
*w:*      $1 \times n$   
*b:*      $1 \times n$   
*m:*      $1 \times 1$

## Diagnostics

`_gschurd()` and `_gschurdgroupby()` abort with error if *A* or *B* is a view.

`gschurd()`, `_gschurd()`, `gschurdgroupby()`, and `_gschurdgroupby()` return missing results if *A* or *B* contains missing values.

## Also see

- [M-1] [LAPACK](#) — The LAPACK linear-algebra routines
- [M-5] [ghessenbergd\(\)](#) — Generalized Hessenberg decomposition
- [M-5] [geigensystem\(\)](#) — Generalized eigenvectors and eigenvalues
- [M-4] [matrix](#) — Matrix functions

## Syntax

```

real matrix    halton(real scalar n, real scalar d)
real matrix    halton(real scalar n, real scalar d, real scalar start)
real matrix    halton(real scalar n, real scalar d, real scalar start,
                        real scalar hammersley)

void           _halton(real matrix x)
void           _halton(real matrix x, real scalar start)
void           _halton(real matrix x, real scalar start,
                        real scalar hammersley)

real colvector ghalton(real scalar n, real scalar base, real scalar u)
    
```

## Description

`halton( $n$ ,  $d$ )` returns an  $n \times d$  matrix containing a Halton set of length  $n$  and dimension  $d$ .

`halton( $n$ ,  $d$ ,  $start$ )` does the same thing, but the first row of the returned matrix contains the sequences starting at index  $start$ . The default is  $start = 1$ .

`halton( $n$ ,  $d$ ,  $start$ ,  $hammersley$ )`, with  $hammersley \neq 0$ , returns a Hammersley set of length  $n$  and dimension  $d$  with the first row of the returned matrix containing the sequences starting at index  $start$ .

`_halton( $x$ )` modifies the  $n \times d$  matrix  $x$  so that it contains a Halton set of dimension  $d$  of length  $n$ .

`_halton( $x$ ,  $start$ )` does the same thing, but the first row of the returned matrix contains the sequences starting at index  $start$ . The default is  $start = 1$ .

`_halton( $x$ ,  $start$ ,  $hammersley$ )`, with  $hammersley \neq 0$ , returns a Hammersley set of length  $n$  and dimension  $d$  with the first row of the returned matrix containing the sequences starting at index  $start$ .

`ghalton( $n$ ,  $base$ ,  $u$ )` returns an  $n \times 1$  vector containing a (generalized) Halton sequence using base  $base$  and starting from scalar  $0 < u < 1$ , or a nonnegative index  $u = 0, 1, 2, \dots$ .

## Remarks

The Halton sequences are generated from the first  $d$  primes and generally have more uniform coverage over the unit cube of dimension  $d$  than that of sequences generated from pseudouniform random numbers. However, Halton sequences based on large primes ( $d > 10$ ) can be highly correlated, and their coverage can be worse than that of the pseudorandom uniform sequences.

The Hammersley set contains the sequence  $(2 * i - 1) / (2 * n)$ ,  $i = 1, \dots, n$ , in the first dimension and Halton sequences for dimensions 2,  $\dots$ ,  $d$ .

`_halton()` modifies  $x$  and can be used when repeated calls are made to generate long sequences in blocks. Here update the *start* index between calls by using  $start = start + \text{rows}(x)$ .

`ghalton()` uses the base  $base$ , preferably a prime, and generates a Halton sequence using  $0 < u < 1$  or a nonnegative index as the starting value. If  $u$  is uniform  $(0, 1)$ , the sequence is a randomized Halton sequence. If  $u$  is a nonnegative integer, the sequence is the standard Halton sequence starting at index  $u + 1$ .

Conformability

`halton(n, d, start, hammersley):`

<i>input:</i>	<i>n:</i>	$1 \times 1$	
	<i>d:</i>	$1 \times 1$	
	<i>start:</i>	$1 \times 1$	(optional)
	<i>hammersley:</i>	$1 \times 1$	(optional)
<i>output:</i>	<i>result:</i>	$n \times d$	

`_halton(x, start, hammersley):`

<i>input:</i>	<i>x:</i>	$n \times d$	
	<i>start:</i>	$1 \times 1$	(optional)
	<i>hammersley:</i>	$1 \times 1$	(optional)
<i>output:</i>	<i>x:</i>	$n \times d$	

`ghalton(n, base, u):`

<i>input:</i>	<i>n:</i>	$1 \times 1$	
	<i>base:</i>	$1 \times 1$	
	<i>u:</i>	$1 \times 1$	
<i>output:</i>	<i>result:</i>	$n \times 1$	

Diagnostics

The maximum dimension,  $d$ , is 20. The scalar index *start* must be a positive integer, and the scalar  $u$  must be such that  $0 < u < 1$  or a nonnegative integer.

John Henry Halton (1931– ) was born in Brussels, Belgium. He studied mathematics and physics at Cambridge and then at Oxford, where he was a doctoral student of J. M. Hammersley. His career has included positions in government, industry, and academia in both Britain and the United States, latterly as a Professor of Computer Sciences at the University of Wisconsin in Madison and the University of North Carolina in Chapel Hill. Halton's work arises from problems in pure mathematics, theoretical physics, statistics, probability theory, and engineering, and has led to devising and rigorously analyzing algorithms of a combinatorial, probabilistic, and geometric nature. It encompasses tests for statistical relationships between random variables, the hydrodynamic theory of lubrication, probabilistic properties of the Traveling Salesman Problem, and most theoretical aspects of the Monte Carlo method, including the theory and use of both pseudorandom and quasirandom sequences and sets.

John Michael Hammersley (1920–2004) was born in Helensburgh, Dunbartonshire, Scotland. His mathematical studies at Cambridge were interspersed with military service in the Royal Artillery. Hammersley contributed to the use of radar in predicting the flight paths of enemy aircraft and was promoted to the rank of major. After graduation, he moved to Oxford where his research covered several areas of mathematics and statistics. Hammersley is best known for achievements in the study of spatial disorder, especially percolation models, and of stochastic processes generally. He was a pioneer in the use of Monte Carlo methods and was senior author of an important early monograph ([Hammersley and Handscomb 1964](#)). Hammersley contributed to reform of mathematical teaching in Britain, emphasizing the value of solving concrete problems. He was elected a Fellow of the Royal Society.

## References

- Grimmett, G., and D. Welsh. 2007. John Michael Hammersley: 21 March 1920–2 May 2004. *Biographical Memoirs of Fellows of the Royal Society* 53: 163–183.
- Hammersley, J. M., and D. C. Handscomb. 1964. *Monte Carlo Methods*. London: Methuen.

## Also see

[M-4] [mathematical](#) — Important mathematical functions

Syntax

```
real scalar hash1(x [ , n [ , byteorder ] ] )
```

where

*x*: of any type except struct and of any dimension.

*n*: *real scalar*;  $1 \leq n \leq 2,147,483,647$  or . (missing). Optional; default . (missing).

*byteorder*: *real scalar*; 1 (HILO), 2 (LOHI), . (missing, natural byte order). Optional; default . (missing).

Description

`hash1(x)` returns Jenkins’ one-at-a-time hash calculated over the bytes of *x*;  $0 \leq \text{hash1}(x) \leq 4,294,967,295$ .

`hash1(x, n)` returns Jenkins’ one-at-a-time hash scaled to  $1 \leq \text{hash1}(x, n) \leq n$ , assuming  $n < .$  (missing). `hash1(x, .)` is equivalent to `hash1(x)`.

`hash1(x, n, byteorder)` returns `hash1(x, n)` performed on the bytes of *x* ordered as they would be on a HILO computer (*byteorder* = 1), or as they would be on a LOHI computer (*byteorder* = 2), or as they are on this computer (*byteorder* ≥ .). See [M-5] [byteorder\(\)](#) for a definition of byte order.

In all cases, the values returned by `hash1()` are integers.

Remarks

Calculation is significantly faster using the natural byte order of the computer. Argument *byteorder* is included for those rare cases when it is important to calculate the same hash value across different computers, which in the case of `hash1()` is mainly for testing. `hash1()`, being a one-at-a-time method, is not sufficient for constructing digital signatures. It is sufficient for constructing hash tables; see [M-5] [asarray\(\)](#), in which case, byte order is irrelevant. Also note that because strings occur in the same order on all computers, the value of *byteorder* is irrelevant when *x* is a string.

For instance,

```
: hash1("this"), hash1("this",.,1), hash1("this",.,2)
      1              2              3
1 | 2385389520   2385389520   2385389520

: hash1(15), hash1(15,.,1), hash1(15,.,2)
      1              2              3
1 | 463405819    3338064604    463405819
```

The computer on which this example was run is evidently *byteorder* = 2, meaning LOHI, or least-significant byte first.

In a Mata context, it is the two-argument form of `hash1()` that is most useful. In that form, the full result is mapped onto  $[1, n]$ :

$$\text{hash}(x, n) = \text{floor}((\text{hash}(x)/4294967295)*n) + 1$$

For instance,

```
: hash1("this", 10)
6
: hash1(15, 10)
2
```

The result of `hash(x, 10)` could be used directly to index a  $10 \times 1$  array.

## Conformability

```
hash1(x, n, byteorder):
      x:      r × c
      n:      1 × 1      (optional)
byteorder:  1 × 1      (optional)
result:     1 × 1
```

## Diagnostics

None.

Note that `hash1(x[, ...])` never returns a missing result, even if *x* is or contains a missing value. In the missing case, the hash value is calculated of the missing value. Also note that *x* can be a vector or a matrix, in which case the result is calculated over the elements aligned rowwise as if they were a single element. Thus `hash1(("a", "b")) == hash1("ab")`.

## References

- Jenkins, B. 1997. *Dr. Dobbs's Journal*. Algorithm alley: Hash functions. <http://www.ddj.com/184410284>.  
 ——. unknown. A hash function for hash table lookup. <http://www.burtleburtle.net/bob/hash/doobs.html>.

## Also see

- [M-5] [asarray\(\)](#) — Associative arrays  
 [M-4] [programming](#) — Programming functions

Syntax

```
void hessenbergd(numeric matrix A, H, Q)
void _hessenbergd(numeric matrix A, Q)
```

Description

`hessenbergd(A, H, Q)` calculates the Hessenberg decomposition of a square, numeric matrix, *A*, returning the [upper Hessenberg form](#) matrix in *H* and the orthogonal (unitary) matrix in *Q*. *Q* is orthogonal if *A* is real and unitary if *A* is complex.

`_hessenbergd(A, Q)` does the same as `hessenbergd()` except that it returns *H* in *A*.

`_hessenbergd_la()` is the interface into the LAPACK routines used to implement the above function; see [\[M-1\] LAPACK](#). Its direct use is not recommended.

Remarks

The Hessenberg decomposition of a matrix, **A**, can be written as

$$\mathbf{Q}' \times \mathbf{A} \times \mathbf{Q} = \mathbf{H}$$

where **H** is upper Hessenberg; **Q** is orthogonal if **A** is real or unitary if **A** is complex.

A matrix **H** is in upper Hessenberg form if all entries below its first subdiagonal are zero. For example, a 5 × 5 upper Hessenberg matrix looks like

	1	2	3	4	5
1	x	x	x	x	x
2	x	x	x	x	x
3	0	x	x	x	x
4	0	0	x	x	x
5	0	0	0	x	x

For instance,

```
: A
      1      2      3      4      5
1      3      2      1     -2     -5
2      4      2      1      0      3
3      4      4      0      1     -1
4      5      6      7     -2      4
5      6      7      1      2     -1

: hessenbergd(A, H=., Q=.)
```



: H		1	2	3	4	5
1		3	2.903464745	-.552977683	-4.78764119	-1.530555451
2		-9.643650761	7.806451613	2.878001755	5.1085876	5.580422694
3		0	-3.454023879	-6.119229633	-.2347200215	1.467932097
4		0	0	1.404136249	-1.715823624	-.9870601994
5		0	0	0	-2.668128952	-.971398356

: Q		1	2	3	4	5
1		1	0	0	0	0
2		0	-.4147806779	-.0368006164	-.4047768558	-.8140997488
3		0	-.4147806779	-.4871239484	-.5692309155	.5163752637
4		0	-.5184758474	.8096135604	-.0748449196	.2647771074
5		0	-.6221710168	-.3253949238	.7117092805	-.0221645995

Many algorithms use a Hessenberg decomposition in the process of finding another decomposition with more structure.

## Conformability

`hessenbergd(A, H, Q):`

*input:*

*A:*  $n \times n$

*output:*

*H:*  $n \times n$

*Q:*  $n \times n$

`_hessenbergd(A, Q):`

*input:*

*A:*  $n \times n$

*output:*

*A:*  $n \times n$

*Q:*  $n \times n$

## Diagnostics

`_hessenbergd()` aborts with error if *A* is a view.

`hessenbergd()` and `_hessenbergd()` return missing results if *A* contains missing values.

Karl Adolf Hessenberg (1904–1959) was born in Frankfurt am Main, Germany. He was an electrical engineer and gained degrees from the Technische Hochschule Darmstadt. His doctoral dissertation, approved in 1942, was on computation of the eigenvalues and eigensolutions of linear systems of equations. In concurrent work, he introduced what are now called Hessenberg matrices. The mathematician Gerhard Hessenberg was a near relative.

## Also see

[M-1] [LAPACK](#) — The LAPACK linear-algebra routines

[M-5] [schurd\(\)](#) — Schur decomposition

[M-4] [matrix](#) — Matrix functions

# Title

[M-5] Hilbert() — Hilbert matrices

## Syntax

*real matrix*   `Hilbert(real scalar  $n$ )`  
*real matrix*   `invHilbert(real scalar  $n$ )`

## Description

`Hilbert( $n$ )` returns the  $n \times n$  Hilbert matrix, defined as matrix  $H$  with elements  $H_{ij} = 1/(i+j-1)$ .  
`invHilbert( $n$ )` returns the inverse of the  $n \times n$  Hilbert matrix, defined as the matrix with elements  $-1^{i+j} (i+j-1) \times \text{comb}(n+i-1, n-j) \times \text{comb}(n+j-1, n-i) \times \text{comb}(i+j-2, i-1)^2$ .

## Remarks

`Hilbert( $n$ )` and `invHilbert( $n$ )` are used in testing Mata. Hilbert matrices are notoriously ill conditioned. The determinants of the first five Hilbert matrices are 1, 1/12, 1/2,160, 1/6,048,000, and 1/266,716,800,000.

## Conformability

`Hilbert( $n$ )`, `invHilbert( $n$ )`:  
*n*:             $1 \times 1$   
*result*:        $\text{trunc}(n) \times \text{trunc}(n)$

## Diagnostics

None.

David Hilbert (1862–1943) was born near Königsberg, Prussia (now Kaliningrad, Russia), and studied mathematics at the university there. He joined the staff from 1886 to 1895, when he moved to Göttingen, where he stayed despite tempting offers to move. Hilbert was one of the outstanding mathematicians of his time, producing major work in several fields, including invariant theory, algebraic number theory, the foundations of geometry, functional analysis, integral equations, and the calculus of variations. In 1900 he identified 23 key problems in an address to the Second International Congress of Mathematicians in Paris that continues to influence directions in research ([Hilbert 1902](#)). Hilbert’s most noteworthy contribution is the concept of a Hilbert space. His work on what are now known as Hilbert matrices was published in [1894](#).

## References

- Choi, M.-D. 1983. Tricks or treats with the Hilbert matrix. *American Mathematical Monthly* 90: 301–312.
- Hilbert, D. 1894. Ein Beitrag zur Theorie des Legendreschen Polynoms. *Acta Mathematica* 18: 155–159.
- . 1902. Mathematical problems. *Bulletin of the American Mathematical Society* 8: 437–479.
- Reid, C. 1970. *Hilbert*. Berlin: Springer.

## Also see

[\[M-4\]](#) **standard** — Functions to create standard matrices

# Title

[M-5] **I()** — Identity matrix

## Syntax

*real matrix*    **I**(*real scalar n*)

*real matrix*    **I**(*real scalar m*, *real scalar n*)

## Description

**I**(*n*) returns the  $n \times n$  identity matrix.

**I**(*m*, *n*) returns an  $m \times n$  matrix with 1s down its principal diagonal and 0s elsewhere.

## Remarks

**I**() must be typed in uppercase.

## Conformability

**I**(*n*):

<i>n</i> :	$1 \times 1$
<i>result</i> :	$n \times n$

**I**(*m*, *n*):

<i>m</i> :	$1 \times 1$
<i>n</i> :	$1 \times 1$
<i>result</i> :	$m \times n$

## Diagnostics

**I**(*n*) aborts with error if *n* is less than 0 or is missing. *n* is interpreted as **trunc**(*n*).

**I**(*m*, *n*) aborts with error if *m* or *n* are less than 0 or if they are missing. *m* and *n* are interpreted as **trunc**(*m*) and **trunc**(*n*).

## Also see

[M-4] **standard** — Functions to create standard matrices

# Title

[M-5] `inbase()` — Base conversion

## Syntax

*string matrix* `inbase(real scalar base, real matrix  $x$  [, real scalar  $fdigits$  [,  $err$ ]])`  
*real matrix* `frombase(real scalar base, string matrix  $s$ )`

## Description

`inbase(base,  $x$ )` returns a string matrix containing the values of  $x$  in base  $base$ .

`inbase(base,  $x$ ,  $fdigits$ )` does the same;  $fdigits$  specifies the maximum number of digits to the right of the base point to appear in the returned result when  $x$  has a fractional part. `inbase(base,  $x$ )` is equivalent to `inbase(base,  $x$ , 8)`.

`inbase(base,  $x$ ,  $fdigits$ ,  $err$ )` is the same as `inbase(base,  $x$ ,  $fdigits$ )`, except that it returns in  $err$  the difference between  $x$  and the converted result.

$x = \text{frombase}(base, s)$  is the inverse of  $s = \text{inbase}(base, x)$ . It returns base  $base$  number  $s$  as a number. We are tempted to say, “as a number in base 10”, but that is not exactly true. It returns the result as a *real*, that is, as an IEEE base-2 double-precision float that, when you display it, is displayed in base 10.

## Remarks

Remarks are presented under the following headings:

*Positive integers*

*Negative integers*

*Numbers with nonzero fractional parts*

*Use of the functions*

### Positive integers

`inbase(2, 1691)` is 11010011011; that is, 1691 base 10 equals 11010011011 base 2. `frombase(2, "11010011011")` is 1691.

`inbase(3, 1691)` is 2022122; that is, 1691 base 10 equals 2022122 base 3. `frombase(3, "2022122")` is 1691.

`inbase(16, 1691)` is 69b; that is, 1691 base 10 equals 1691 base 16. `frombase(16, "69b")` is 1691. (The base-16 digits are 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, a, b, c, d, e, f.)

`inbase(62, 1691)` is rh; that is, 1691 base 10 equals rh base 62. `frombase(62, "rh")` is 1691. (The base-62 digits are 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, a, b, ..., z, A, B, ..., Z.)

There is a one-to-one correspondence between the integers in different bases. The error of the conversion is always zero.

## Negative integers

Negative integers are no different from positive integers. For instance, `inbase(2, -1691)` is `-11010011011`; that is, `-1691` base 10 equals `-11010011011` base 2. `frombase(2, "-11010011011")` is `-1691`.

The error of the conversion is always zero.

## Numbers with nonzero fractional parts

`inbase(2, 3.5)` is `11.1`; that is, `3.5` base 10 equals `11.1` base 2. `frombase(2, "11.1")` is `3.5`.

`inbase(3, 3.5)` is `10.11111111`.

`inbase(3, 3.5, 20)` is `10.11111111111111111111`.

`inbase(3, 3.5, 30)` is `10.111111111111111111111111111111`.

Therefore, `3.5` base 10 equals `1.1111...` in base 3. There is no exact representation of one-half in base 3. The errors of the above three conversions are `.0000762079`, `1.433399e-10`, and `2.45650e-15`. Those are the values that would be returned in *err* if `inbase(3, 3.5, fdigits, err)` were coded.

`frombase(3, "10.11111111")` is `3.499923792`.

`frombase(3, "10.11111111111111111111")` is `3.4999999998566`.

`frombase(3, "10.111111111111111111111111111111")` is `3.49999999999999734`.

`inbase(16, 3.5)` is `3.8`; that is, `3.5` base 10 equals `3.8` base 16. The error is zero. `frombase(16, "3.8")` is `3.5`.

`inbase(62, 3.5)` is `3.v`; that is, `3.5` base 10 equals `3.v` base 62. `frombase(62, "3.v")` is `3.5`. The error is zero.

In `inbase(base, x, fdigits)`, *fdigits* specifies the maximum number of digits to appear to the right of the base point. *fdigits* is required to be greater than or equal to 1. `inbase(16, 3.5, fdigits)` will be `3.8` regardless of the value of *fdigits* because more digits are unnecessary.

The error that is returned in `inbase(base, x, fdigits, err)` can be an understatement. For instance, `inbase(16, .1, 14, err)` is `0.19999999999999a` and returned in *err* is 0 even though there is no finite-digit representation of `0.1` base 10 in base 16. That is because the `.1` you specified in the call was not actually `0.1` base 10. The computer that you are using is binary, and it converted the `.1` you typed to

`0.000110011001100110011001100110011001100110011001100110011010` base 2

before `inbase()` was ever called. `0.19999999999999a` base 16 is an exact representation of that number.

## Use of the functions

These functions are used mainly for teaching, especially on the sources and avoidance of roundoff error; see [Gould \(2006\)](#).

The functions can have a use in data processing, however, when used with integer arguments. You have a dataset with 10-digit identification numbers. You wish to record the 10-digit number, but more densely. You could convert the number to base 62. The largest 10-digit ID number possible is 9999999999, or aUKYOz base 62. You can record the ID numbers in a six-character string by using `inbase()`. If you needed the original numbers back, you could use `frombase()`.

In a similar way, Stata internally uses base 36 for naming temporary files, and that was important when filenames were limited to eight characters. Base 36 allows Stata to generate up to 2,821,109,907,455 filenames before wrapping of filenames occurs.

## Conformability

inbase(base, x, fdigits, err):  
input:  
base: 1 × 1  
x: r × c  
fdigits: 1 × 1 (optional)  
output:  
err: r × c (optional)  
result: r × c

frombase(base, s):  
base: 1 × 1  
s: r × c  
result: r × c

## Diagnostics

The digits used by `inbase()/frombase()` to encode/decode results are

0 0	10 a	20 k	30 u	40 E	50 0	60 Y
1 1	11 b	21 l	31 v	41 F	51 P	61 Z
2 2	12 c	22 m	32 w	42 G	52 Q	
3 3	13 d	23 n	33 x	43 H	53 R	
4 4	14 e	24 o	34 y	44 I	54 S	
5 5	15 f	25 p	35 z	45 J	55 T	
6 6	16 g	26 q	36 A	46 K	56 U	
7 7	17 h	27 r	37 B	47 L	57 V	
8 8	18 i	28 s	38 C	48 M	58 W	
9 9	19 j	29 t	39 D	49 N	59 X	

When  $base \leq 36$ , `frombase()` treats A, B, C, ..., as if they were a, b, c, ....

`inbase(base, x, fdigits, err)` returns . (missing) if  $base < 2$ ,  $base > 62$ ,  $base$  is not an integer, or  $x$  is missing. If  $fdigits$  is less than 1 or  $fdigits$  is missing, results are as if  $fdigits = 8$  were specified.

`frombase(base, s)` returns . (missing) if  $base < 2$ ,  $base > 62$ ,  $base$  is not an integer, or  $s$  is missing; if  $s$  is not a valid base  $base$  number; or if the converted value of  $s$  is greater than 8.988e+307 in absolute value.



---

## Reference

Gould, W. W. 2006. [Mata Matters: Precision](#). *Stata Journal* 6: 550–560.

## Also see

[M-4] [mathematical](#) — Important mathematical functions

# Title

[M-5] `indexnot()` — Find character not in list

## Syntax

*real matrix*    `indexnot(string matrix  $s_1$ , string matrix  $s_2$ )`

## Description

`indexnot( $s_1$ ,  $s_2$ )` returns the position of the first character of  $s_1$  not found in  $s_2$ , or it returns 0 if all characters of  $s_1$  are found in  $s_2$ .

## Conformability

`indexnot( $s_1$ ,  $s_2$ )`:  
     $s_1$ :         $r_1 \times c_1$   
     $s_2$ :         $r_2 \times c_2$ ,     $s_1$  and  $s_2$  r-conformable  
    *result*:     $\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

`indexnot( $s_1$ ,  $s_2$ )` returns 0 if all characters of  $s_1$  are found in  $s_2$ .

## Also see

[M-4] [string](#) — String manipulation functions

## Syntax

*real vector*   **invorder**(*real vector p*)

*real vector*   **revorder**(*real vector p*)

where  $p$  is assumed to be a [permutation vector](#).

## Description

**invorder**( $p$ ) returns the permutation vector that undoes the permutation performed by  $p$ .

**revorder**( $p$ ) returns the permutation vector that is the reverse of the permutation performed by  $p$ .

## Remarks

See [\[M-1\] permutation](#) for a description of permutation vectors. To summarize,

1. Permutation vectors  $p$  are used to permute the rows or columns of a matrix  $X$ :  $r \times c$ .  
 If  $p$  is intended to permute the rows of  $X$ , the permuted  $X$  is obtained via  $Y = X[p, \cdot]$ .  
 If  $p$  is intended to permute the columns of  $X$ , the permuted  $X$  is obtained via  $Y = X[\cdot, p]$ .
2. If  $p$  is intended to permute the rows of  $X$ , it is called a row-permutation vector. Row-permutation vectors are  $r \times 1$  column vectors.
3. If  $p$  is intended to permute the columns of  $X$ , it is called a column-permutation vector. Column-permutation vectors are  $1 \times c$  row vectors.
4. Row-permutation vectors contain a permutation of the integers 1 to  $r$ .
5. Column-permutation vectors contain a permutation of the integers 1 to  $c$ .

Let us assume that  $p$  is a row-permutation vector, so that

$$Y = X[p, \cdot]$$

**invorder**( $p$ ) returns the row-permutation vector that undoes  $p$ :

$$X = Y[\text{invorder}(p), \cdot]$$

That is, using the matrix notation of [\[M-1\] permutation](#),

$$Y = PX \quad \text{implies} \quad X = P^{-1}Y$$

If  $p$  is the permutation vector corresponding to permutation matrix  $P$ , **invorder**( $p$ ) is the permutation vector corresponding to permutation matrix  $P^{-1}$ .

`revorder(p)` returns the permutation vector that reverses the order of  $p$ . For instance, say that row-permutation vector  $p$  permutes the rows of  $X$  so that the diagonal elements are in ascending order. Then `revorder(p)` would permute the rows of  $X$  so that the diagonal elements would be in descending order.

## Conformability

`invorder(p)`, `revorder(p)`:

$p$ :	$r \times 1$	or	$1 \times c$
<i>result</i> :	$r \times 1$	or	$1 \times c$

## Diagnostics

`invorder(p)` and `revorder(p)` can abort with error or can produce meaningless results when  $p$  is not a permutation vector.

## Also see

[M-1] [permutation](#) — An aside on permutation matrices and vectors

[M-4] [manipulation](#) — Matrix manipulation

# Title

[M-5] **invsym()** — Symmetric real matrix inversion

## Syntax

```
real matrix    invsym(real matrix A)  
real matrix    invsym(real matrix A, real vector order)  
  
void           _invsym(real matrix A)  
void           _invsym(real matrix A, real vector order)
```

## Description

`invsym(A)` returns a generalized inverse of real, symmetric, positive-definite matrix *A*.

`invsym(A, order)` does the same but allows you to specify which columns are to be swept first.

`_invsym(A)` and `_invsym(A, order)` do the same thing as `invsym(A)` and `invsym(A, order)` except that *A* is replaced with the generalized inverse result rather than the result being returned. `_invsym()` uses less memory than `invsym()`.

`invsym()` and `_invsym()` are the routines Stata uses for calculating inverses of symmetric matrices.

Also see [M-5] **luinv()**, [M-5] **qrinv()**, and [M-5] **pinv()** for general matrix inversion.

## Remarks

Remarks are presented under the following headings:

*Definition of generalized inverse*

*Specifying the order in which columns are dropped*

*Determining the rank, or counting the number of dropped columns*

*Extracting linear dependencies*

## Definition of generalized inverse

When the matrix is of full rank and positive definite, the generalized inverse equals the inverse, that is, assuming *A* is  $n \times n$ ,

$$\text{invsym}(A) * A = A * \text{invsym}(A) = I(n)$$

or, at least the above restriction is true up to roundoff error. When *A* is not full rank, the generalized inverse `invsym()` satisfies (ignoring roundoff error)

$$A * \text{invsym}(A) * A = A$$

$$\text{invsym}(A) * A * \text{invsym}(A) = \text{invsym}(A)$$

In the generalized case, there are an infinite number of inverse matrices that can satisfy the above restrictions. The one `invsym()` chooses is one that sets entire columns (and therefore rows) to 0, thus treating *A* as if it were of reduced dimension. Which columns (rows) are selected is determined on the basis of minimizing roundoff error.

In the above we talk as if determining whether a matrix is of full rank is an easy calculation. That is not true. Because of the roundoff error in the manufacturing and recording of  $A$  itself, columns that ought to be perfectly collinear will not be and yet you will still want `invsym()` to behave as if they were. `invsym()` tolerates a little deviation from collinearity in making the perfectly collinear determination.

### Specifying the order in which columns are dropped

Left to make the decision itself, `invsym()` will choose which columns to drop (to set to 0) to minimize the overall roundoff error of the generalized inverse calculation. If column 1 and column 3 are collinear, then `invsym()` will choose to drop column 1 or column 3.

There are occasions, however, when you would like to ensure that a particular column or set of columns are not dropped. Perhaps column 1 corresponds to the intercept of a regression model and you would much rather, if one of columns 1 and 3 has to be dropped, that it be column 3.

Order allows you to specify the columns of the matrix that you would prefer not be dropped in the generalized inverse calculation. In the above example, to prevent column 1 from being dropped, you could code

```
invsym(A, 1)
```

If you would like to keep columns 1, 5, and 10 from being dropped, you can code

```
invsym(A, (1,5,10))
```

Specifying columns not to be dropped does not guarantee that they will not be dropped because they still might be collinear with each other or they might equal constants. However, if any other column can be dropped to satisfy your desire, it will be.

### Determining the rank, or counting the number of dropped columns

If a column is dropped, 0 will appear on the corresponding diagonal entry. Hence, the rank of the original matrix can be extracted after inversion by `invsym()`:

```
: Ainv = invsym(A)
: rank = rows(Ainv)-diag0cnt(Ainv)
```

See [M-5] `diag0cnt()`.

### Extracting linear dependencies

The linear dependencies can be read from the rows of  $A \cdot \text{invsym}(A)$ :

```
: A*invsym(A)
```

	1	2	3
1	1	0	-5.20417e-17
2	-1	0	1
3	1.34441e-16	0	1

The above is interpreted to mean

$$x_1 = x_1$$

$$x_2 = -x_1 + x_3$$

$$x_3 = x_3$$

ignoring roundoff error.

## Conformability

`invsym(A)`, `invsym(A, order)`:

*A*:  $n \times n$   
*order*:  $1 \times k$  or  $k \times 1, k \leq n$  (optional)  
*result*:  $n \times n$

`_invsym(A)`, `_invsym(A, order)`:

*A*:  $n \times n$   
*order*:  $1 \times k$  or  $k \times 1, k \leq n$  (optional)  
*output*:  
*A*:  $n \times n$

## Diagnostics

`invsym(A)`, `invsym(A, order)`, `_invsym(A)`, and `_invsym(A, order)` assume that *A* is symmetric; they do not check. If *A* is nonsymmetric, they treat it as if it were symmetric and equal to its upper triangle.

`invsym()` and `_invsym()` return a result containing missing values if *A* contains missing values.

`_invsym()` aborts with error if *A* is a view. Both functions abort with argument-out-of-range error if *order* is specified and contains values less than 1, greater than `rows(A)`, or the same value more than once.

`invsym()` and `_invsym()` return a matrix of zeros if *A* is not positive definite.

## Also see

[M-5] `cholinv()` — Symmetric, positive-definite matrix inversion

[M-5] `luinv()` — Square matrix inversion

[M-5] `qrinv()` — Generalized inverse of matrix via QR decomposition

[M-5] `pinv()` — Moore–Penrose pseudoinverse

[M-5] `diag0cnt()` — Count zeros on diagonal

[M-4] `matrix` — Matrix functions

[M-4] `solvers` — Functions to solve  $AX=B$  and to obtain *A* inverse

# Title

[M-5] `invtokens()` — Concatenate string rowvector into string scalar

## Syntax

```
string scalar   invtokens(string rowvector s)

string scalar   invtokens(string rowvector s, string scalar c)
```

## Description

`invtokens(s)` returns the elements of *s*, concatenated into a string scalar with the elements separated by spaces. `invtokens(s)` is equivalent to `invtokens(s, " ")`.

`invtokens(s, c)` returns the elements of *s*, concatenated into a string scalar with the elements separated by *c*.

## Remarks

`invtokens(s)` is the inverse of `tokens()` (see [M-5] `tokens()`); `invtokens()` returns the string obtained by concatenating the elements of *s* into a space-separated list.

`invtokens(s, c)` places *c* between the elements of *s* even when the elements of *s* are equal to "". For instance,

```
: s = ("alpha", "", "gamma", "")
: invtokens(s, ";")
alpha;;gamma;
```

To remove separators between empty elements, use `select()` (see [M-5] `select()`) to remove the empty elements from *s* beforehand:

```
: s2 = select(s, strlen(s):>0)
: s2
      1      2
1 | alpha  gamma |
: invtokens(s2, ";")
alpha;gamma
```

## Conformability

```
invtokens(s, c):
  s:      1 × p
  c:      1 × 1   (optional)
result:  1 × 1
```



## Diagnostics

If  $s$  is  $1 \times 0$ , `invtokens( $s,c$ )` returns `""`.

## Also see

[M-5] `tokens()` — Obtain tokens from string

[M-4] `string` — String manipulation functions

# Title

[M-5] **isdiagonal()** — Whether matrix is diagonal

## Syntax

*real scalar*   `isdiagonal(numeric matrix A)`

## Description

`isdiagonal(A)` returns 1 if  $A$  has only zeros off the principal diagonal and returns 0 otherwise. `isdiagonal()` may be used with either real or complex matrices.

## Remarks

See [M-5] **diag()** for making diagonal matrices out of vectors or out of nondiagonal matrices; see [M-5] **diagonal()** for extracting the diagonal of a matrix into a vector.

## Conformability

`isdiagonal(A)`:  
     $A: r \times c$   
    *result*:  $1 \times 1$

## Diagnostics

`isdiagonal(A)` returns 1 if  $A$  is void.

## Also see

[M-5] **diag()** — Create diagonal matrix

[M-5] **diagonal()** — Extract diagonal into column vector

[M-4] **utility** — Matrix utility functions

# Title

[M-5] `isfleeing()` — Whether argument is temporary

## Syntax

*real scalar*   `isfleeing(polymorphic matrix A)`

where  $A$  is an argument passed to your function.

## Description

`isfleeing(A)` returns 1 if  $A$  was constructed for the sole purpose of passing to your function, and returns 0 otherwise. If an argument is fleeting, then you may change its contents and be assured that the caller will not care or even know.

## Remarks

Let us assume that you have written function `myfunc(A)` that takes  $A: r \times c$  and returns an  $r \times c$  matrix. Just to fix ideas, we will pretend that the code for `myfunc()` reads

```
real matrix myfunc(real matrix A)
{
    real scalar      i
    real matrix      B

    B=A
    for (i=1; i<=rows(B); i++) B[i,i] = 1
    return(B)
}
```

Function `myfunc(A)` returns a matrix equal to  $A$ , but with ones along the diagonal. Now let's imagine `myfunc()` in use. A snippet of the code might read

```
...
C = A*myfunc(D)*C
...
```

Here  $D$  is passed to `myfunc()`, and the argument  $D$  is said not to be fleeting. Now consider another code snippet:

```
...
D = A*myfunc(D+E)*D
...
```

In this code snippet, the argument passed to `myfunc()` is  $D+E$  and that argument is fleeting. It is fleeting because it was constructed for the sole purpose of being passed to `myfunc()`, and once `myfunc()` concludes, the matrix containing  $D+E$  will be discarded.

Arguments that are fleeting can be reused to save memory.

Look carefully at the code for `myfunc()`. It makes a copy of the matrix that it was passed. It did that to avoid damaging the matrix that it was passed. Making that copy, however, is unnecessary if the argument received was fleeting, because damaging something that would have been discarded anyway does not matter. Had we not made the copy, we would have saved not only computer time but also memory. Function `myfunc()` could be recoded to read

```
real matrix myfunc(real matrix A)
{
    real scalar    i
    real matrix    B

    if (isfleeting(A)) {
        for (i=1; i<=rows(A); i++) A[i,i] = 1
        return(A)
    }
    B=A
    for (i=1; i<=rows(B); i++) B[i,i] = 1
    return(B)
}
```

Here we wrote separate code for the fleeting and nonfleeting cases. That is not always necessary. We could use a pointer here to combine the two code blocks:

```
real matrix myfunc(real matrix A)
{
    real scalar    i
    real matrix    B
    pointer scalar  p

    if (isfleeting(A)) p = &A
    else {
        B = A
        p = &B
    }
    for (i=1; i<=rows(*p); i++) (*p)[i,i] = 1
    return(*p)
}
```

Many official library functions come in two varieties: `_foo(A, ...)`, which replaces `A` with the calculated result, and `foo(A, ...)`, which returns the result leaving `A` unmodified. Invariably, the code for `foo()` reads

```
function foo(A, ...)
{
    matrix B

    if (isfleeting(A)) {
        _foo(A, ...)
        return(A)
    }
    _foo(B=A, ...)
    return(B)
}
```

This makes function `foo()` whoppingly efficient. If `foo()` is called with a temporary argument—an argument that could be modified without the caller being aware of it—then no extra copy of the matrix is ever made.

## Conformability

```
isfleeing(A):
```

<i>A:</i>	$r \times c$
<i>result:</i>	$1 \times 1$

## Diagnostics

`isfleeing(A)` returns 1 if *A* is fleeting and not a view. The value returned is indeterminate if *A* is not an argument of the function in which it appears, and therefore the value of `isfleeing()` is also indeterminate when used interactively.

## Also see

[\[M-4\] programming](#) — Programming functions

# Title

[M-5] **isreal()** — Storage type of matrix

## Syntax

*real scalar*    **isreal**(*transmorphic matrix X*)

*real scalar*    **iscomplex**(*transmorphic matrix X*)

*real scalar*    **isstring**(*transmorphic matrix X*)

*real scalar*    **ispointer**(*transmorphic matrix X*)

## Description

**isreal**(*X*) returns 1 if *X* is a **real** and returns 0 otherwise.

**iscomplex**(*X*) returns 1 if *X* is a **complex** and returns 0 otherwise.

**isstring**(*X*) returns 1 if *X* is a **string** and returns 0 otherwise.

**ispointer**(*X*) returns 1 if *X* is a **pointer** and returns 0 otherwise.

## Remarks

These functions base their results on storage type. **isreal()** is not the way to check whether a number is real, since it might be stored as a complex and yet still be a real number, such as  $2 + 0i$ . To determine whether *x* is real, you want to use **isrealvalues**(*X*); see [M-5] **isrealvalues()**.

## Conformability

**isreal**(*X*), **iscomplex**(*X*), **isstring**(*X*), **ispointer**(*X*):

<i>X</i> :	$r \times c$
<i>result</i> :	$1 \times 1$

## Diagnostics

These functions return 1 or 0; they cannot fail.

## Also see

[M-5] **isrealvalues()** — Whether matrix contains only real values

[M-5] **eltype()** — Element type and organizational type of object

[M-4] **utility** — Matrix utility functions

# Title

[M-5] **isrealvalues()** — Whether matrix contains only real values

## Syntax

*real scalar*    **isrealvalues**(*numeric matrix X*)

## Description

**isrealvalues**(*X*) returns 1 if *X* is of storage type real or if *X* is of storage type complex and all elements are real or missing; 0 is returned otherwise.

## Remarks

**isrealvalues**(*X*) is logically equivalent to  $X == \text{Re}(X)$  but is significantly faster and requires less memory.

## Conformability

**isrealvalues**(*X*):  
    *X*:         $r \times c$   
    *result*:     $1 \times 1$

## Diagnostics

**isrealvalues**(*X*) returns 1 if *X* is void and complex.

## Also see

[M-5] **isreal()** — Storage type of matrix

[M-4] **utility** — Matrix utility functions

# Title

[M-5] issymmetric() — Whether matrix is symmetric (Hermitian)

## Syntax

*real scalar*    `issymmetric(transmorphic matrix A)`

*real scalar*    `issymmetriconly(transmorphic matrix A)`

## Description

`issymmetric(A)` returns 1 if  $A=A'$  and returns 0 otherwise. (Also see `mreldifsym()` in [M-5] `reldif()`.)

`issymmetriconly(A)` returns 1 if  $A==\text{transposeonly}(A)$  and returns 0 otherwise.

## Remarks

`issymmetric(A)` and `issymmetriconly(A)` return the same result except when  $A$  is complex.

In the complex case, `issymmetric(A)` returns 1 if  $A$  is equal to its conjugate transpose, that is, if  $A$  is Hermitian, which is the complex analog of symmetric.  $A$  is symmetric (Hermitian) if its off-diagonal elements are conjugates of each other and its diagonal elements are real.

`issymmetriconly(A)`, on the other hand, uses the mechanical definition of symmetry:  $A$  is symmetriconly [*sic*] if its off-diagonal elements are equal. `issymmetriconly()` is uninteresting, mathematically speaking, but can be useful in certain data-management programming situations.

## Conformability

`issymmetric(A)`, `issymmetriconly(A)`:

*A:*         $r \times c$   
*result:*     $1 \times 1$

## Diagnostics

`issymmetric(A)` returns 0 if  $A$  is not square. If  $A$  is  $0 \times 0$ , it is symmetric.

`issymmetriconly(A)` returns 0 if  $A$  is not square. If  $A$  is  $0 \times 0$ , it is symmetriconly.

Charles Hermite (1822–1901) was born in Dieuze in eastern France and early showed an aptitude for mathematics, publishing two papers before entering university. He started studying at the Ecole Polytechnique but left after 1 year because of difficulties from a defect in his right foot. However, Hermite was soon appointed to a post at the same institution and later at the Sorbonne. He made outstanding contributions to number theory, algebra, and especially analysis, and he published the first proof that  $e$  is transcendental. Hermite’s name carries on in Hermite polynomials, Hermite’s differential equation, Hermite’s formula of interpolation, and Hermitian matrices.



## Reference

James, I. 2002. *Remarkable Mathematicians: From Euler to von Neumann*. Cambridge: Cambridge University Press.

## Also see

[M-5] **makesymmetric()** — Make square matrix symmetric (Hermitian)

[M-5] **reldif()** — Relative/absolute difference

[M-4] **utility** — Matrix utility functions

# Title

[M-5] **isview()** — Whether matrix is view

## Syntax

*real scalar*    **isview**(*transmorphic matrix X*)

## Description

**isview**(*X*) returns 1 if *X* is a view and 0 otherwise.

See [M-6] **Glossary** for a definition of a **view**.

## Remarks

View matrices are created by **st\_view()**; see [M-5] **st\_view()**.

## Conformability

**isview**(*X*):  
          *X*:         $r \times c$   
          *result*:     $1 \times 1$

## Diagnostics

None.

## Also see

[M-5] **eltype()** — Element type and organizational type of object

[M-4] **utility** — Matrix utility functions

# Title

[M-5] J() — Matrix of constants

## Syntax

*transmorphic matrix* J(*real scalar r*, *real scalar c*, *scalar val*)

*transmorphic matrix* J(*real scalar r*, *real scalar c*, *matrix mat*)

## Description

J(*r*, *c*, *val*) returns an  $r \times c$  matrix with each element equal to *val*.  
J(*r*, *c*, *mat*) returns an  $(r*\text{rows}(\text{mat})) \times (c*\text{cols}(\text{mat}))$  matrix with elements equal to *mat*.  
The first, J(*r*, *c*, *val*), is how J() is commonly used. The first is nothing more than a special case of the second, J(*r*, *c*, *mat*), when *mat* is  $1 \times 1$ .

## Remarks

Remarks are presented under the following headings:

*First syntax: J(r, c, val), val a scalar*  
*Second syntax: J(r, c, mat), mat a matrix*

### First syntax: J(r, c, val), val a scalar

J(*r*, *c*, *val*) creates matrices of constants. For example, J(2, 3, 0) creates

	1	2	3
1	0	0	0
2	0	0	0

J() must be typed in uppercase.  
J() can create any type of matrix:

Function	Returns
J(2, 3, 4)	$2 \times 3$ real matrix, each element = 4
J(2, 3, 4+5i)	$2 \times 3$ complex matrix, each element = $4 + 5i$
J(2, 3, "hi")	$2 \times 3$ string matrix, each element = "hi"
J(2, 3, &x)	$2 \times 3$ pointer matrix, each element = address of x

Also, J() can create void matrices:

J(0, 0, .)	0 × 0 real
J(0, 1, .)	0 × 1 real
J(1, 0, .)	1 × 0 real
J(0, 0, 1i)	0 × 0 complex
J(0, 1, 1i)	0 × 1 complex
J(1, 0, 1i)	1 × 0 complex
J(0, 0, "")	0 × 0 string
J(0, 1, "")	0 × 1 string
J(1, 0, "")	1 × 0 string
J(0, 0, NULL)	0 × 0 pointer
J(0, 1, NULL)	0 × 1 pointer
J(1, 0, NULL)	1 × 0 pointer

When J(*r*, *c*, *val*) is used to create a void matrix, the particular value of the third argument does not matter. Its element type, however, determines the type of matrix produced. Thus, J(0, 0, .), J(0, 0, 1), and J(0, 0, 1/3) all create the same result: a 0 × 0 real matrix. Similarly, J(0, 0, ""), J(0, 0, "name"), and J(0, 0, "?") all create the same result: a 0 × 0 string matrix. See [\[M-2\] void](#) to learn how void matrices are used.

Second syntax: J(*r*, *c*, *mat*), *mat* a matrix

J(*r*, *c*, *mat*) is a generalization of J(*r*, *c*, *val*). When the third argument is a matrix, that matrix is replicated in the result. For instance, if X is (1,2\3,4), then J(2, 3, X) creates

	1	2	3	4	5	6
1	1	2	1	2	1	2
2	3	4	3	4	3	4
3	1	2	1	2	1	2
4	3	4	3	4	3	4

J(*r*, *c*, *val*) is a special case of J(*r*, *c*, *mat*); it just happens that *mat* is 1 × 1.

The matrix created has *r*\*rows(*mat*) rows and *c*\*cols(*mat*) columns.

Note that J(*r*, *c*, *mat*) creates a void matrix if any of *r*, *c*, rows(*mat*), or cols(*mat*) are zero.

## Conformability

$J(r, c, val)$ :

$r$ :  $1 \times 1$   
 $c$ :  $1 \times 1$   
 $val$ :  $1 \times 1$   
 $result$ :  $r \times c$

$J(r, c, mat)$ :

$r$ :  $1 \times 1$   
 $c$ :  $1 \times 1$   
 $mat$ :  $m \times n$   
 $result$ :  $r*m \times c*n$

## Diagnostics

$J(r, c, val)$  and  $J(r, c, mat)$  abort with error if  $r < 0$  or  $c < 0$ , or if  $r \geq .$  or  $c \geq ..$  Arguments  $r$  and  $c$  are interpreted as `trunc(r)` and `trunc(c)`.

## Also see

[M-5] [missingof\(\)](#) — Appropriate missing value

[M-4] [standard](#) — Functions to create standard matrices

# Title

[M-5] **Kmatrix()** — Commutation matrix

## Syntax

*real matrix* **Kmatrix**(*real scalar m*, *real scalar n*)

## Description

**Kmatrix**(*m*, *n*) returns the  $mn \times mn$  commutation matrix **K** for which  $\mathbf{K} * \text{vec}(X) = \text{vec}(X')$ , where  $X$  is an  $m \times n$  matrix.

## Remarks

Commutation matrices are frequently used in computing derivatives of functions of matrices. Section 9.2 of [Lütkepohl \(1996\)](#) lists many useful properties of commutation matrices.

## Conformability

<b>Kmatrix</b> ( <i>m</i> , <i>n</i> ):	
<i>m</i> :	$1 \times 1$
<i>n</i> :	$1 \times 1$
<i>result</i> :	$mn \times mn$

## Diagnostics

**Kmatrix**(*m*, *n*) aborts with error if either *m* or *n* is less than 0 or is missing. *m* and *n* are interpreted as **trunc**(*m*) and **trunc**(*n*).

## Reference

Lütkepohl, H. 1996. *Handbook of Matrices*. New York: Wiley.

## Also see

[M-5] **Dmatrix()** — Duplication matrix

[M-5] **Lmatrix()** — Elimination matrix

[M-5] **vec()** — Stack matrix columns

[M-4] **standard** — Functions to create standard matrices

## Syntax

```
void _flopin(numeric matrix A)
void lapack_function(...)
void _flopout(numeric matrix A)
```

where *lapack\_function* may be

LA_DGBMV()	
LA_DGEBAK()	LA_ZGEBAK()
LA_DGEBAL()	LA_ZGEBAL()
LA_DGEES()	LA_ZGEES()
LA_DGEEV()	LA_ZGEEV()
LA_DGEHRD()	LA_ZGEHRD()
LA_DGGBAK()	LA_ZGGBAK()
LA_DGGBAL()	LA_ZGGBAL()
LA_DGGHRD()	LA_ZGGHRD()
LA_DHGEQZ()	LA_ZHGEQZ()
LA_DHSEIN()	LA_ZHSEIN()
LA_DHSEQR()	LA_ZHSEQR()
LA_DLAMCH()	
LA_DORGHR()	
LA_DSYEVX()	
LA_DTGSSEN()	LA_ZTGSSEN()
LA_DTGEVC()	LA_ZTGEVC()
LA_DTREVC()	LA_ZTREVC()
LA_DTRSEN()	LA_ZTRSEN()
	LA_ZUNGHR()

## Description

LA\_DGBMV(), LA\_DGEBAK(), LA\_ZGEBAK(), LA\_DGEBAL(), LA\_ZGEBAL(), ... are LAPACK functions in original, as-is form; see [\[M-1\] LAPACK](#). These functions form the basis for many of Mata's linear-algebra capabilities. Mata functions such as [cholesky\(\)](#), [svd\(\)](#), and [eigensystem\(\)](#) are implemented using these functions; see [\[M-4\] matrix](#). Those functions are easier to use. The LA\_\*( ) functions provide more capability.

\_flop<sub>in</sub>() and \_flop<sub>out</sub>() convert matrices to and from the form required by the LA\_\*( ) functions.

## Remarks

LAPACK stands for Linear Algebra PACKage and is a freely available set of Fortran 90 routines for solving systems of simultaneous equations, eigenvalue problems, and singular-value problems. The original Fortran routines have six-letter names like DGEHRD, DORGHR, and so on. The Mata functions `LA_DGEHRD()`, `LA_DORGHR()`, etc., are a subset of the LAPACK double-precision real and complex routine. All LAPACK double-precision functions will eventually be made available.

Documentation for the LAPACK routines can be found at <http://www.netlib.org/lapack/>, although we recommend obtaining *LAPACK Users' Guide* by [Anderson et al. \(1999\)](#).

Remarks are presented under the following headings:

*Mapping calling sequence from Fortran to Mata*

*Flopping: Preparing matrices for LAPACK*

*Warning on the use of `rows()` and `cols()` after `_flopint()`*

*Warning: It is your responsibility to check info*

*Example*

## Mapping calling sequence from Fortran to Mata

LAPACK functions are named with first letter S, D, C, or Z. S means single-precision real, D means double-precision real, C means single-precision complex, and Z means double-precision complex. Mata provides the D\* and Z\* functions. The LAPACK documentation is in terms of S\* and C\*. Thus, to find the documentation for `LA_DGEHRD`, you must look up `SGEHRD` in the original documentation.

The documentation (Anderson et al. 1999, 227) reads, in part,

```
SUBROUTINE SGEHRD(N, ILO, IHI, A, LDA, TAU, WORK, LWORK, INFO)
INTEGER  IHI, ILO, INFO, LDA, LWORK, N
REAL     A(LDA, *), TAU(*), WORK(LWORK)
```

and the documentation states that `SGEHRD` reduces a real, general matrix, **A**, to upper Hessenberg form, **H**, by an orthogonal similarity transformation:  $\mathbf{Q}' \times \mathbf{A} \times \mathbf{Q} = \mathbf{H}$ .

The corresponding Mata function, `LA_DGEHRD()`, has the same arguments. In Mata, arguments `ihi`, `ilo`, `info`, `lda`, `lwork`, and `n` are *real scalars*. Argument **A** is a *real matrix*, and arguments `tau` and `work` are *real vectors*.

You can read the rest of the original documentation to find out what is to be placed (or returned) in each argument. It turns out that **A** is assumed to be dimensioned `LDA × something` and that the routine works on **A**(1,1) (using Fortran notation) through **A**(*N*,*N*). The routine also needs work space, which you are to supply in vector **WORK**. In the standard LAPACK way, LAPACK offers you a choice: you can preallocate **WORK**, in which case you have to choose a fairly large dimension for it, or you can do a query to find out how large the dimension needs to be for this particular problem. If you preallocate, the documentation reveals that the **WORK** must be of size *N*, and you set `LWORK` equal to *N*. If you wish to query, then you make **WORK** of size 1 and set `LWORK` equal to `-1`. The LAPACK routine will then return in the first element of **WORK** the optimal size. Then you call the function again with **WORK** allocated to be the optimal size and `LWORK` set to equal the optimal size.

Concerning Mata, the above works. You can follow the LAPACK documentation to the letter. Use `J()` to allocate matrices or vectors. Alternatively, you can specify all sizes as missing value (`.`), and Mata will fill in the appropriate value based on the assumption that you are using the entire matrix.



Thus, in `LA_DGEHRD()`, you could specify `lda` as missing, and the function would run as if you had specified `lda` equal to `cols(A)`. You could specify `n` as missing, and the function would run as if you had specified `n` as `rows(A)`.

Work areas, however, are treated differently. You can follow the standard LAPACK convention outlined above; or you can specify the sizes of work areas (`lwork`) and specify the work areas themselves (`work`) as missing values, and Mata will allocate the work areas for you. The allocation will be as you specified.

One feature provided by some LAPACK functions is not supported by the Mata implementation. If a function allows a function pointer, you may not avail yourself of that option.

## Flopping: Preparing matrices for LAPACK

The LAPACK functions provided in Mata are the original LAPACK functions. Mata, which is C based, stores matrices rowwise. LAPACK, which is Fortran based, stores matrices columnwise. Mata and Fortran also disagree on how complex matrices are to be organized.

Functions `_flopin()` and `_flopout()` handle these issues. Coding `_flopin(A)` changes matrix `A` from the Mata convention to the LAPACK convention. Coding `_flopout(A)` changes `A` from the LAPACK convention to the Mata convention.

The `LA_*`() functions do not do this for you because LAPACK often takes two or three LAPACK functions run in sequence to achieve the desired result, and it would be a waste of computer time to switch conventions between calls.

## Warning on the use of `rows()` and `cols()` after `_flopin()`

Be careful using the `rows()` and `cols()` functions. `rows()` of a flopped matrix returns the logical number of columns and `cols()` of a flopped matrix returns the logical number of rows!

The danger of confusion is especially great when using `J()` to allocate work areas. If a LAPACK function requires a work area of  $r \times c$ , your code,

```
_LA_function(..., J(c, r, .), ...)
```

## Warning: It is your responsibility to check info

The LAPACK functions do not abort with error on failure. They instead store 0 in `info` (usually the last argument) if successful and store an error code if not successful. The error code is usually negative and indicates the argument that is a problem.

## Example

The following example uses the LAPACK function `DGEHRD` to obtain the Hessenberg form of matrix `A`. We will begin with

	1	2	3	4
1	1	2	3	4
2	4	5	6	7
3	7	8	9	10
4	8	9	10	11

The first step is to use `_flopin()` to put **A** in LAPACK order:

```
: _flopin(A)
```

Next we make a work-space query to get the optimal size of the work area.

```
: LA_DGEHRD(., 1, 4, A, ., tau=., work=., lwork=-1, info=0)
: lwork = work[1,1]
: lwork
128
```

After putting the work-space size in `lwork`, we can call `LA_DGEHRD()` again to perform the Hessenberg decomposition:

```
: LA_DGEHRD(., 1, 4, A, ., tau=., work=., lwork, info=0)
```

LAPACK function `DGEHRD` saves the result in the upper triangle and the first subdiagonal of **A**. We must use `_flopout()` to change that back to Mata order, and finally, we extract the result:

```
: _flopout(A)
: A = A-sublowertriangle(A, 2)
: A
```

	1	2	3	4
1	1	-5.370750529	.0345341258	.3922322703
2	-11.35781669	25.18604651	-4.40577178	-.6561483899
3	0	-1.660145888	-.1860465116	.1760901813
4	0	0	-8.32667e-16	-5.27356e-16

## Reference

Anderson, E., Z. Bai, C. Bischof, S. Blackford, J. Demmel, J. J. Dongarra, J. Du Croz, A. Greenbaum, S. Hammarling, A. McKenney, and D. Sorensen. 1999. *LAPACK Users' Guide*. 3rd ed. Philadelphia: Society for Industrial and Applied Mathematics.

## Also see

- [M-1] [LAPACK](#) — The LAPACK linear-algebra routines
- [R] [copyright lapack](#) — LAPACK copyright notification
- [M-4] [matrix](#) — Matrix functions

# Title

[M-5] **liststruct()** — List structure’s contents

## Syntax

*void* **liststruct**(*struct whatever matrix x*)

## Description

**liststruct()** lists *x*’s contents, where *x* is an instance of structure *whatever*.

## Remarks

**liststruct()** is often useful in debugging.

The dimension and type of all elements are listed, and the values of scalars are also shown.

## Conformability

**liststruct**(*x*):  
          *x*:           $r \times c$   
          *result*:      *void*

## Diagnostics

None.

## Also see

[\[M-2\] \*\*struct\*\*](#) — Structures

[\[M-4\] \*\*io\*\*](#) — I/O functions

# Title

[M-5] **Lmatrix()** — Elimination matrix

## Syntax

*real matrix* **Lmatrix**(*real scalar n*)

## Description

**Lmatrix**( $n$ ) returns the  $n(n+1)/2 \times n^2$  elimination matrix  $L$  for which  $L \cdot \text{vec}(X) = \text{vech}(X)$ , where  $X$  is an  $n \times n$  symmetric matrix.

## Remarks

Elimination matrices are frequently used in computing derivatives of functions of symmetric matrices. Section 9.6 of [Lütkepohl \(1996\)](#) lists many useful properties of elimination matrices.

## Conformability

<b>Lmatrix</b> ( $n$ ):	
$n$ :	$1 \times 1$
$result$ :	$n(n+1)/2 \times n^2$

## Diagnostics

**Lmatrix**( $n$ ) aborts with error if  $n$  is less than 0 or is missing.  $n$  is interpreted as **trunc**( $n$ ).

## Reference

Lütkepohl, H. 1996. *Handbook of Matrices*. New York: Wiley.

## Also see

[M-5] **Dmatrix()** — Duplication matrix

[M-5] **Kmatrix()** — Commutation matrix

[M-5] **vec()** — Stack matrix columns

[M-4] **standard** — Functions to create standard matrices

# Title

[M-5] **logit()** — Log odds and complementary log-log

## Syntax

*real matrix*   `logit(real matrix X)`

*real matrix*   `invlogit(real matrix X)`

*real matrix*   `cloglog(real matrix X)`

*real matrix*   `invcloglog(real matrix X)`

## Description

`logit(X)` returns the log of the odds ratio of the elements of  $X$ ,  $\ln\{x/(1-x)\}$ .

`invlogit(X)` returns the inverse of the `logit()` of the elements of  $X$ ,  $\exp(x)/\{1+\exp(x)\}$ .

`cloglog(X)` returns the complementary log-log of the elements of  $X$ ,  $\ln\{-\ln(1-x)\}$ .

`invcloglog(X)` returns the elementwise inverse of `cloglog()` of the elements of  $X$ ,  $1 - \exp\{-\exp(x)\}$ .

## Conformability

All functions return a matrix of the same dimension as input containing element-by-element calculated results.

## Diagnostics

`logit(X)` and `cloglog(X)` return missing when  $x \leq 0$  or  $x \geq 1$ .

## Also see

[\[M-4\] statistical](#) — Statistical functions

## Syntax

*numeric matrix*    `lowertriangle(numeric matrix A [ , numeric scalar d ])`

*numeric matrix*    `uppertriangle(numeric matrix A [ , numeric scalar d ])`

*void*                `_lowertriangle(numeric matrix A [ , numeric scalar d ])`

*void*                `_uppertriangle(numeric matrix A [ , numeric scalar d ])`

where argument *d* is optional.

## Description

`lowertriangle()` returns the lower triangle of *A*.

`uppertriangle()` returns the upper triangle of *A*.

`_lowertriangle()` replaces *A* with its lower triangle.

`_uppertriangle()` replaces *A* with its upper triangle.

## Remarks

Remarks are presented under the following headings:

*Optional argument d*

*Nonsquare matrices*

### Optional argument d

Optional argument *d* specifies the treatment of the diagonal. Specifying *d* >= ., or not specifying *d* at all, means no special treatment; if

$$A = \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}$$

then

$$\text{lowertriangle}(A, .) = \begin{bmatrix} 1 & 0 & 0 \\ 4 & 5 & 0 \\ 7 & 8 & 9 \end{bmatrix}$$

If a nonmissing value is specified for  $d$ , however, that value is substituted for each element of the diagonal, for example,

$$\text{lowertriangle}(A, 1) = \begin{bmatrix} 1 & 0 & 0 \\ 4 & 1 & 0 \\ 7 & 8 & 1 \end{bmatrix}$$

## Nonsquare matrices

`lowertriangle()` and `uppertriangle()` may be used with nonsquare matrices. If

$$A = \begin{bmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{bmatrix}$$

then

$$\text{lowertriangle}(A) = \begin{bmatrix} 1 & 0 & 0 \\ 5 & 6 & 0 \\ 9 & 10 & 11 \end{bmatrix}$$

and

$$\text{uppertriangle}(A) = \begin{bmatrix} 1 & 2 & 3 & 4 \\ 0 & 6 & 7 & 8 \\ 0 & 0 & 11 & 12 \end{bmatrix}$$

`_lowertriangle()` and `_uppertriangle()`, however, may not be used with nonsquare matrices.

## Conformability

`lowertriangle(A, d):`

$A:$   $r \times c$   
 $d:$   $1 \times 1$  (optional)  
 $result:$   $r \times \min(r, c)$

`uppertriangle(A, d):`

$A:$   $r \times c$   
 $d:$   $1 \times 1$  (optional)  
 $result:$   $\min(r, c) \times c$

`_lowertriangle(A, d), _uppertriangle(A, d):`

*input:*

$A:$   $n \times n$   
 $d:$   $1 \times 1$  (optional)

*output:*

$A:$   $n \times n$

## Diagnostics

None.

## Also see

[M-4] [manipulation](#) — Matrix manipulation



# Title

[M-5] **lud()** — LU decomposition

## Syntax

*void* `lud(numeric matrix A, L, U, p)`

*void* `_lud(numeric matrix L, U, p)`

*void* `_lud_la(numeric matrix A, q)`

where

1.  $A$  may be real or complex and need not be square.
2. The types of  $L$ ,  $U$ ,  $p$ , and  $q$  are irrelevant; results are returned there.

## Description

`lud(A, L, U, p)` returns the LU decomposition (with partial pivoting) of  $A$  in  $L$  and  $U$  along with a permutation vector  $p$ . The returned results are such that  $A=L[p, .]*U$  up to roundoff error.

`_lud(L, U, p)` is similar to `lud()`, except that it conserves memory. The matrix to be decomposed is passed in  $L$ , and the same storage location is overwritten with the calculated  $L$  matrix.

`_lud_la(A, q)` is the [M-1] **LAPACK** routine that the above functions use to calculate the LU decomposition. See [LAPACK routine](#) below.

## Remarks

Remarks are presented under the following headings:

[LU decomposition](#)  
[LAPACK routine](#)

## LU decomposition

The LU decomposition of matrix  $A$  can be written

$$P'A = LU$$

where  $P'$  is a [permutation matrix](#) that permutes the rows of  $A$ .  $L$  is lower triangular and  $U$  is upper triangular. The decomposition can also be written

$$A = PLU$$

because, given that  $P$  is a permutation matrix,  $P^{-1} = P'$ .

Rather than returning  $P$  directly, returned is  $p$  corresponding to  $P$ . Lowercase  $p$  is a column vector that contains the subscripts of the rows in the desired order. That is,

$$PL = L[p, .]$$

The advantage of this is that  $p$  requires less memory than  $P$  and the reorganization, should it be desired, can be performed more quickly; see [M-1] **permutation**. In any case, the formula defining the LU decomposition can be written

$$A = L[p, .] * U$$

One can also write

$$B = LU, \text{ where } B[p, .] = A$$

## LAPACK routine

`_lud_la(A, q)` is the interface into the [M-1] **LAPACK** routines that the above functions use to calculate the LU decomposition. You may use it directly if you wish.

Matrix  $A$  is decomposed, and the decomposition is placed back in  $A$ .  $U$  is stored in the upper triangle (including the diagonal) of  $A$ .  $L$  is stored in the lower triangle of  $A$ , and it is understood that  $L$  is supposed to have ones on its diagonal.  $q$  is a column vector recording the row swaps that account for the pivoting. This is the same information as stored in  $p$ , but in a different format.

$q$  records the row swaps to be made. For instance,  $q = (1\ 2\ 2)$  means that (start at the end) the third row is to be swapped with the second row, then the second row is to stay where it is, and finally the first row is to stay where it is.  $q$  can be converted into  $p$  by the following logic:

```
p = 1::rows(q)
for (i=rows(q); i>=1; i--) {
  hold = p[i]
  p[i] = p[q[i]]
  p[q[i]] = hold
}
```

## Conformability

`lud(A, L, U, p):`

*input:*

$A:$        $r \times c$

*output:*

$L:$        $r \times m, \quad m = \min(r, c)$

$U:$        $m \times c$

$p:$        $r \times 1$

`_lud(L, U, p):`

*input:*

$L: \quad r \times c$

*output:*

$L: \quad r \times m, \quad m = \min(r, c)$

$U: \quad m \times c$

$p: \quad r \times 1$

`_lud_la(A, q):`

*input:*

$A: \quad r \times c$

*output:*

$A: \quad r \times c$

$q: \quad r \times 1$

## Diagnostics

`lud(A, L, U, p)` returns missing results if  $A$  contains missing values;  $L$  will have missing values below the diagonal, 1s on the diagonal, and 0s above the diagonal;  $U$  will have missing values on and above the diagonal and 0s below. Thus if there are missing values,  $U[1, 1]$  will contain missing.

`_lud(L, U, p)` sets  $L$  and  $U$  as described above if  $A$  contains missing values.

`_lud_la(A, q)` aborts with error if  $A$  is a view.

## Also see

[M-5] `det()` — Determinant of matrix

[M-5] `lusolve()` — Solve  $AX=B$  for  $X$  using LU decomposition

[M-4] `matrix` — Matrix functions

# Title

[M-5] **luinv()** — Square matrix inversion

## Syntax

```
numeric matrix    luinv(numeric matrix A)
numeric matrix    luinv(numeric matrix A, real scalar tol)

void              _luinv(numeric matrix A)
void              _luinv(numeric matrix A, real scalar tol)

real scalar       _luinv_la(numeric matrix A, b)
```

## Description

`luinv(A)` and `luinv(A, tol)` return the inverse of real or complex, square matrix *A*.

`_luinv(A)` and `_luinv(A, tol)` do the same thing except that, rather than returning the inverse matrix, they overwrite the original matrix *A* with the inverse.

In all cases, optional argument *tol* specifies the tolerance for determining singularity; see *Remarks* below.

`_luinv_la(A, b)` is the interface to the [M-1] **LAPACK** routines that do the work. The output *b* is a real scalar, which is 1 if the LAPACK routine used a blocked algorithm and 0 otherwise.

## Remarks

These routines calculate the inverse of *A*. The inverse matrix  $A^{-1}$  of *A* satisfies the conditions

$$AA^{-1} = I$$

$$A^{-1}A = I$$

*A* is required to be square and of full rank. See [M-5] **qrinv()** and [M-5] **pinv()** for generalized inverses of nonsquare or rank-deficient matrices. See [M-5] **invsym()** for inversion of real, symmetric matrices.

`luinv(A)` is logically equivalent to `lusolve(A, I(rows(A)))`; see [M-5] **lusolve()** for details and for use of the optional *tol* argument.

## Conformability

```
luinv(A, tol):
    A:      n × n
    tol:    1 × 1   (optional)
    result: n × n
```

`_luinv(A, tol):`

*input:*

$A:$       $n \times n$   
 $tol:$      $1 \times 1$     (optional)

*output:*

$A:$       $n \times n$

`_luinv_la(A, b):`

*input:*

$A:$       $n \times n$

*output:*

$A:$       $n \times n$

$b:$       $1 \times 1$

*result:*      $1 \times 1$

## Diagnostics

The inverse returned by these functions is real if  $A$  is real and is complex if  $A$  is complex. If you use these functions with a singular matrix, returned will be a matrix of missing values. The determination of singularity is made relative to *tol*. See [Tolerance](#) under *Remarks* in [M-5] [lusolve\(\)](#) for details.

`luinv(A)` and `_luinv(A)` return a matrix containing missing if  $A$  contains missing values.

`_luinv(A)` aborts with error if  $A$  is a view.

`_luinv_la(A, b)` should not be used directly; use `_luinv()`.

See [M-5] [lusolve\(\)](#) and [M-1] [tolerance](#) for information on the optional *tol* argument.

## Also see

[M-5] [invsym\(\)](#) — Symmetric real matrix inversion

[M-5] [cholinv\(\)](#) — Symmetric, positive-definite matrix inversion

[M-5] [qrinv\(\)](#) — Generalized inverse of matrix via QR decomposition

[M-5] [pinv\(\)](#) — Moore–Penrose pseudoinverse

[M-5] [lusolve\(\)](#) — Solve  $AX=B$  for  $X$  using LU decomposition

[M-5] [lud\(\)](#) — LU decomposition

[M-4] [matrix](#) — Matrix functions

[M-4] [solvers](#) — Functions to solve  $AX=B$  and to obtain  $A$  inverse

## Syntax

<i>numeric matrix</i>	<code>lusolve(numeric matrix A, numeric matrix B)</code>
<i>numeric matrix</i>	<code>lusolve(numeric matrix A, numeric matrix B, real scalar tol)</code>
<i>void</i>	<code>_lusolve(numeric matrix A, numeric matrix B)</code>
<i>void</i>	<code>_lusolve(numeric matrix A, numeric matrix B, real scalar tol)</code>
<i>real scalar</i>	<code>_lusolve_la(numeric matrix A, numeric matrix B)</code>
<i>real scalar</i>	<code>_lusolve_la(numeric matrix A, numeric matrix B, real scalar tol)</code>

## Description

`lusolve(A, B)` solves  $AX=B$  and returns  $X$ . `lusolve()` returns a matrix of missing values if  $A$  is singular.

`lusolve(A, B, tol)` does the same thing but allows you to specify the tolerance for declaring that  $A$  is singular; see [Tolerance](#) under *Remarks* below.

`_lusolve(A, B)` and `_lusolve(A, B, tol)` do the same thing except that, rather than returning the solution  $X$ , they overwrite  $B$  with the solution and, in the process of making the calculation, they destroy the contents of  $A$ .

`_lusolve_la(A, B)` and `_lusolve_la(A, B, tol)` are the interfaces to the [\[M-1\] LAPACK](#) routines that do the work. They solve  $AX=B$  for  $X$ , returning the solution in  $B$  and, in the process, using as workspace (overwriting)  $A$ . The routines return 1 if  $A$  was singular and 0 otherwise. If  $A$  was singular,  $B$  is overwritten with a matrix of missing values.

## Remarks

The above functions solve  $AX=B$  via LU decomposition and are accurate. An alternative is `qrsolve()` (see [\[M-5\] qrsolve\(\)](#)), which uses QR decomposition. The difference between the two solutions is not, practically speaking, accuracy. When  $A$  is of full rank, both routines return equivalent results, and the LU approach is quicker, using approximately  $O(2/3n^3)$  operations rather than  $O(4/3n^3)$ , where  $A$  is  $n \times n$ .

The difference arises when  $A$  is singular. Then the LU-based routines documented here return missing values. The QR-based routines documented in [\[M-5\] qrsolve\(\)](#) return a generalized (least squares) solution.

For more information on LU and QR decomposition, see [\[M-5\] lud\(\)](#) and see [\[M-5\] qrd\(\)](#).

Remarks are presented under the following headings:

[Derivation](#)  
[Relationship to inversion](#)  
[Tolerance](#)

## Derivation

We wish to solve for  $X$

$$AX = B \quad (1)$$

Perform LU decomposition on  $A$  so that we have  $A = PLU$ . Then (1) can be written

$$PLUX = B$$

or, premultiplying by  $P'$  and remembering that  $P'P = I$ ,

$$LUX = P'B \quad (2)$$

Define

$$Z = UX \quad (3)$$

Then (2) can be rewritten

$$LZ = P'B \quad (4)$$

It is easy to solve (4) for  $Z$  because  $L$  is a lower-triangular matrix. Once  $Z$  is known, it is easy to solve (3) for  $X$  because  $U$  is upper triangular.

## Relationship to inversion

Another way to solve

$$AX = B$$

is to obtain  $A^{-1}$  and then calculate

$$X = A^{-1}B$$

It is, however, better to solve  $AX = B$  directly because fewer numerical operations are required, and the result is therefore more accurate and obtained in less computer time.

Indeed, rather than thinking about how solving a system of equations can be implemented via inversion, it is more productive to think about how inversion can be implemented via solving a system of equations. Obtaining  $A^{-1}$  amounts to solving

$$AX = I$$

Thus `lusolve()` (or any other solve routine) can be used to obtain inverses. The inverse of  $A$  can be obtained by coding

```
: Ainv = lusolve(A, I(rows(A)))
```

In fact, we provide `luinv()` (see [M-5] [luinv\(\)](#)) for obtaining inverses via LU decomposition, but `luinv()` amounts to making the above calculation, although a little memory is saved because the matrix  $I$  is never constructed.

Hence, everything said about `lusolve()` applies equally to `luinv()`.

# Tolerance

The default tolerance used is

$$\eta = (1\text{e-}13) * \text{trace}(\text{abs}(U)) / n$$

where  $U$  is the upper-triangular matrix of the LU decomposition of  $A$ :  $n \times n$ .  $A$  is declared to be singular if any diagonal element of  $U$  is less than or equal to  $\eta$ .

If you specify  $\text{tol} > 0$ , the value you specify is used to multiply  $\eta$ . You may instead specify  $\text{tol} \leq 0$ , and then the negative of the value you specify is used in place of  $\eta$ ; see [M-1] [tolerance](#).

So why not specify  $\text{tol} = 0$ ? You do not want to do that because, as matrices become close to being singular, results can become inaccurate. Here is an example:

```

: rseed(12345)
: A = lowertriangle(runiform(4,4))
: A[3,3] = 1e-15
: trux = runiform(4,1)
: b = A*trux
: /* the above created an Ax=b problem, and we have placed the true
> value of x in trux. We now obtain the solution via lusolve()
> and compare trux with the value obtained:
> */
: x = lusolve(A, b, 0)
: trux, x

```

1	.7997150919	.7997150919
2	.9102488109	.9102488109
3	.442547889	.3593109488
4	.756650276	.8337468202

← The discussed numerical instability can cause this output to vary a little across different computers

We would like to see the second column being nearly equal to the first—the estimated  $x$  being nearly equal to the true  $x$ —but there are substantial differences.

Even though the difference between  $x$  and  $x_{\text{true}}$  is substantial, the difference between them is small in the prediction space:

```

: A*trux-b, A*x-b

```

	1	2
1	0	-2.77556e-17
2	0	0
3	0	0
4	0	0

What made this problem so difficult was the line  $A[3,3] = 1\text{e-}15$ . Remove that and you would find that the maximum difference between  $x$  and  $\text{trux}$  would be  $2.22045\text{e-}16$ .

The degree to which the residuals  $A*x-b$  are a reliable measure of the accuracy of  $x$  depends on the condition number of the matrix, which can be obtained by [M-5] [cond\(\)](#), which for  $A$ , is  $3.23984\text{e+}15$ . If the matrix is well conditioned, small residuals imply an accurate solution for  $x$ . If the matrix is ill conditioned, small residuals are not a reliable indicator of accuracy.

Another way to check the accuracy of  $x$  is to set  $\text{tol} = 0$  and to see how well  $x$  could be obtained were  $x = x$ :



```
: x = lusolve(A, b, 0)
: x2 = lusolve(A, A*x, 0)
```

If  $x$  and  $x2$  are virtually the same, then you can safely assume that  $x$  is the result of a numerically accurate calculation. You might compare  $x$  and  $x2$  with `mreldif(x2,x)`; see [M-5] `reldif()`. In our example, `mreldif(x2,x)` is .03, a large difference.

If  $A$  is ill conditioned, then small changes in  $A$  or  $B$  can lead to radical differences in the solution for  $X$ .

## Conformability

`lusolve(A, B, tol)`:

*input:*

$A:$       $n \times n$   
 $B:$       $n \times k$   
 $tol:$      $1 \times 1$  (optional)

*output:*

*result:*      $n \times k$

`_lusolve(A, B, tol)`:

*input:*

$A:$       $n \times n$   
 $B:$       $n \times k$   
 $tol:$      $1 \times 1$  (optional)

*output:*

$A:$       $0 \times 0$   
 $B:$       $n \times k$

`_lusolve_la(A, B, tol)`:

*input:*

$A:$       $n \times n$   
 $B:$       $n \times k$   
 $tol:$      $1 \times 1$  (optional)

*output:*

$A:$       $0 \times 0$   
 $B:$       $n \times k$   
*result:*      $1 \times 1$

## Diagnostics

`lusolve(A, B, ...)`, `_lusolve(A, B, ...)`, and `_lusolve_la(A, B, ...)` return a result containing missing if  $A$  or  $B$  contain missing values. The functions return a result containing all missing values if  $A$  is singular.

`_lusolve(A, B, ...)` and `_lusolve_la(A, B, ...)` abort with error if  $A$  or  $B$  is a view.

`_lusolve_la(A, B, ...)` should not be used directly; use `_lusolve()`.

## Also see

[M-5] `luinv()` — Square matrix inversion

[M-5] `lud()` — LU decomposition

[M-5] `solverlower()` — Solve  $AX=B$  for  $X$ ,  $A$  triangular

[M-5] `cholsolve()` — Solve  $AX=B$  for  $X$  using Cholesky decomposition

[M-5] `qrsolve()` — Solve  $AX=B$  for  $X$  using QR decomposition

[M-5] `svsolve()` — Solve  $AX=B$  for  $X$  using singular value decomposition

[M-4] `matrix` — Matrix functions

[M-4] `solvers` — Functions to solve  $AX=B$  and to obtain  $A$  inverse

# Title

[M-5] **makesymmetric()** — Make square matrix symmetric (Hermitian)

## Syntax

*numeric matrix*    `makesymmetric(numeric matrix A)`

*void*                `_makesymmetric(numeric matrix A)`

## Description

`makesymmetric(A)` returns  $A$  made into a symmetric (Hermitian) matrix by reflecting elements below the diagonal.

`_makesymmetric(A)` does the same thing but stores the result back in  $A$ .

## Remarks

If  $A$  is real, elements below the diagonal are copied into their corresponding above-the-diagonal position.

If  $A$  is complex, the conjugate of the elements below the diagonal are copied into their corresponding above-the-diagonal positions, and the imaginary part of the diagonal is set to zero.

Whether  $A$  is real or complex, roundoff error can make matrix calculations that are supposed to produce symmetric matrices produce matrices that vary a little from symmetry, and `makesymmetric()` can be used to correct the situation.

## Conformability

`makesymmetric(A):`

*A:*             $n \times n$   
*result:*        $n \times n$

`_makesymmetric(A):`

*A:*             $n \times n$

## Diagnostics

`makesymmetric(A)` and `_makesymmetric(A)` abort with error if  $A$  is not square. Also, `_makesymmetric()` aborts with error if  $A$  is a view.

## Also see

[M-5] [issymmetric\(\)](#) — Whether matrix is symmetric (Hermitian)

[M-4] [manipulation](#) — Matrix manipulation

# Title

[M-5] **matexpsym()** — Exponentiation and logarithms of symmetric matrices

## Syntax

*numeric matrix*     **matexpsym**(*numeric matrix A*)

*numeric matrix*     **matlogsym**(*numeric matrix A*)

*void*                 **\_matexpsym**(*numeric matrix A*)

*void*                 **\_matlogsym**(*numeric matrix A*)

## Description

**matexpsym**(*A*) returns the matrix exponential of the symmetric (Hermitian) matrix *A*.

**matlogsym**(*A*) returns the matrix natural logarithm of the symmetric (Hermitian) matrix *A*.

**\_matexpsym**(*A*) and **\_matlogsym**(*A*) do the same thing as **matexpsym**() and **matlogsym**(), but instead of returning the result, they store the result in *A*.

## Remarks

Do not confuse **matexpsym**(*A*) with **exp**(*A*), nor **matlogsym**(*A*) with **log**(*A*).

**matexpsym**(2\***matlogsym**(*A*)) produces the same result as *A*\**A*. **exp**() and **log**() return elementwise results.

Exponentiated results and logarithms are obtained by extracting the eigenvectors and eigenvalues of *A*, performing the operation on the eigenvalues, and then rebuilding the matrix. That is, first *X* and *L* are found such that

$$AX = X * \text{diag}(L) \tag{1}$$

For symmetric (Hermitian) matrix *A*, *X* is orthogonal, meaning  $X'X = XX' = I$ . Thus

$$A = X * \text{diag}(L) * X' \tag{2}$$

**matexpsym**(*A*) is then defined

$$A = X * \text{diag}(\exp(L)) * X' \tag{3}$$

and **matlogsym**(*A*) is defined

$$A = X * \text{diag}(\log(L)) * X' \tag{4}$$

(1) is obtained via **symeigensystem**(); see [M-5] **eigensystem**().

## Conformability

`matexpsym(A)`, `matlogsym(A)`:

*A*:  $n \times n$

*result*:  $n \times n$

`_matexpsym(A)`, `_matlogsym(A)`:

*input*:

*A*:  $n \times n$

*output*:

*A*:  $n \times n$

## Diagnostics

`matexpsym(A)`, `matlogsym(A)`, `_matexpsym(A)`, and `_matlogsym(A)` return missing results if *A* contains missing values.

Also:

1. These functions do not check that *A* is symmetric or Hermitian. If *A* is a real matrix, only the lower triangle, including the diagonal, is used. If *A* is a complex matrix, only the lower triangle and the real parts of the diagonal elements are used.
2. These functions return a matrix of the same storage type as *A*.

For `symatlog(A)`, this means that if *A* is real and the result cannot be expressed as a real, a matrix of missing values is returned. If you want the generalized solution, code `matlogsym(C(A))`. This is the same rule as with scalars: `log(-1)` evaluates to missing, but `log(C(-1))` is `3.14159265i`.

3. These functions are guaranteed to return a matrix that is numerically symmetric, Hermitian, or [symmetriconly](#) if theory states that the matrix should be symmetric, Hermitian, or symmetriconly. See [\[M-5\] matpowersym\(\)](#) for a discussion of this issue.

For the functions here, real function `exp(x)` is defined for all real values of *x* (ignoring overflow), and thus the matrix returned by `matexpsym()` will be symmetric (Hermitian).

The same is not true for `matlogsym()`. `log(x)` is not defined for  $x = 0$ , so if any of the eigenvalues of *A* are 0 or very small, a matrix of missing values will result. Also, `log(x)` is complex for  $x < 0$ , and thus if any of the eigenvalues are negative, the resulting matrix will be (1) missing if *A* is real stored as real, (2) `symmetriconly` if *A* contains reals stored as complex, and (3) general if *A* is complex.

## Also see

[\[M-5\] matpowersym\(\)](#) — Powers of a symmetric matrix

[\[M-5\] eigensystem\(\)](#) — Eigenvectors and eigenvalues

[\[M-4\] matrix](#) — Matrix functions

# Title

[M-5] **matpowersym()** — Powers of a symmetric matrix

## Syntax

*numeric matrix*    `matpowersym(numeric matrix A, real scalar p)`

*void*                `_matpowersym(numeric matrix A, real scalar p)`

## Description

`matpowersym(A, p)` returns  $A^p$  for symmetric matrix or Hermitian matrix  $A$ . The matrix returned is real if  $A$  is real and complex if  $A$  is complex.

`_matpowersym(A, p)` does the same thing, but instead of returning the result, it stores the result in  $A$ .

## Remarks

Do not confuse `matpowersym(A, p)` and  $A.^p$ . If  $p=2$ , the first returns  $A*A$  and the second returns  $A$  with each element squared.

Powers can be positive, negative, integer, or noninteger. Thus `matpowersym(A, .5)` is a way to find the square-root matrix  $R$  such that  $R*R=A$ , and `matpowersym(A, -1)` is a way to find the inverse. For inversion, you could obtain the result more quickly using other routines.

Powers are obtained by extracting the eigenvectors and eigenvalues of  $A$ , raising the eigenvalues to the specified power, and then rebuilding the matrix. That is, first  $X$  and  $L$  are found such that

$$AX = X * \text{diag}(L) \tag{1}$$

For symmetric (Hermitian) matrix  $A$ ,  $X$  is orthogonal, meaning  $X'X = XX' = I$ . Thus

$$A = X * \text{diag}(L)X' \tag{2}$$

$A^p$  is then defined

$$A^p = X * \text{diag}(L.^p) * X' \tag{3}$$

(1) is obtained via `symeigensystem()`; see [M-5] [eigensystem\(\)](#).

## Conformability

`matpowersym(A, p):`

$A$ :	$n \times n$
$p$ :	$1 \times 1$
<i>result</i> :	$n \times n$

`_matpowersym(A, p):`

*input:*

$A: \quad n \times n$

$p: \quad 1 \times 1$

*output:*

$A: \quad n \times n$

## Diagnostics

`matpowersym(A, p)` and `_matpowersym(A, p)` return missing results if  $A$  contains missing values.

Also:

1. These functions do not check that  $A$  is symmetric or Hermitian. If  $A$  is a real matrix, only the lower triangle, including the diagonal, is used. If  $A$  is a complex matrix, only the lower triangle and the real parts of the diagonal elements are used.
2. These functions return a matrix of the same storage type as  $A$ . That means that if  $A$  is real and  $A^p$  cannot be expressed as a real, a matrix of missing values is returned. If you want the generalized solution, code `matpowersym(C(A), p)`. This is the same rule as with scalars:  $(-1)^.5$  is missing, but  $C(-1)^.5$  is  $1i$ .
3. These functions are guaranteed to return a matrix that is numerically symmetric, Hermitian, or [symmetriconly](#) if theory states that the matrix should be symmetric, Hermitian, or symmetriconly.

Concerning theory, the returned result is not necessarily symmetric (Hermitian). The eigenvalues  $L$  of a symmetric (Hermitian) matrix are real. If  $L:~^p$  are real, then the returned matrix will be symmetric (Hermitian), but otherwise, it will not. Think of a negative eigenvalue and  $p = .5$ : this results in a complex eigenvalue for  $A^p$ . Then if the original matrix was real (the eigenvectors were real), the resulting matrix will be symmetriconly. If the original matrix was complex (the eigenvectors were complex), the resulting matrix will have no special structure.

## Also see

[M-5] [matexpsym\(\)](#) — Exponentiation and logarithms of symmetric matrices

[M-5] [eigensystem\(\)](#) — Eigenvectors and eigenvalues

[M-4] [matrix](#) — Matrix functions

## Syntax

*real rowvector*    `mean(X[, w])`

*real matrix*        `variance(X[, w])`

*real matrix*        `quadvariance(X[, w])`

*real matrix*        `meanvariance(X[, w])`

*real matrix*        `quadmeanvariance(X[, w])`

*real matrix*        `correlation(X[, w])`

*real matrix*        `quadcorrelation(X[, w])`

where

*X*:        *real matrix X* (rows are observations, columns are variables)

*w*:        *real colvector w* and is optional

## Description

`mean(X, w)` returns the weighted-or-unweighted column means of data matrix *X*. `mean()` uses quad precision in forming sums and so is very accurate.

`variance(X, w)` returns the weighted-or-unweighted variance matrix of *X*. In the calculation, means are removed and those means are calculated in quad precision, but quad precision is not otherwise used.

`quadvariance(X, w)` returns the weighted-or-unweighted variance matrix of *X*. Calculation is highly accurate; quad precision is used in forming all sums.

`meanvariance(X, w)` returns `mean(X,w)\variance(X,w)`.

`quadmeanvariance(X, w)` returns `mean(X,w)\quadvariance(X,w)`.

`correlation(X, w)` returns the weighted-or-unweighted correlation matrix of *X*. `correlation()` obtains the variance matrix from `variance()`.

`quadcorrelation(X, w)` returns the weighted-or-unweighted correlation matrix of *X*. `quadcorrelation()` obtains the variance matrix from `quadvariance()`.

In all cases, *w* specifies the weight. Omit *w*, or specify *w* as 1 to obtain unweighted means.

In all cases, rows of *X* or *w* that contain missing values are omitted from the calculation, which amounts to casewise deletion.



## Remarks

1. There is no `quadmean()` function because `mean()`, in fact, is `quadmean()`. The fact that `mean()` defaults to the quad-precision calculation reflects our judgment that the extra computational cost in computing means in quad precision is typically justified.
2. The fact that `variance()` and `correlation()` do not default to using quad precision for their calculations reflects our judgment that the extra computational cost is typically not justified. The emphasis on this last sentence is on the word typically.

It is easier to justify means in part because the extra computational cost is less: there are only  $k$  means but  $k(k+1)/2$  variances and covariances.

3. If you need just the mean or just the variance matrix, call `mean()` or `variance()` (or `quadvariance()`). If you need both, there is a CPU-time savings to be had by calling `meanvariance()` instead of the two functions separately (or `quadmeanvariance()` instead of calling `mean()` and `quadvariance()`).

The savings is not great—one `mean()` calculation is saved—but the greater `rows(X)`, the greater the savings.

Upon getting back the combined result, it can be efficiently broken into its components via

```
: var    = meanvariance(X)
: means  = var[1,.]
: var    = var[12,1  ..,1]
```

## Conformability

`mean(X, w):`

*X:*         $n \times k$   
*w:*         $n \times 1$     or     $1 \times 1$     (optional,  $w = 1$  assumed)  
*result:*     $1 \times k$

`variance(X, w), quadvariance(X, w), correlation(X, w), quadcorrelation(X, w):`

*X:*         $n \times k$   
*w:*         $n \times 1$     or     $1 \times 1$     (optional,  $w = 1$  assumed)  
*result:*     $k \times k$

`meanvariance(X, w), quadmeanvariance(X, w):`

*X:*         $n \times k$   
*w:*         $n \times 1$     or     $1 \times 1$     (optional,  $w = 1$  assumed)  
*result:*     $(k+1) \times k$

## Diagnostics

All functions omit from the calculation rows that contain missing values unless all rows contain missing values. Then the returned result contains all missing values.

## Also see

[M-4] [statistical](#) — Statistical functions

[M-5] mindouble() — Minimum and maximum nonmissing value

Syntax

```
real scalar mindouble()  
  
real scalar maxdouble()  
  
real scalar smallestdouble()
```

Description

mindouble() returns the largest negative, nonmissing value.  
maxdouble() returns the largest positive, nonmissing value.  
smallestdouble() returns the smallest full-precision value of  $e$ ,  $e > 0$ . The largest full-precision value of  $e$ ,  $e < 0$  is  $-\text{smallestdouble}()$ .

Remarks

All nonmissing values  $x$  fulfill  $\text{mindouble}() \leq x \leq \text{maxdouble}()$ .  
All missing values  $m$  fulfill  $m > \text{maxdouble}()$   
Missing values also fulfill  $m \geq .$   
On all computers on which Stata and Mata are currently implemented, which are computers following IEEE standards:

Function	Exact hexadecimal value	Approximate decimal value
mindouble()	-1.fffffffffffffX+3ff	-1.7977e+308
smallestdouble()	+1.0000000000000X-3fe	2.2251e-308
epsilon(1)	+1.0000000000000X-034	2.2205e-016
maxdouble()	+1.fffffffffffffX+3fe	8.9885e+307

The smallest missing value ( $. < .a < \dots < .z$ ) is  $+1.0000000000000X+3ff$ .  
Do not confuse smallestdouble() with the more interesting value epsilon(1). smallestdouble() is the smallest full-precision value of  $e$ ,  $e > 0$ . epsilon(1) is the smallest value of  $e$ ,  $e+1 > 1$ ; see [M-5] epsilon().

## Conformability

```
mindouble(), maxdouble(), smallestdouble():  
  result:      1 × 1
```

## Diagnostics

None.

## Reference

Linhart, J. M. 2008. [Mata Matters: Overflow, underflow and the IEEE floating-point format](#). *Stata Journal* 8: 255–268.

## Also see

[M-5] [epsilon\(\)](#) — Unit roundoff error (machine precision)

[M-4] [utility](#) — Matrix utility functions

## Syntax

*void minindex(real vector v, real scalar k, i, w)*

*void maxindex(real vector v, real scalar k, i, w)*

Results are returned in *i* and *w*.

*i* will be a *real colvector*.

*w* will be a  $K \times 2$  *real matrix*,  $K \leq |k|$ .

## Description

`minindex(v, k, i, w)` returns in *i* and *w* the indices of the *k* minimums of *v*.

`maxindex(v, k, i, w)` does the same, except that it returns the indices of the *k* maximums.

`minindex()` may be called with  $k < 0$ ; it is then equivalent to `maxindex()`.

`maxindex()` may be called with  $k < 0$ ; it is then equivalent to `minindex()`.

## Remarks

Remarks are presented under the following headings:

*Use of functions when v has all unique values*

*Use of functions when v has repeated (tied) values*

*Summary*

Remarks are cast in terms of `minindex()` but apply equally to `maxindex()`.

## Use of functions when v has all unique values

Consider  $v = (3, 1, 5, 7, 6)$ .

1. `minindex(v, 1, i, w)` returns  $i = 2$ , which means that  $v[2]$  is the minimum value in *v*.
2. `minindex(v, 2, i, w)` returns  $i = (2, 1)'$ , which means that  $v[2]$  is the minimum value of *v* and that  $v[1]$  is the second minimum.

...

5. `minindex(v, 5, i, w)` returns  $i = (2, 1, 3, 5, 4)'$ , which means that the ordered values in *v* are  $v[2]$ ,  $v[1]$ ,  $v[3]$ ,  $v[5]$ , and  $v[4]$ .
6. `minindex(v, 6, i, w)`, `minindex(v, 7, i, w)`, and so on, return the same as (5), because there are only five minimums in a five-element vector.

When *v* has unique values, the values returned in *w* are irrelevant.

- In (1),  $w$  will be (1, 1).
- In (2),  $w$  will be (1, 1\2, 1).
- ...
- In (5),  $w$  will be (1, 1\2, 1\3, 1\4, 1\5, 1).

The second column of  $w$  records the number of tied values. Since the values in  $v$  are unique, the second column of  $w$  will be ones. If you have a problem where you are uncertain whether the values in  $v$  are unique, code

```
if (!allof(w[,2], 1)) {
    /* uniqueness assumption false */
}
```

## Use of functions when $v$ has repeated (tied) values

Consider  $v = (3, 2, 3, 2, 3, 3)$ .

1. `minindex(v, 1, i, w)` returns  $i = (2, 4)'$ , which means that there is one minimum value and that it is repeated in two elements of  $v$ , namely,  $v[2]$  and  $v[4]$ .

Here,  $w$  will be (1, 2), but you can ignore that. There are two values in  $i$  corresponding to the same minimum.

When  $k=1$ , `rows(i)` equals the number of observations in  $v$  corresponding to the minimum, as does  $w[1,2]$ .

2. `minindex(v, 2, i, w)` returns  $i = (2, 4, 1, 3, 5, 6)'$  and  $w = (1, 2\3, 4)$ .

Begin with  $w$ . The first row of  $w$  is (1, 2), which states that the indices of the first minimums of  $v$  start at  $i[1]$  and consist of two elements. Thus the indices of the first minimums are  $i[1]$  and  $i[2]$  (the minimums are  $v[i[1]]$  and  $v[i[2]]$ , which of course are equal).

The second row of  $w$  is (3, 4), which states that the indices of the second minimums of  $v$  start at  $i[3]$  and consist of four elements:  $i[3]$ ,  $i[4]$ ,  $i[5]$ , and  $i[6]$  (which are 1, 3, 5, and 6).

In summary, `rows(w)` records the number of minimums returned.  $w[m,1]$  records where in  $i$  the  $m$ th minimum begins (it begins at  $i[w[m,1]]$ ).  $w[m,2]$  records the total number of tied values. Thus one could step across the minimums and the tied values by coding

```
minindex(v, k, i, w)
for (m=1; m<=rows(w); m++) {
    for (j=w[m,1]; j<w[m,1]+w[m,2]; j++) {
        /* i[j] is the index in v of an mth minimum */
    }
}
```

3. `minindex(v, 3, i, w)`, `minindex(v, 4, i, w)`, and so on, return the same as (2) because, with  $v = (3, 2, 3, 2, 3, 3)$ , there are only two minimums.

Summary

Consider `minindex(v, k, i, w)`. Returned will be

$$\begin{aligned}
 w &= \begin{bmatrix} i1 & n1 \\ i2 & n2 \\ \cdot & \cdot \\ \cdot & \cdot \end{bmatrix} && w : K \times 2, \quad K \leq |k| \\
 i &= \begin{bmatrix} j1 \\ j2 \\ j3 \\ j4 \\ \cdot \\ \cdot \\ \cdot \end{bmatrix} && \left. \begin{array}{l} \leftarrow i[i1] \text{ is start of first minimums} \\ \leftarrow i[i2] \text{ is start of second minimums} \\ \text{etc.} \end{array} \right\} \begin{array}{l} \text{has } n1 \text{ values} \\ \text{has } n2 \text{ values} \end{array} \\
 &&& i : 1 \times m, \quad m = n1 + n2 + \dots
 \end{aligned}$$

*j1, j2, ...,* are indices into *v*.

Conformability

`minindex(v, k, i, w)`, `maxindex(v, k, i, w)`:

*input:*

$$\begin{array}{ll}
 v: & n \times 1 \text{ or } 1 \times n \\
 k: & 1 \times 1
 \end{array}$$

*output:*

$$\begin{array}{ll}
 i: & L \times 1, \quad L \geq K \\
 w: & K \times 2, \quad K \leq |k|
 \end{array}$$

Diagnostics

`minindex(v, k, i, w)` and `maxindex(v, k, i, w)` abort with error if *i* or *w* is a view.

In `minindex(v, k, i, w)` and `maxindex(v, k, i, w)`, missing values in *v* are ignored in obtaining minimums and maximums.

In the examples above, we have shown input vector *v* as a row vector. It can also be a column vector; it makes no difference.

In `minindex(v, k, i, w)`, input argument *k* specifies the number of minimums to be obtained. *k* may be zero. If *k* is negative, *−k* maximums are obtained.

Similarly, in `maxindex(v, k, i, w)`, input argument *k* specifies the number of maximums to be obtained. *k* may be zero. If *k* is negative, *−k* minimums are obtained.

`minindex()` and `maxindex()` are designed for use when *k* is small relative to `length(v)`; otherwise, see `order()` in [M-5] [sort\(\)](#).

## Also see

[M-5] `minmax()` — Minimums and maximums

[M-4] `utility` — Matrix utility functions

## Syntax

<i>real colvector</i>	<code>rowmin(real matrix X)</code>
<i>real rowvector</i>	<code>colmin(real matrix X)</code>
<i>real scalar</i>	<code>min(real matrix X)</code>
<i>real colvector</i>	<code>rowmax(real matrix X)</code>
<i>real rowvector</i>	<code>colmax(real matrix X)</code>
<i>real scalar</i>	<code>max(real matrix X)</code>
<i>real matrix</i>	<code>rowminmax(real matrix X)</code>
<i>real matrix</i>	<code>colminmax(real matrix X)</code>
<i>real rowvector</i>	<code>minmax(real matrix X)</code>
<i>real matrix</i>	<code>rowminmax(real matrix X, real scalar usemiss)</code>
<i>real matrix</i>	<code>colminmax(real matrix X, real scalar usemiss)</code>
<i>real rowvector</i>	<code>minmax(real matrix X, real scalar usemiss)</code>
<i>real colvector</i>	<code>rowmaxabs(numeric matrix A)</code>
<i>real rowvector</i>	<code>colmaxabs(numeric matrix A)</code>

## Description

These functions return the indicated minimums and maximums of  $X$ .

`rowmin(X)` returns the minimum of each row of  $X$ , `colmin(X)` returns the minimum of each column, and `min(X)` returns the overall minimum. Elements of  $X$  that contain missing are ignored.

`rowmax(X)` returns the maximum of each row of  $X$ , `colmax(X)` returns the maximum of each column, and `max(X)` returns the overall maximum. Elements of  $X$  that contain missing are ignored.

`rowminmax(X)` returns the minimum and maximum of each row of  $X$  in an  $r \times 2$  matrix; `colminmax(X)` returns the minimum and maximum of each column in a  $2 \times c$  matrix; and `minmax(X)` returns the overall minimum and maximum. Elements of  $X$  that contain missing are ignored.

The two-argument versions of `rowminmax()`, `colminmax()`, and `minmax()` allow you to specify how missing values are to be treated. Specifying a second argument with value 0 is the same as using the single-argument versions of the functions. In the two-argument versions, if the second argument is not zero, missing values are treated like all other values in determining the minimums and maximums: `nonmissing < . < .a < .b < ... < .z`.



`rowmaxabs(A)` and `colmaxabs(A)` return the same result as `rowmax(abs(A))` and `colmax(abs(A))`. The advantage is that matrix `abs(A)` is never formed or stored, and so these functions use less memory.

## Conformability

`rowmin(X), rowmax(X):`

*X:*  $r \times c$   
*result:*  $r \times 1$

`colmin(X), colmax(X):`

*X:*  $r \times c$   
*result:*  $1 \times c$

`min(X), max(X):`

*X:*  $r \times c$   
*result:*  $1 \times 1$

`rowminmax(X, usemiss):`

*X:*  $r \times c$   
*usemiss:*  $1 \times 1$   
*result:*  $r \times 2$

`colminmax(X, usemiss)`

*X:*  $r \times c$   
*usemiss:*  $1 \times 1$   
*result:*  $2 \times c$

`minmax(X, usemiss)`

*X:*  $r \times c$   
*usemiss:*  $1 \times 1$   
*result:*  $1 \times 2$

`rowmaxabs(A):`

*A:*  $r \times c$   
*result:*  $r \times 1$

`colmaxabs(A):`

*A:*  $r \times c$   
*result:*  $1 \times c$

## Diagnostics

`row*()` functions return missing value for the corresponding minimum or maximum when the entire row contains missing.

`col*()` functions return missing value for the corresponding minimum or maximum when the entire column contains missing.

`min()` and `max()` return missing value when the entire matrix contains missing.

## Also see

[M-5] [minindex\(\)](#) — Indices of minimums and maximums

[M-4] [mathematical](#) — Important mathematical functions

[M-4] [utility](#) — Matrix utility functions

## Syntax

```

real rowvector  colmissing(numeric matrix X)
real colvector  rowmissing(numeric matrix X)
real scalar     missing(numeric matrix X)

real rowvector  colnonmissing(numeric matrix X)
real colvector  rownonmissing(numeric matrix X)
real scalar     nonmissing(numeric matrix X)
real scalar     hasmissing(numeric matrix X)

```

## Description

These functions return the indicated count of missing or nonmissing values.

`colmissing(X)` returns the count of missing values of each column of  $X$ , `rowmissing(X)` returns the count of missing values of each row, and `missing(X)` returns the overall count.

`colnonmissing(X)` returns the count of nonmissing values of each column of  $X$ , `rownonmissing(X)` returns the count of nonmissing values of each row, and `nonmissing(X)` returns the overall count.

`hasmissing(X)` returns 1 if  $X$  has a missing value or 0 if  $X$  does not have a missing value.

## Remarks

```

colnonmissing(X) = rows(X) :- colmissing(X)
rownonmissing(X) = cols(X) :- rowmissing(X)
nonmissing(X) = rows(X)*cols(X) - missing(X)

```

## Conformability

`colmissing(X)`, `colnonmissing(X)`:

```

X:       $r \times c$ 
result:  $1 \times c$ 

```

`rowmissing(X)`, `rownonmissing(X)`:

```

X:       $r \times c$ 
result:  $r \times 1$ 

```

`missing(X)`, `nonmissing(X)`, `hasmissing(X)`:

```

X:       $r \times c$ 
result:  $1 \times 1$ 

```

# Diagnostics

None.

# Also see

[M-5] [editmissing\(\)](#) — Edit matrix for missing values

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] **missingof()** — Appropriate missing value

## Syntax

*transmorphic scalar*    `missingof(transmorphic matrix A)`

## Description

`missingof(A)` returns a missing of the same element type as *A*:

- if *A* is real, a real missing is returned;
- if *A* is complex, a complex missing is returned;
- if *A* is pointer, NULL is returned;
- if *A* is string, "" is returned.

## Remarks

`missingof()` is useful when creating empty matrices of the same type as another matrix; for example,

`newmat = J(rows(x), cols(x), missingof(x))`

## Conformability

`missingof(A)`  
          *A*:         $r \times c$   
          *result*:     $1 \times 1$

## Diagnostics

None.

## Also see

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] `mod()` — Modulus

## Syntax

*real matrix* `mod`(*real matrix* *x*, *real matrix* *y*)

## Description

`mod(x, y)` returns the elementwise modulus of *x* with respect to *y*. `mod()` is defined

$$\text{mod}(x, y) = x - y * \text{trunc}(x/y)$$

## Conformability

`mod(x, y)`:

<i>x</i> :	$r_1 \times c_1$
<i>y</i> :	$r_2 \times c_2$ , <i>x</i> and <i>y</i> r-conformable
<i>result</i> :	$\max(r_1, r_2) \times \max(c_1, c_2)$ (elementwise calculation)

## Diagnostics

`mod(x, y)` returns missing when either argument is missing or when *y* = 0.

## Also see

[\[M-4\] scalar](#) — Scalar mathematical functions

## Syntax

If you are reading this entry for the first time, skip down to [Description](#) and to [Remarks](#), and more especially, to [Mathematical statement of the `moptimize\(\)` problem](#) under Remarks.

Syntax is presented under the following headings:

- [Step 1: Initialization](#)
- [Step 2: Definition of maximization or minimization problem](#)
- [Step 3: Perform optimization or perform a single function evaluation](#)
- [Step 4: Post, display, or obtain results](#)
- [Utility functions for use in all steps](#)
- [Definition of  \$M\$](#)
- [Setting the sample](#)
- [Specifying dependent variables](#)
- [Specifying independent variables](#)
- [Specifying constraints](#)
- [Specifying weights or survey data](#)
- [Specifying clusters and panels](#)
- [Specifying optimization technique](#)
- [Specifying initial values](#)
- [Performing one evaluation of the objective function](#)
- [Performing optimization of the objective function](#)
- [Tracing optimization](#)
- [Specifying convergence criteria](#)
- [Accessing results](#)
- [Stata evaluators](#)
- [Advanced functions](#)
- [Syntax of evaluators](#)
- [Syntax of type \*lf\* evaluators](#)
- [Syntax of type \*d\* evaluators](#)
- [Syntax of type \*lf\\*\* evaluators](#)
- [Syntax of type \*gf\* evaluators](#)
- [Syntax of type \*q\* evaluators](#)
- [Passing extra information to evaluators](#)
- [Utility functions](#)

### Step 1: Initialization

```
M = moptimize_init()
```

## Step 2: Definition of maximization or minimization problem

*In each of the functions, the last argument is optional. If specified, the function sets the value and returns void. If not specified, no change is made, and instead what is currently set is returned.*

(varies) `moptimize_init_which(M, { "max" | "min" })`

(varies) `moptimize_init_evaluator(M, &functionname())`

(varies) `moptimize_init_evaluator(M, "programname")`

(varies) `moptimize_init_evaluatortype(M, evaluatortype)`

(varies) `moptimize_init_negH(M, { "off" | "on" })`

(varies) `moptimize_init_touse(M, "tousevarname")`

(varies) `moptimize_init_ndepvars(M, D)`

(varies) `moptimize_init_depvar(M, j, y)`

(varies) `moptimize_init_eq_n(M, m)`

(varies) `moptimize_init_eq_indepvars(M, i, X)`

(varies) `moptimize_init_eq_cons(M, i, { "on" | "off" })`

(varies) `moptimize_init_eq_offset(M, i, o)`

(varies) `moptimize_init_eq_exposure(M, i, t)`

(varies) `moptimize_init_eq_name(M, i, name)`

(varies) `moptimize_init_eq_colnames(M, i, names)`

(varies) `moptimize_init_eq_coefs(M, i, b0)`

(varies) `moptimize_init_constraints(M, Cc)`

(varies) `moptimize_init_search(M, { "on" | "off" })`

(varies) `moptimize_init_search_random(M, { "off" | "on" })`

(varies) `moptimize_init_search_repeat(M, nr)`

(varies) `moptimize_init_search_bounds(M, i, minmax)`

(varies) `moptimize_init_search_rescale(M, { "on" | "off" })`



(varies) `moptimize_init_weight(M, w)`

(varies) `moptimize_init_weighttype(M, weighttype)`

(varies) `moptimize_init_cluster(M, c)`

(varies) `moptimize_init_svy(M, {"off" | "on"})`

(varies) `moptimize_init_by(M, by)`

(varies) `moptimize_init_nuserinfo(M, n_user)`

(varies) `moptimize_init_userinfo(M, l, Z)`

(varies) `moptimize_init_technique(M, technique)`

(varies) `moptimize_init_vcetype(M, vcetype)`

(varies) `moptimize_init_nmsimplexdeltas(M, delta)`

(varies) `moptimize_init_gnweightmatrix(M, W)`

(varies) `moptimize_init_singularHmethod(M, singularHmethod)`

(varies) `moptimize_init_conv_maxiter(M, maxiter)`

(varies) `moptimize_init_conv_warning(M, {"on" | "off"})`

(varies) `moptimize_init_conv_ptol(M, ptol)`

(varies) `moptimize_init_conv_vtol(M, vtol)`

(varies) `moptimize_init_conv_nrtol(M, nrtol)`

(varies) `moptimize_init_conv_ignorenrtol(M, {"off" | "on"})`

(varies) `moptimize_init_iterid(M, id)`

(varies) `moptimize_init_valueid(M, id)`

(varies) `moptimize_init_tracelevel(M, tracelevel)`

(varies) `moptimize_init_trace_ado(M, {"off" | "on"})`

(varies) `moptimize_init_trace_dots(M, {"off" | "on"})`

(varies) `moptimize_init_trace_value(M, {"on" | "off"})`

(varies) `moptimize_init_trace_tol(M, {"off" | "on"})`

(varies) `moptimize_init_trace_step(M, {"off" | "on"})`

(varies) `moptimize_init_trace_coefdiffs(M, {"off" | "on"})`

(varies) `moptimize_init_trace_coefs(M, {"off" | "on"})`

(varies) `moptimize_init_trace_gradient(M, {"off" | "on"})`

(varies) `moptimize_init_trace_Hessian(M, {"off" | "on"})`

(varies) `moptimize_init_evaluations(M, { "off" | "on" })`

(varies) `moptimize_init_verbose(M, { "on" | "off" })`

### Step 3: Perform optimization or perform a single function evaluation

*void* `moptimize(M)`

*real scalar* `_moptimize(M)`

*void* `moptimize_evaluate(M)`

*real scalar* `_moptimize_evaluate(M)`

### Step 4: Post, display, or obtain results

*void* `moptimize_result_post(M [, vcetype])`

*void* `moptimize_result_display([M [, vcetype]])`

*real scalar* `moptimize_result_value(M)`

*real scalar* `moptimize_result_value0(M)`

*real rowvector* `moptimize_result_eq_coefs(M [, i])`

*real rowvector* `moptimize_result_coefs(M)`

*string matrix* `moptimize_result_colstripe(M [, i])`

*real matrix* `moptimize_result_scores(M)`

*real rowvector* `moptimize_result_gradient(M [, i])`

*real matrix* `moptimize_result_Hessian(M [, i])`

*real matrix* `moptimize_result_V(M [, i])`

*string scalar* `moptimize_result_Vtype(M)`

*real matrix* `moptimize_result_V_oim(M [, i])`

*real matrix* `moptimize_result_V_opg(M [, i])`

*real matrix* `moptimize_result_V_robust(M [, i])`

*real scalar*     `moptimize_result_iterations(M)`  
*real scalar*     `moptimize_result_converged(M)`  
*real colvector* `moptimize_result_iterationlog(M)`  
*real rowvector* `moptimize_result_evaluations(M)`  
*real scalar*     `moptimize_result_errorcode(M)`  
*string scalar*   `moptimize_result_errortext(M)`  
*real scalar*     `moptimize_result_returncode(M)`  
*void*             `moptimize_ado_cleanup(M)`

## Utility functions for use in all steps

*void*             `moptimize_query(M)`

*real matrix*     `moptimize_util_eq_indices(M, i [, i2])`

*(varies)*        `moptimize_util_depvar(M, j)`  
*returns y set by moptimize\_init\_depvar(M, j, y), which is usually a real colvector*

*real colvector* `moptimize_util_xb(M, b, i)`

*real scalar*     `moptimize_util_sum(M, real colvector v)`

*real rowvector* `moptimize_util_vecsum(M, i, real colvector s, real scalar value)`

*real matrix*     `moptimize_util_matsum(M, i, i2, real colvector s,`  
*real scalar value)*

*real matrix*     `moptimize_util_matbysum(M, i, real colvector a, real colvector b,`  
*real scalar value)*

*real matrix*     `moptimize_util_matbysum(M, i, i2, real colvector a,`  
*real colvector b, real colvector c, real scalar value)*

*pointer scalar* `moptimize_util_by(M)`

## Definition of M

*M*, if it is declared, should be declared transmorphic. *M* is obtained from `moptimize_init()` and then passed as an argument to the other `moptimize()` functions.

`moptimize_init()` returns *M*, called an `moptimize()` problem handle. The function takes no arguments. *M* holds the information about the optimization problem.

## Setting the sample

Various `moptimize_init_*`() functions set values for dependent variables, independent variables, etc. When you set those values, you do that either by specifying Stata variable names or by specifying Mata matrices containing the data themselves. Function `moptimize_init_touse()` specifies the sample to be used when you specify Stata variable names.

`moptimize_init_touse(M, "tousevarname")` specifies the name of the variable in the Stata dataset that marks the observations to be included. Observations for which the Stata variable is nonzero are included. The default is "", meaning all observations are to be used.

You need to specify *tousevarname* only if you specify Stata variable names in the other `moptimize_init_*`() functions, and even then it is not required. Setting *tousevar* when you specify the data themselves via Mata matrices, whether views or not, has no effect.

## Specifying dependent variables

*D* and *j* index dependent variables:

<i>index</i>	Description
<i>D</i>	number of dependent variables, $D \geq 0$
<i>j</i>	dependent variable index, $1 \leq j \leq D$

*D* and *j* are real scalars.

You set the dependent variables one at a time. In a particular optimization problem, you may have no dependent variables or have more dependent variables than [equations](#).

`moptimize_init_depvar(M, j, y)` sets the *j*th dependent variable to be *y*. *y* may be a string scalar containing a Stata variable name that in turn contains the values of the *j*th dependent variable, or *y* may be a real colvector directly containing the values.

`moptimize_init_ndepvars(M, D)` sets the total number of dependent variables. You can set *D* before defining dependent variables, and that speeds execution slightly, but it is not necessary because *D* is automatically set to the maximum *j*.

## Specifying independent variables

Independent variables are defined within parameters or, equivalently, equations. The words parameter and equation mean the same thing. *m*, *i*, and *i2* index parameters:

<i>index</i>	Description
<i>m</i>	number of parameters (equations), $m \geq 1$
<i>i</i>	equation index, $1 \leq i \leq m$
<i>i2</i>	equation index, $1 \leq i2 \leq m$

*m*, *i*, and *i2* are real scalars.

The function to be optimized is  $f(p_1, p_2, \dots, p_m)$ . The *i*th parameter (equation) is defined as

$$pi = Xi \times bi' + oi + \ln(ti) :+ b0i$$

where

<i>pi</i> : $Ni \times 1$	( <i>i</i> th parameter)
<i>Xi</i> : $Ni \times ki$	( <i>Ni</i> observations on <i>ki</i> independent variables)
<i>bi</i> : $1 \times ki$	(coefficients to be fit)
<i>oi</i> : $Ni \times 1$	(exposure/offset in offset form, optional)
<i>ti</i> : $Ni \times 1$	(exposure/offset in exposure form, optional)
<i>b0i</i> : $1 \times 1$	(constant or intercept, optional)

Any of the terms may be omitted. The most common forms for a parameter are  $pi = Xi \times bi' + b0i$  (standard model),  $pi = Xi \times bi'$  (no-constant model), and  $pi = b0i$  (constant-only model).

In addition, define *b*:  $1 \times K$  as the entire coefficient vector, which is to say,

$$b = (b1, [b01,] \quad b2, [b02,] \quad \dots)$$

That is, because *bi* is  $1 \times ki$  for  $i = 1, 2, \dots, m$ , then *b* is  $1 \times K$ , where  $K = \sum_i ki + ci$ , where *ci* is 1 if equation *i* contains an intercept and is 0 otherwise. Note that *bi* does not contain the constant or intercept, if there is one, but *b* contains all the coefficients, including the intercepts. *b* is called *the full set of coefficients*.

Parameters are defined one at a time by using the following functions:

`moptimize_init_eq_n(M, m)` sets the number of parameters. Use of this function is optional; *m* will be automatically determined from the other `moptimize_init_eq_*`() functions you issue.

`moptimize_init_eq_indepvars(M, i, X)` sets *X* to be the data (independent variables) for the *i*th parameter. *X* may be a  $1 \times ki$  string rowvector containing Stata variable names, or *X* may be a string scalar containing the same names in space-separated format, or *X* may be an  $Ni \times ki$  real matrix containing the data for the independent variables. Specify *X* as "" to omit term  $Xi \times bi'$ , for instance, as when fitting a constant-only model. The default is "".

`moptimize_init_eq_cons(M, i, { "on" | "off" })` specifies whether the equation for the *i*th parameter includes *b0i*, a constant or intercept. Specify "on" to include *b0i*, "off" to exclude it. The default is "on".

`moptimize_init_eq_offset(M, i, o)` specifies  $oi$  in the equation for the  $i$ th parameter. *o* may be a string scalar containing a Stata variable name, or *o* may be an  $Ni \times 1$  real colvector containing the offsets. The default is "", meaning term  $oi$  is omitted. Parameters may not have both  $oi$  and  $\ln(ti)$  terms.

`moptimize_init_eq_exposure(M, i, t)` specifies  $ti$  in term  $\ln(ti)$  of the equation for the  $i$ th parameter. *t* may be a string scalar containing a Stata variable name, or *t* may be an  $Ni \times 1$  real colvector containing the exposure values. The default is "", meaning term  $\ln(ti)$  is omitted.

`moptimize_init_eq_name(M, i, name)` specifies a string scalar, *name*, to be used in the output to label the  $i$ th parameter. The default is to use an automatically generated name.

`moptimize_init_eq_colnames(M, i, names)` specifies a  $1 \times ki$  string rowvector, *names*, to be used in the output to label the coefficients for the  $i$ th parameter. The default is to use automatically generated names.

## Specifying constraints

Linear constraints may be placed on the coefficients, *b*, which may be either within equation or between equations.

`moptimize_init_constraints(M, Cc)` specifies an  $R \times K + 1$  real matrix, *Cc*, that places *R* linear restrictions on the  $1 \times K$  full set of coefficients, *b*. Think of *Cc* as being (*C*, *c*), *C*:  $R \times K$  and *c*:  $R \times 1$ . Optimization will be performed subject to the constraint  $Cb' = c$ . The default is no constraints.

## Specifying weights or survey data

You may specify weights, and once you do, everything is automatic, assuming you implement your [evaluator](#) by using the provided [utility functions](#).

`moptimize_init_weight(M, w)` specifies the weighting variable or data. *w* may be a string scalar containing a Stata variable name, or *w* may be a real colvector directly containing the weight values. The default is "", meaning no weights.

`moptimize_init_weighttype(M, weighttype)` specifies how *w* is to be treated. *weighttype* may be "fweight", "aweight", "pweight", or "iweight". You may set *w* first and then *weighttype*, or the reverse. If you set *w* without setting *weighttype*, then "fweight" is assumed. If you set *weighttype* without setting *w*, then *weighttype* is ignored. The default *weighttype* is "fweight".

Alternatively, you may inherit the full set of [survey settings](#) from Stata by using `moptimize_init_svy()`. If you do this, do not use `moptimize_init_weight()`, `moptimize_init_weighttype()`, or `moptimize_init_cluster()`. When you use the survey settings, everything is nearly automatic, assuming you use the provided [utility functions](#) to implement your [evaluator](#). The proviso is that your evaluator must be of [evaluator type](#) lf, lf\*, gf, or q.

`moptimize_init_svy(M, {"off" | "on"})` specifies whether Stata's survey settings should be used. The default is "off". Using the survey settings changes the default [vcetype](#) to "svy", which is equivalent to "robust".

## Specifying clusters and panels

Clustering refers to possible nonindependence of the observations within groups called clusters. A cluster variable takes on the same value within a cluster and different values across clusters. After setting the cluster variable, there is nothing special you have to do, but be aware that clustering is allowed only if you use a type lf, lf\*, gf, or q [evaluator](#). `moptimize_init_cluster()` allows you to set a cluster variable.

Panels refer to likelihood functions or other objective functions that can only be calculated at the panel level, for which there is no observation-by-observation decomposition. Unlike clusters, these panel likelihood functions are difficult to calculate and require the use of type d or gf [evaluator](#). A panel variable takes on the same value within a panel and different values across panels. `moptimize_init_by()` allows you to set a panel variable.

You may set both a cluster variable and a panel variable, but be careful because, for most likelihood functions, panels are mathematically required to be nested within cluster.

`moptimize_init_cluster(M, c)` specifies a cluster variable. *c* may be a string scalar containing a Stata variable name, or *c* may be a real colvector directly containing the cluster values. The default is "", meaning no clustering. If clustering is specified, the default *vcetype* becomes "robust".

`moptimize_init_by(M, by)` specifies a panel variable and specifies that only panel-level calculations are meaningful. *by* may be a string scalar containing a Stata variable name, or *by* may be a real colvector directly containing the panel ID values. The default is "", meaning no panels. If panels are specified, the default *vcetype* remains unchanged, but if the opg variance estimator is used, the opg calculation is modified so that it is clustered at the panel level.

## Specifying optimization technique

Technique refers to the numerical methods used to solve the optimization problem. The default is Newton–Raphson maximization.

`moptimize_init_which(M, { "max" | "min" })` sets whether the maximum or minimum of the objective function is to be found. The default is "max".

`moptimize_init_technique(M, technique)` specifies the technique to be used to find the coefficient vector *b* that maximizes or minimizes the objective function. Allowed values are

<i>technique</i>	Description
"nr"	modified Newton–Raphson
"dfp"	Davidon–Fletcher–Powell
"bfgs"	Broyden–Fletcher–Goldfarb–Shanno
"bhhh"	Berndt–Hall–Hall–Hausman
"nm"	Nelder–Mead
"gn"	Gauss–Newton (quadratic optimization)

The default is "nr".

You can switch between "nr", "dfp", "bfgs", and "bhhh" by specifying two or more of them in a space-separated list. By default, `moptimize()` will use an algorithm for five iterations before switching to the next algorithm. To specify a different number of iterations, include

the number after the technique. For example, specifying `moptimize_init_technique(M, "bhhh 10 nr 1000")` requests that `moptimize()` perform 10 iterations using the Berndt–Hall–Hall–Hausman algorithm, followed by 1,000 iterations using the modified Newton–Raphson algorithm, and then switch back to Berndt–Hall–Hall–Hausman for 10 iterations, and so on. The process continues until [convergence](#) or until *maxiter* is exceeded.

`moptimize_init_singularHmethod(M, singularHmethod)` specifies the action to be taken during optimization if the Hessian is found to be singular and the *technique* requires the Hessian be of full rank. Allowed values are

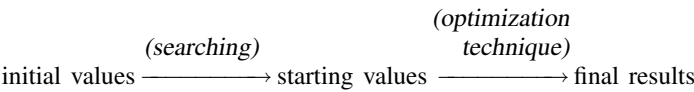
<i>singularHmethod</i>	Description
"m-marquardt"	modified Marquardt algorithm
"hybrid"	mixture of steepest descent and Newton

The default is "m-marquardt".  
"hybrid" is equivalent to ml’s difficult option; see [\[R\] ml](#).

`moptimize_init_nmsimplexdeltas(M, delta)` is for use with Nelder–Mead, also known as technique nm. This function sets the values of *delta* to be used, along with the initial parameters, to build the simplex required by Nelder–Mead. Use of this function is required only in the Nelder–Mead case. The values in *delta* must be at least 10 times larger than *ptol*. The initial simplex will be  $\{p, p + (d_1, 0, \dots, 0), p + (0, d_2, 0, \dots, 0), \dots, p + (0, 0, \dots, 0, d_K)\}$ .

### Specifying initial values

Initial values are values you optionally specify that via a search procedure result in starting values that are then used for the first iteration of the optimization technique. That is,



Initial values are specified [parameter](#) by parameter.

`moptimize_init_eq_coefs(M, i, b0)` sets the initial values of the coefficients for the *i*th parameter to be *b0*:  $1 \times (ki + ci)$ . The default is (0, 0, ..., 0).

The following functions control whether searching is used to improve on the initial values to produce better starting values. In addition to searching a predetermined set of hardcoded starting values, there are two other methods that can be used to improve on the initial values: random and rescaling. By default, random is off and rescaling is on. You can use one, the other, or both.

`moptimize_init_search(M, {"on"|"off"})` determines whether any attempts are to be made to improve on the initial values via a search technique. The default is "on". If you specify "off", the initial values become the starting values.

`moptimize_init_search_random(M, {"off"|"on"})` determines whether the random method of improving initial values is to be attempted. The default is "off". Use of the random method is recommended when the initial values are or might be infeasible. Infeasible means that the function cannot be evaluated, which mechanically corresponds to the user-written evaluator returning a missing value. The random method is seldom able to improve on feasible initial values. It works well when the initial values are or might be infeasible.



`moptimize_init_search_repeat(M, nr)` controls how many times random values are tried if the random method is turned on. The default is 10.

`moptimize_init_search_bounds(M, i, minmax)` specifies the bounds for the random search. *minmax* is a  $1 \times 2$  real rowvector containing the minimum and maximum values for the *i*th parameter (equation). The default is `(., .)`, meaning no lower and no upper bounds.

`moptimize_init_search_rescale(M, { "on" | "off" })` determines whether rescaling is attempted. The default is "on". Rescaling is a deterministic (not random) method. It also usually improves initial values, and usually reduces the number of subsequent iterations required by the optimization technique.

## Performing one evaluation of the objective function

`moptimize_evaluate(M)` and `_moptimize_evaluate(M)` perform one evaluation of the function evaluated at the [initial values](#). Results can be accessed by using `moptimize_result_*`(), including first- and second-derivative-based results.

`moptimize_evaluate()` and `_moptimize_evaluate()` do the same thing, differing only in that `moptimize_evaluate()` aborts with a nonzero return code if things go badly, whereas `_moptimize_evaluate()` returns the real scalar [error code](#). An infeasible initial value is an error.

The evaluation is performed at the [initial values](#), not the [starting values](#), and this is true even if search is turned on. If you want to perform an evaluation at the starting values, then perform [optimization](#) with *maxiter* set to 0.

## Performing optimization of the objective function

`moptimize(M)` and `_moptimize(M)` perform optimization. Both routines do the same thing; they differ only in their behavior when things go badly. `moptimize()` returns nothing and aborts with error. `_moptimize()` returns a real scalar [error code](#). `moptimize()` is best for interactive use and often adequate for use in programs that do not want to consider the possibility that optimization might fail.

The optimization process is as follows:

1. The [initial values](#) are used to create [starting values](#). The value of the function at the starting values is calculated. If that results in a missing value, the starting values are declared infeasible. `moptimize()` aborts with return code 430; `_moptimize()` returns a nonzero error code, which maps to 430 via `moptimize_result_returncode()`. This step is called iteration 0.
2. The starting values are passed to the [technique](#) to produce better values. Usually this involves the technique calculating first and second derivatives, numerically or analytically, and then stepping multiple times in the appropriate direction, but techniques can vary on this. In general, the technique performs what it calls one iteration, the result of which is to produce better values. Those new values then become the starting values and the process repeats.

An iteration is said to fail if the new coefficient vector is infeasible (results in a missing value). Then attempts are made to recover and, if those attempts are successful, optimization continues. If they fail, `moptimize()` aborts with error and `_moptimize()` returns a nonzero error code.

Other problems may arise, such as singular Hessians or the inability to find better values. Various fix-ups are made and optimization continues. These are not failures.

This step is called iterations 1, 2, and so on.

3. Step 2 continues either until the process `converges` or until the `maximum number of iterations` (*maxiter*) is exceeded. Stopping due to *maxiter* is not considered an error. Upon completion, programmers should check `moptimize_result_converged()`.

If optimization succeeds, which is to say, if `moptimize()` does not abort or `_moptimize()` returns 0, you can use the `moptimize_result_*`() functions to access results.

### Tracing optimization

`moptimize()` and `_moptimize()` will produce output like

```
Iteration 0:  f(p) = .....
Iteration 1:  f(p) = .....
```

You can change the *f(p)* to be “log likelihood” or whatever else you want. You can also change “Iteration”.

`moptimize_init_iterid(M, id)` sets the string to be used to label the iterations in the iteration log. *id* is a string scalar. The default is “Iteration”.

`moptimize_init_valueid(M, id)` sets the string to be used to label the objective function value in the iteration log. *id* is a string scalar. The default is “f(p)”.

Additional results can be displayed during optimization, which can be useful when you are debugging your `evaluator`. This is called tracing the execution.

`moptimize_init_tracelevel(M, tracelevel)` specifies the output to be displayed during the optimization process. Allowed values are

<i>tracelevel</i>	To be displayed each iteration
"none"	nothing
"value"	function value
"tolerance"	previous + convergence values
"step"	previous + stepping information
"coefdiffs"	previous + parameter relative differences
"paramdiffs"	same as "coefdiffs"
"coefs"	previous + parameter values
"params"	same as "coefs"
"gradient"	previous + gradient vector
"hessian"	previous + Hessian matrix

The default is “value”.

Setting *tracelevel* is a shortcut. The other trace functions allow you to turn on and off individual features. In what follows, the documented defaults are the defaults when *tracelevel* is “value”.

`moptimize_init_trace_ado(M, { "off" | "on" })` traces the execution of evaluators written as ado-files. This topic is not discussed in this manual entry. The default is “off”.

`moptimize_init_trace_dots(M, {"off" | "on"})` displays a dot each time your evaluator is called. The default is "off".

`moptimize_init_trace_value(M, {"on" | "off"})` displays the function value at the start of each iteration. The default is "on".

`moptimize_init_trace_tol(M, {"off" | "on"})` displays the value of the calculated result that is compared with the effective [convergence](#) criterion at the end of each iteration. The default is "off".

`moptimize_init_trace_step(M, {"off" | "on"})` displays the steps within iteration. Listed are the value of objective function along with the word forward or backward. The default is "off".

`moptimize_init_trace_coefdiffs(M, {"off" | "on"})` displays the coefficient relative differences from the previous iteration that are greater than the coefficient tolerance *ptol*. The default is "off".

`moptimize_init_trace_coefs(M, {"off" | "on"})` displays the coefficients at the start of each iteration. The default is "off".

`moptimize_init_trace_gradient(M, {"off" | "on"})` displays the gradient vector at the start of each iteration. The default is "off".

`moptimize_init_trace_hessian(M, {"off" | "on"})` displays the Hessian matrix at the start of each iteration. The default is "off".

## Specifying convergence criteria

Convergence is based on several rules controlled by four parameters: *maxiter*, *ptol*, *vtol*, and *nrtol*. The first rule is not a convergence rule, but a stopping rule, and it is controlled by *maxiter*.

`moptimize_init_conv_maxiter(M, maxiter)` specifies the maximum number of iterations. If this number is exceeded, optimization stops and results are posted where they are accessible by using the `moptimize_result_*`(*)* functions, just as if convergence had been achieved. `moptimize_result_converged()`, however, is set to 0 rather than 1. The default *maxiter* is Stata's `c(maxiter)`, which is usually 16,000.

`moptimize_init_conv_warning(M, {"on" | "off"})` specifies whether the warning message "convergence not achieved" is to be displayed when this stopping rule is invoked. The default is "on".

Usually, convergence occurs before the stopping rule comes into effect. The convergence criterion is a function of three real scalar values: *ptol*, *vtol*, and *nrtol*. Let

*b* = full set of coefficients  
*b\_prior* = value of *b* from prior iteration  
*v* = value of objective function  
*v\_prior* = value of *v* from prior iteration  
*g* = gradient vector from this iteration  
*H* = Hessian matrix from this iteration

Define, for maximization,

$$\begin{aligned} C\_ptol: & \quad \text{mreldif}(b, b\_prior) \leq ptol \\ C\_vtol: & \quad \text{reldif}(v, v\_prior) \leq vtol \\ C\_nrtol: & \quad g \times \text{invsym}(-H) \times g' < nrtol \\ C\_concave: & \quad -H \text{ is positive semidefinite} \end{aligned}$$

For minimization, think in terms of maximization of  $-f(p)$ . Convergence is declared when

$$(C\_ptol \mid C\_vtol) \ \& \ C\_nrtol \ \& \ C\_concave$$

The above applies in cases of derivative-based optimization, which currently is all [techniques](#) except "nm" (Nelder–Mead). In the Nelder–Mead case, the criterion is

$$\begin{aligned} C\_ptol: & \quad \text{mreldif}(\text{vertices of } R) \leq ptol \\ C\_vtol: & \quad \text{reldif}(R) \leq vtol \end{aligned}$$

where  $R$  is the minimum and maximum values on the simplex. Convergence is declared when  $C\_ptol \mid C\_vtol$ .

The values of *ptol*, *vtol*, and *nrtol* are set by the following functions:

`moptimize_init_conv_ptol(M, ptol)` sets *ptol*. The default is 1e–6.

`moptimize_init_conv_vtol(M, vtol)` sets *vtol*. The default is 1e–7.

`moptimize_init_conv_nrtol(M, nrtol)` sets *nrtol*. The default is 1e–5.

`moptimize_init_conv_ignorenrtol(M, {"off" | "on"})` sets whether *C\_nrtol* should always be treated as true, which in effect removes the *nrtol* criterion from the convergence rule. The default is "off".

## Accessing results

Once you have successfully performed [optimization](#), or you have successfully performed a [single function evaluation](#), you may display results, post results to Stata, or access individual results.

To display results, use `moptimize_result_display()`.

`moptimize_result_display(M)` displays estimation results. Standard errors are shown using the default *vcetype*.

`moptimize_result_display(M, vcetype)` displays estimation results. Standard errors are shown using the specified *vcetype*.

Also there is a third syntax for use after results have been posted to Stata, which we will discuss below.

`moptimize_result_display()` without arguments (not even *M*) displays the estimation results currently posted in Stata.

*vcetype* specifies how the variance–covariance matrix of the estimators (VCE) is to be calculated. Allowed values are

<i>vcetype</i>	Description
"	use default for <a href="#">technique</a>
"oim"	observed information matrix
"opg"	outer product of gradients
"robust"	Huber/White/sandwich estimator
"svy"	survey estimator; equivalent to <code>robust</code>

The default *vcetype* is `oim` except for technique `bhhh`, where it is `opg`.  
If survey, `pweights`, or clusters are used, the default becomes `robust` or `svy`.

As an aside, if you set `moptimize_init_vcetype()` during initialization, that changes the default.

`moptimize_init_vcetype(M, vcetype)`, *vcetype* being a string scalar, resets the default *vcetype*.

To post results to Stata, use `moptimize_result_post()`.

`moptimize_result_post(M)` posts estimation results to Stata where they can be displayed with Mata function `moptimize_result_post()` (without arguments) or with Stata command `ereturn display` (see [\[P\] ereturn](#)). The posted VCE will be of the default *vcetype*.

`moptimize_result_post(M, vcetype)` does the same thing, except the VCE will be of the specified *vcetype*.

The remaining `moptimize_result_*`() functions simply return the requested result. It does not matter whether results have been posted or previously displayed.

`moptimize_result_value(M)` returns the real scalar value of the objective function.

`moptimize_result_value0(M)` returns the real scalar value of the objective function at the [starting values](#).

`moptimize_result_eq_coefs(M [, i])` returns the  $1 \times (ki + ci)$  coefficient rowvector for the *i*th equation. If  $i \geq .$  or argument *i* is omitted, the  $1 \times K$  [full set of coefficients](#) is returned.

`moptimize_result_coefs(M)` returns the  $1 \times K$  [full set of coefficients](#).

`moptimize_result_colstripe(M [, i])` returns a  $(ki + ci) \times 2$  string matrix containing, for the *i*th equation, the equation names in the first column and the coefficient names in the second. If  $i \geq .$  or argument *i* is omitted, the result is  $K \times 2$ .

`moptimize_result_scores(M)` returns an  $N \times m$  ([evaluator types](#) `lf` and `lf*`), or an  $N \times K$  (evaluator type `gf`), or an  $L \times K$  (evaluator type `q`) real matrix containing the observation-by-observation scores. For all other evaluator types, `J(0,0,.)` is returned. For evaluator types `lf` and `lf*`, scores are defined as the derivative of the objective function with respect to the [parameters](#). For evaluator type `gf`, scores are defined as the derivative of the objective function with respect to the [coefficients](#). For evaluator type `q`, scores are defined as the derivatives of the *L* [independent elements](#) with respect to the coefficients.

`moptimize_result_gradient(M [, i])` returns the  $1 \times (ki + ci)$  gradient rowvector for the *i*th equation. If  $i \geq .$  or argument *i* is omitted, the  $1 \times K$  gradient corresponding to the [full set](#)

of coefficients is returned. Gradient is defined as the derivative of the objective function with respect to the coefficients.

`moptimize_result_Hessian(M [, i])` returns the  $(ki + ci) \times (ki + ci)$  Hessian matrix for the *i*th equation. If  $i \geq .$  or argument *i* is omitted, the  $K \times K$  Hessian corresponding to the full set of coefficients is returned. The Hessian is defined as the second derivative of the objective function with respect to the coefficients.

`moptimize_result_V(M [, i])` returns the appropriate  $(ki + ci) \times (ki + ci)$  submatrix of the full variance matrix calculated according to the default *vcetype*. If  $i \geq .$  or argument *i* is omitted, the full  $K \times K$  variance matrix corresponding to the full set of coefficients is returned.

`moptimize_result_Vtype(M)` returns a string scalar containing the default *vcetype*.

`moptimize_result_V_oim(M [, i])` returns the appropriate  $(ki + ci) \times (ki + ci)$  submatrix of the full variance matrix calculated as the inverse of the negative Hessian matrix (the observed information matrix). If  $i \geq .$  or argument *i* is omitted, the full  $K \times K$  variance matrix corresponding to the full set of coefficients is returned.

`moptimize_result_V_opg(M [, i])` returns the appropriate  $(ki + ci) \times (ki + ci)$  submatrix of the full variance matrix calculated as the inverse of the outer product of the gradients. If  $i \geq .$  or argument *i* is omitted, the full  $K \times K$  variance matrix corresponding to the full full set of coefficients is returned. If `moptimize_result_V_opg()` is used with evaluator types other than lf, lf\*, gf, or q, an appropriately dimensioned matrix of zeros is returned.

`moptimize_result_V_robust(M [, i])` returns the appropriate  $(ki + ci) \times (ki + ci)$  submatrix of the full variance matrix calculated via the sandwich estimator. If  $i \geq .$  or argument *i* is omitted, the full  $K \times K$  variance matrix corresponding to the full set of coefficients is returned. If `moptimize_result_V_robust()` is used with evaluator types other than lf, lf\*, gf, or q, an appropriately dimensioned matrix of zeros is returned.

`moptimize_result_iterations(M)` returns a real scalar containing the number of iterations performed.

`moptimize_result_converged(M)` returns a real scalar containing 1 if convergence was achieved and 0 otherwise.

`moptimize_result_iterationlog(M)` returns a real colvector containing the values of the objective function at the end of each iteration. Up to the last 20 iterations are returned, one to a row.

`moptimize_result_errorcode(M)` returns the real scalar containing the error code from the most recently run optimization or function evaluation. The error code is 0 if there are no errors. This function is useful only after `_moptimize()` or `_moptimize_evaluate()` because the nonunderscore versions aborted with error if there were problems.

`moptimize_result_errortext(M)` returns a string scalar containing the error text corresponding to `moptimize_result_errorcode()`.

`moptimize_result_returncode(M)` returns a real scalar containing the Stata return code corresponding to `moptimize_result_errorcode()`.

The following error codes and their corresponding Stata return codes are for `moptimize()` only. To see other [error codes](#) and their corresponding Stata return codes, see [\[M-5\] optimize\(\)](#).

Error code	Return code	Error text
400	1400	could not find feasible values
401	491	Stata program evaluator returned an error
402	198	views are required when the evaluator is a Stata program
403	198	Stata program evaluators require a touse variable

## Stata evaluators

The following function is useful only when your evaluator is a Stata program instead of a Mata function.

`moptimize_ado_cleanup(M)` removes all the global macros with the `ML_` prefix. A temporary weight variable is also dropped if weights were specified.

## Advanced functions

These functions are not really advanced, they are just seldomly used.

`moptimize_init_verbose(M, {"on"|"off"})` specifies whether error messages are to be displayed. The default is "on".

`moptimize_init_evaluations(M, {"off"|"on"})` specifies whether the system is to count the number of times the [evaluator](#) is called. The default is "off".

`moptimize_result_evaluations(M)` returns a  $1 \times 3$  real rowvector containing the number of times the [evaluator](#) was called, assuming `moptimize_init_evaluations()` was set on. Contents are the number of times called for the purposes of 1) calculating the objective function, 2) calculating the objective function and its first derivative, and 3) calculating the objective function and its first and second derivatives. If `moptimize_init_evaluations()` was set off, returned is (0,0,0).

## Syntax of evaluators

An *evaluator* is a program you write that calculates the value of the function being optimized and optionally calculates the function's first and second derivatives. The evaluator you write is called by the `moptimize()` functions.

There are five styles in which the evaluator can be written, known as types `lf`, `d`, `lf*`, `gf`, and `q`. *evaluator**type*, optionally specified in `moptimize_init_evaluator`*type*`()`, specifies the style in which the evaluator is written. Allowed values are

<i>evaluator</i> <i>type</i>	Description
"lf"	<i>function()</i> returns $N \times 1$ colvector value
"d0"	<i>function()</i> returns scalar value
"d1"	same as "d0" and returns gradient rowvector
"d2"	same as "d1" and returns Hessian matrix
"d1debug"	same as "d1" but checks gradient
"d2debug"	same as "d2" but checks gradient and Hessian
"lf0"	<i>function()</i> returns $N \times 1$ colvector value
"lf1"	same as "lf0" and return equation-level score matrix
"lf2"	same as "lf1" and returns Hessian matrix
"lf1debug"	same as "lf1" but checks gradient
"lf2debug"	same as "lf2" but checks gradient and Hessian
"gf0"	<i>function()</i> returns $N \times 1$ colvector value
"gf1"	same as "gf0" and returns score matrix
"gf2"	same as "gf1" and returns Hessian matrix
"gf1debug"	same as "gf1" but checks gradient
"gf2debug"	same as "gf2" but checks gradient and Hessian
"q0"	<i>function()</i> returns colvector value
"q1"	same as "q0" and returns score matrix
"q1debug"	same as "q1" but checks gradient

The default is "lf" if not set.  
"q" evaluators are used with technique "gn".  
Returned gradients are  $1 \times K$  rowvectors.  
Returned Hessians are  $K \times K$  matrices.

Examples of each of the evaluator types are outlined below.

You must tell `moptimize()` the identity and type of your evaluator, which you do by using the `moptimize_init_evaluator()` and `moptimize_init_evaluortype()` functions.

`moptimize_init_evaluator(M, &functionname())` sets the identity of the evaluator function that you write in Mata.

`moptimize_init_evaluator(M, "programname")` sets the identity of the evaluator program that you write in Stata.

`moptimize_init_evaluortype(M, evaluatortype)` informs `moptimize()` of the style of evaluator you have written. *evaluator**type* is a string scalar from the table above. The default is "lf".

`moptimize_init_negH(M, { "off" | "on" })` sets whether the evaluator you have written returns  $H$  or  $-H$ , the Hessian or the negative of the Hessian, if it returns a Hessian at all. This is for backward compatibility with prior versions of Stata's `ml` command (see [R] `ml`). Modern evaluators return  $H$ . The default is "off".



## Syntax of type lf evaluators

`lfeval(M, b,  $\overline{fv}$ ):`

*inputs:*

*M*: problem definition  
*b*: coefficient vector

*outputs:*

*fv*:  $N \times 1$ ,  $N = \#$  of observations

Notes:

1. The objective function is  $f() = \text{colsum}(fv)$ .
2. In the case where  $f()$  is a log-likelihood function, the values of the log likelihood must be summable over the observations.
3. For use with any [technique](#) except `gn`.
4. May be used with robust, clustering, and survey.
5. Returns *fv* containing missing ( $fv = .$ ) if evaluation is not possible.

## Syntax of type d evaluators

`deval(M, todo, b,  $\overline{fv}$ ,  $\overline{g}$ ,  $\overline{H}$ ):`

*inputs:*

*M*: problem definition  
*todo*: real scalar containing 0, 1, or 2  
*b*: coefficient vector

*outputs:*

*fv*: real scalar  
*g*:  $1 \times K$ , gradients,  $K = \#$  of coefficients  
*H*:  $K \times K$ , Hessian

Notes:

1. The objective function is  $f() = fv$ .
2. For use with any log-likelihood function, or any function.
3. For use with any [technique](#) except `gn` and `bhhh`.
4. Cannot be used with robust, clustering, or survey.
5. `deval()` must always fill in *fv*, and fill in *g* if  $todo \geq 1$ , and fill in *H* if  $todo = 2$ . For type `d0`, *todo* will always be 0. For type `d1` and `d1debug`, *todo* will be 0 or 1. For type `d2` and `d2debug`, *todo* will be 0, 1, or 2.
6. Returns  $fv = .$  if evaluation is not possible.

## Syntax of type lf\* evaluators

`lfeval(M, todo, b,  $\overline{fv}$ ,  $\overline{S}$ ,  $\overline{H}$ ):`

*inputs:*

*M*: problem definition  
*todo*: real scalar containing 0, 1, or 2  
*b*: coefficient vector

*outputs:*

*fv*:  $N \times 1$ ,  $N = \#$  of observations  
*S*:  $N \times m$ , scores,  $m = \#$  of equations (parameters)  
*H*:  $K \times K$ , Hessian,  $K = \#$  of coefficients

Notes:

1. The objective function is  $f() = \text{colsum}(fv)$ .
2. Type `lf*` is a variation of type `lf` that allows the user to supply analytic derivatives. Although `lf*` could be used with an arbitrary function, it is intended for use when  $f()$  is a log-likelihood function and the log-likelihood values are summable over the observations.
3. For use with any [technique](#) except `gn`.
4. May be used with robust, clustering, and survey.
5. Always returns *fv*, returns *S* if *todo*  $\geq 1$ , and returns *H* if *todo* = 2. For type `lf0`, *todo* will always be 0. For type `lf1` and `lf1debug`, *todo* will be 0 or 1. For type `lf2` and `lf2debug`, *todo* will be 0, 1, or 2.
6. Returns *fv* containing missing ( $fv = .$ ) if evaluation is not possible.

## Syntax of type gf evaluators

`gfeval(M, todo, b,  $\overline{fv}$ ,  $\overline{S}$ ,  $\overline{H}$ ):`

*inputs:*

*M*: problem definition  
*todo*: real scalar containing 0, 1, or 2  
*b*: coefficient vector

*outputs:*

*fv*:  $L \times 1$ , values,  $L = \#$  of independent elements  
*S*:  $L \times K$ , scores,  $K = \#$  of coefficients  
*H*:  $K \times K$ , Hessian

Notes:

1. The objective function is  $f() = \text{colsum}(fv)$ .
2. Type `gf` is a variation on type `lf*` that relaxes the requirement that the log-likelihood function be summable over the observations. `gf` is especially useful for fitting panel-data models with [technique](#) `bhhh`. Then  $L$  is the number of panels.
3. For use with any `gf` is especially useful for fitting panel-data models with except `gn`.
4. May be used with robust, clustering, and survey.
5. Always returns  $fv$ , returns  $S$  if  $todo \geq 1$ , and returns  $H$  if  $todo = 2$ . For type `gf0`,  $todo$  will always be 0. For type `gf1` and `gf1debug`,  $todo$  will be 0 or 1. For type `gf2` and `gf2debug`,  $todo$  will be 0, 1, or 2.
6. Returns  $fv = .$  if evaluation is not possible.

## Syntax of type q evaluators

`qeval(M, todo, b,  $\bar{r}$ ,  $\bar{S}$ )`

*inputs:*

*M*: problem definition  
*todo*: real scalar containing 0 or 1  
*b*: coefficient vector

*outputs:*

*r*:  $L \times 1$  of independent elements  
*S*:  $L \times m$ , scores,  $m = \#$  of parameters

Notes:

1. Type `q` is for quadratic optimization. The objective function is  $f() = r'Wr$ , where  $r$  is returned by `qeval()` and  $W$  has been previously set by using `moptimize_init_gnweightmatrix()`, described below.
2. For use only with [techniques](#) `gn` and `nm`.
3. Always returns  $r$  and returns  $S$  if  $todo = 1$ . For type `q0`,  $todo$  will always be 0. For type `q1` and `q1debug`,  $todo$  will be 0 or 1. There is no type `q2`.
4. Returns  $r$  containing missing, or  $r = .$  if evaluation is not possible.

Use `moptimize_init_gnweightmatrix()` during initialization to set matrix  $W$ .

`moptimize_init_gnweightmatrix(M, W)` sets real matrix  $W$ :  $L \times L$ , which is used only by type `q` evaluators. The objective function is  $r'Wr$ . If  $W$  is not set and if observation weights  $w$  are set by using `moptimize_init_weight()`, then  $W = \text{diag}(w)$ . If  $w$  is not set, then  $W$  is the identity matrix.

`moptimize()` does not produce a robust VCE when you set  $W$  with `moptimize_init_gnweight()`.

## Passing extra information to evaluators

In addition to the arguments the evaluator receives, you may arrange that extra information be sent to the evaluator. Specify the extra information to be sent by using `moptimize_init_userinfo()`.

`moptimize_init_userinfo(M, l, Z)` specifies that the *l*th piece of extra information is *Z*. *l* is a real scalar. The first piece of extra information should be 1; the second piece, 2; and so on. *Z* can be anything. No copy of *Z* is made.

`moptimize_init_nuserinfo(M, n_user)` specifies the total number of extra pieces of information to be sent. Setting *n\_user* is optional; it will be automatically determined from the `moptimize_init_userinfo()` calls you issue.

Inside your evaluator, you access the information by using `moptimize_util_userinfo()`.

`moptimize_util_userinfo(M, l)` returns the *Z* set by `moptimize_init_userinfo()`.

## Utility functions

There are various utility functions that are helpful in writing [evaluators](#) and in processing results returned by the `moptimize_result_*`() functions.

The first set of utility functions are useful in writing evaluators, and the first set return results that all evaluators need.

`moptimize_util_depvar(M, j)` returns an  $N_j \times 1$  colvector containing the values of the *j*th dependent variable, the values set by `moptimize_init_depvar(M, j, ...)`.

`moptimize_util_xb(M, b, i)` returns the  $N_i \times 1$  colvector containing the value of the *i*th [parameter](#), which is usually  $X_i \times b_i' :+ b_0i$ , but might be as complicated as  $X_i \times b_i' + o_i + \ln(ti) :+ b_0i$ .

Once the inputs of an evaluator have been processed, the following functions assist in making the calculations required of evaluators.

`moptimize_util_sum(M, v)` returns the “sum” of colvector *v*. This function is for use in evaluators that require you to return an overall objective function value rather than observation-by-observation results. Usually, `moptimize_util_sum()` returns `sum(v)`, but in cases where you have specified a weight by using `moptimize_init_weight()` or there is an implied weight due to use of `moptimize_init_svy()`, the appropriately weighted sum is returned. Use `moptimize_util_sum()` to sum log-likelihood values.

`moptimize_util_vecsum(M, i, s, value)` is like `moptimize_util_sum()`, but for use with gradients. The gradient is defined as the vector of partial derivatives of *f*() with respect to the coefficients *bi*. Some evaluator types require that your evaluator be able to return this vector. Nonetheless, it is usually easier to write your evaluator in terms of parameters rather than coefficients, and this function handles the mapping of parameter gradients to the required coefficient gradients.

Input *s* is an  $N_i \times 1$  colvector containing  $df/dpi$  for each observation.  $df/dpi$  is the partial derivative of the objective function, but with respect to the *i*th parameter rather than the *i*th set of coefficients. `moptimize_util_vecsum()` takes *s* and returns the  $1 \times (ki + ci)$  summed gradient. Also weights, if any, are factored into the calculation.

If you have more than one equation, you will need to call `moptimize_util_vecsum()`  $m$  times, once for each equation, and then concatenate the individual results into one vector.

*value* plays no role in `moptimize_util_vecsum()`'s calculations. *value*, however, should be specified as the result obtained from `moptimize_util_sum()`. If that is inconvenient, make *value* any nonmissing value. If the calculation from parameter space to vector space cannot be performed, or if your original parameter space derivatives have any missing values, *value* will be changed to missing. Remember, when a calculation cannot be made, the evaluator is to return a missing value for the objective function. Thus storing the value of the objective function in *value* ensures that your evaluator will return missing if it is supposed to.

`moptimize_util_matsum(M, i, i2, s, value)` is similar to `moptimize_util_vecsum()`, but for Hessians (matrix of second derivatives).

Input *s* is an  $N_i \times 1$  colvector containing  $d^2f/dpidpi^2$  for each observation. `moptimize_util_matsum()` returns the  $(ki + ci) \times (ki + ci)$  summed Hessian. Also weights, if any, are factored into the calculation.

If you have  $m > 1$  equations, you will need to call `moptimize_util_matsum()`  $m \times (m+1)/2$  times and then join the results into one symmetric matrix.

*value* plays no role in the calculation and works the same way it does in `moptimize_util_vecsum()`.

`moptimize_util_matbysum()` is an added helper for making `moptimize_util_matsum()` calculations in cases where you have panel data and the log-likelihood function's values exists only at the panel level. `moptimize_util_matbysum(M, i, a, b, value)` is for making diagonal calculations and `moptimize_util_matbysum(M, i, i2, a, b, c, value)` is for making off-diagonal calculations.

This is an advanced topic; see [Gould, Pitblado, and Poi \(2010, 136–138\)](#) for a full description of it. In applying the chain rule to translate results from parameter space to coefficient space, `moptimize_util_matsum()` can be used to make some of the calculations, and `moptimize_util_matbysum()` can be used to make the rest. *value* plays no role and works just as it did in the other helper functions. `moptimize_util_matbysum()` is for use sometimes when *by* has been set, which is done via `moptimize_init_by(M, by)`. `moptimize_util_matbysum()` is never required unless *by* has been set.

The formula implemented in `moptimize_util_matbysum(M, i, a, b, value)` is

$$\sum_{j=1}^N \left( \sum_{t=1}^{T_j} a_{jt} \right) \left( \sum_{t=1}^{T_j} b_{jt} \mathbf{x}'_{1jt} \right) \left( \sum_{t=1}^{T_j} b_{jt} \mathbf{x}_{1jt} \right)$$

The formula implemented in `moptimize_util_matbysum(M, i, i2, a, b, c, value)` is

$$\sum_{j=1}^N \left( \sum_{t=1}^{T_j} a_{jt} \right) \left( \sum_{t=1}^{T_j} b_{jt} \mathbf{x}'_{1jt} \right) \left( \sum_{t=1}^{T_j} c_{jt} \mathbf{x}_{2jt} \right)$$

`moptimize_util_by()` returns a pointer to the vector of group identifiers that were set using `moptimize_init_by()`. This vector can be used with `panelsetup()` to perform panel level calculations.

The other utility functions are useful inside or outside of evaluators. One of the more useful is `moptimize_util_eq_indices()`, which allows two or three arguments.

`moptimize_util_eq_indices(M, i)` returns a  $1 \times 2$  vector that can be used with [range subscripts](#) to extract the portion relevant for the *i*th equation from any  $1 \times K$  vector, that is, from any vector conformable with the [full coefficient vector](#).

`moptimize_util_eq_indices(M, i, i2)` returns a  $2 \times 2$  matrix that can be used with [range subscripts](#) to extract the portion relevant for the *i*th and *i2*th equations from any  $K \times K$  matrix, that is, from any matrix with rows and columns conformable with the full variance matrix.

For instance, let *b* be the  $1 \times K$  full coefficient vector, perhaps obtained by being passed into an evaluator, or perhaps obtained from *b* = `moptimize_result_coefs(M)`. Then `b[moptimize_util_eq_indices(M, i)]` is the  $1 \times (ki + ci)$  vector of coefficients for the *i*th equation.

Let *V* be the  $K \times K$  full variance matrix obtained by *V* = `moptimize_result_V(M)`. Then `V[moptimize_util_eq_indices(M, i, i)]` is the  $(ki + ci) \times (ki + ci)$  variance matrix for the *i*th equation. `V[moptimize_util_eq_indices(M, i, j)]` is the  $(ki + ci) \times (kj + cj)$  covariance matrix between the *i*th and *j*th equations.

Finally, there is one more utility function that may help when you become confused: `moptimize_query()`.

`moptimize_query(M)` displays in readable form everything you have set via the `moptimize_init_*` functions, along with the status of the system.

## Description

The `moptimize()` functions find coefficients ( $\mathbf{b}_1, \mathbf{b}_2, \dots, \mathbf{b}_m$ ) that maximize or minimize  $f(\mathbf{p}_1, \mathbf{p}_2, \dots, \mathbf{p}_m)$ , where  $\mathbf{p}_i = \mathbf{X}_i \times \mathbf{b}_i'$ , a linear combination of  $\mathbf{b}_i$  and the data. The user of `moptimize()` writes a Mata function or Stata program to evaluate  $f(\mathbf{p}_1, \mathbf{p}_2, \dots, \mathbf{p}_m)$ . The data can be in Mata matrices or in the Stata dataset currently residing in memory.

`moptimize()` is especially useful for obtaining solutions for maximum likelihood models, minimum chi-squared models, minimum squared-residual models, and the like.

## Remarks

Remarks are presented under the following headings:

- Relationship of `moptimize()` to Stata's `ml` and to Mata's `optimize()`*
- Mathematical statement of the `moptimize()` problem*
- Filling in `moptimize()` from the mathematical statement*
- The type `lf` evaluator*
- The type `d`, `lf*`, `gf`, and `q` evaluators*
- Example using type `d`*
- Example using type `lf*`*

## Relationship of moptimize() to Stata's ml and to Mata's optimize()

moptimize() is Mata's and Stata's premier optimization routine. This is the routine used by most of the official optimization-based estimators implemented in Stata.

That said, Stata's ml command—see [R] [ml](#)—provides most of the capabilities of Mata's moptimize(), and ml is easier to use. In fact, ml uses moptimize() to perform the optimization, and ml amounts to little more than a shell providing a friendlier interface. If you have a maximum likelihood model you wish to fit, we recommend you use ml instead of moptimize(). Use moptimize() when you need or want to work in the Mata environment, or when you wish to implement a specialized system for fitting a class of models.

Also make note of Mata's optimize() function; see [M-5] [optimize\(\)](#). moptimize() finds coefficients  $(\mathbf{b}_1, \mathbf{b}_2, \dots, \mathbf{b}_m)$  that maximize or minimize  $f(\mathbf{p}_1, \mathbf{p}_2, \dots, \mathbf{p}_m)$ , where  $\mathbf{p}_i = \mathbf{X}_i \times \mathbf{b}_i$ . optimize() handles a simplified form of the problem, namely, finding constant  $(p_1, p_2, \dots, p_m)$  that maximizes or minimizes  $f()$ . moptimize() is the appropriate routine for fitting a Weibull model, but if all you need to estimate are the fixed parameters of the Weibull distribution for some population, moptimize() is overkill and optimize() will prove easier to use.

These three routines are all related. Stata's ml uses moptimize() to do the numerical work. moptimize(), in turn, uses optimize() to perform certain calculations, including the search for parameters. There is nothing inferior about optimize() except that it cannot efficiently deal with models in which parameters are given by linear combinations of coefficients and data.

## Mathematical statement of the moptimize() problem

We mathematically describe the problem moptimize() solves not merely to fix notation and ease communication, but also because there is a one-to-one correspondence between the mathematical notation and the moptimize\*() functions. Simply writing your problem in the following notation makes obvious the moptimize\*() functions you need and what their arguments should be.

In what follows, we are going to simplify the mathematics a little. For instance, we are about to claim  $\mathbf{p}_i = \mathbf{X}_i \times \mathbf{b}_i :+ c_i$ , when in the syntax section, you will see that  $\mathbf{p}_i = \mathbf{X}_i \times \mathbf{b}_i + \mathbf{o}_i + \ln(\mathbf{t}_i) :+ c_i$ . Here we omit  $\mathbf{o}_i$  and  $\ln(\mathbf{t}_i)$  because they are seldom used. We will omit some other details, too. The statement of the problem under [Syntax](#), above, is the full and accurate statement. We will also use typefaces a little differently. In the syntax section, we use italics following programming convention. In what follows, we will use boldface for matrices and vectors, and italics for scalars so that you can follow the math more easily. So in this section, we will write  $\mathbf{b}_i$ , whereas under syntax we would write *b<sub>i</sub>*; regardless of typeface, they mean the same thing.

Function moptimize() finds coefficients

$$\mathbf{b} = ((\mathbf{b}_1, c_1), (\mathbf{b}_2, c_2), \dots, (\mathbf{b}_m, c_m))$$

where

$$\begin{array}{llll} \mathbf{b}_1: 1 \times k_1, & \mathbf{b}_2: 1 \times k_2, & \dots, & \mathbf{b}_m: 1 \times k_m \\ c_1: 1 \times 1, & c_2: 1 \times 1, & \dots, & c_m: 1 \times 1 \end{array}$$

that maximize or minimize function

$$f(\mathbf{p}_1, \mathbf{p}_2, \dots, \mathbf{p}_m; \mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_D)$$

where

$$\begin{aligned} \mathbf{p}_1 &= \mathbf{X}_1 \times \mathbf{b}'_1 :+ c_1, & \mathbf{X}_1 &: N_1 \times k_1 \\ \mathbf{p}_2 &= \mathbf{X}_2 \times \mathbf{b}'_2 :+ c_2, & \mathbf{X}_2 &: N_2 \times k_2 \\ &\vdots & \\ \mathbf{p}_m &= \mathbf{X}_m \times \mathbf{b}'_m :+ c_m, & \mathbf{X}_m &: N_m \times k_m \end{aligned}$$

and where  $\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_D$  are of arbitrary dimension.

Usually,  $N_1 = N_2 = \dots = N_m$ , and the model is said to be fit on data of  $N$  observations. Similarly, column vectors  $\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_D$  are usually called dependent variables, and each is also of  $N$  observations.

As an example, let's write the maximum likelihood estimator for linear regression in the above notation. We begin by stating the problem in the usual way, but in Mata-ish notation:

Given data  $\mathbf{y}$ :  $N \times 1$  and  $\mathbf{X}$ :  $N \times k$ , obtain  $((\mathbf{b}, c), s^2)$  to fit

$$\mathbf{y} = \mathbf{X} \times \mathbf{b}' :+ c + \mathbf{u}$$

where the elements of  $\mathbf{u}$  are distributed  $N(0, s^2)$ . The log-likelihood function is

$$\ln L = \sum_j \ln(\text{normalden}(\mathbf{y}_j - (\mathbf{X}_j \times \mathbf{b}' :+ c), 0, \text{sqrt}(s^2)))$$

where `normalden( $x$ ,  $mean$ ,  $sd$ )` returns the density at  $x$  of the Gaussian normal with the specified mean and standard deviation; see [M-5] `normal()`.

The above is a two-parameter or, equivalently, two-equation model in `moptimize()` jargon. There may be many coefficients, but the likelihood function can be written in terms of two parameters, namely  $\mathbf{p}_1 = \mathbf{X} \times \mathbf{b}' :+ c$  and  $\mathbf{p}_2 = s^2$ . Here is the problem stated in the `moptimize()` notation:

Find coefficients

$$\mathbf{b} = ((\mathbf{b}_1, c_1), (c_2))$$

where

$$\begin{aligned} \mathbf{b}_1 &: 1 \times k \\ c_1 &: 1 \times 1, & c_2 &: 1 \times 1 \end{aligned}$$

that maximize

$$f(\mathbf{p}_1, \mathbf{p}_2; \mathbf{y}) = \sum \ln(\text{normalden}(\mathbf{y} - \mathbf{p}_1, 0, \text{sqrt}(\mathbf{p}_2)))$$

where

$$\begin{aligned} \mathbf{p}_1 &= \mathbf{X} \times \mathbf{b}'_1 :+ c_1, & \mathbf{X} &: N \times k \\ \mathbf{p}_2 &= c_2 \end{aligned}$$

and where  $y$  is  $N \times 1$ .



Notice that, in this notation, the regression coefficients  $(\mathbf{b}_1, c_1)$  play a secondary role, namely, to determine  $\mathbf{p}_1$ . That is, the function,  $f()$ , to be optimized—a log-likelihood function here—is written in terms of  $\mathbf{p}_1$  and  $\mathbf{p}_2$ . The program you will write to evaluate  $f()$  will be written in terms of  $\mathbf{p}_1$  and  $\mathbf{p}_2$ , thus abstracting from the particular regression model being fit. Whether the regression is mpg on weight or log income on age, education, and experience, your program to calculate  $f()$  will remain unchanged. All that will change are the definitions of  $\mathbf{y}$  and  $\mathbf{X}$ , which you will communicate to `moptimize()` separately.

There is another advantage to this arrangement. We can trivially generalize linear regression without writing new code. Note that the variance  $s^2$  is given by  $\mathbf{p}_2$ , and currently, we have  $\mathbf{p}_2 = c_2$ , that is, a constant. `moptimize()` allows parameters to be constant, but it equally allows them to be given by a linear combination. Thus rather than defining  $\mathbf{p}_2 = c_2$ , we could define  $\mathbf{p}_2 = \mathbf{X}_2 \times \mathbf{b}'_2 :+ c_2$ . If we did that, we would have a second linear equation that allowed the variance to vary observation by observation. As far as `moptimize()` is concerned, that problem is the same as the original problem.

## Filling in `moptimize()` from the mathematical statement

The mathematical statement of our sample problem is the following:

Find coefficients

$$\mathbf{b} = ((\mathbf{b}_1, c_1), (c_2))$$

$$\begin{array}{ll} \mathbf{b}_1: 1 \times k \\ c_1: 1 \times 1, & c_2: 1 \times 1 \end{array}$$

that maximize

$$f(\mathbf{p}_1, \mathbf{p}_2; \mathbf{y}) = \sum \ln(\text{normalden}(\mathbf{y} - \mathbf{p}_1, 0, \text{sqrt}(\mathbf{p}_2)))$$

where

$$\begin{array}{ll} \mathbf{p}_1 = \mathbf{X} \times \mathbf{b}'_1 :+ c_1, & \mathbf{X} : N \times k \\ \mathbf{p}_2 = c_2 \end{array}$$

and where  $\mathbf{y}$  is  $N \times 1$ .

The corresponding code to perform the optimization is

```
. sysuse auto
. mata:
: function linregeval(transmorphic M, real rowvector b,
    real colvector lnf)
{
    real colvector  p1, p2
    real colvector  y1
    p1 = moptimize_util_xb(M, b, 1)
    p2 = moptimize_util_xb(M, b, 2)
    y1 = moptimize_util_depvar(M, 1)
    lnf = ln(normalden(y1:-p1, 0, sqrt(p2)))
}
: M = moptimize_init()
: moptimize_init_evaluator(M, &linregeval())
: moptimize_init_depvar(M, 1, "mpg")
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
: moptimize_init_eq_indepvars(M, 2, "")
: moptimize(M)
: moptimize_result_display(M)
```

Here is the result of running the above code:

```
. sysuse auto
(1978 Automobile Data)
. mata:
----- mata (type end to exit) -----
: function linregeval(transmorphic M, real rowvector b, real colvector lnf)
> {
>     real colvector  p1, p2
>     real colvector  y1
>
>     p1 = moptimize_util_xb(M, b, 1)
>     p2 = moptimize_util_xb(M, b, 2)
>     y1 = moptimize_util_depvar(M, 1)
>
>     lnf = ln(normalden(y1:-p1, 0, sqrt(p2)))
> }
: M = moptimize_init()
: moptimize_init_evaluator(M, &linregeval())
: moptimize_init_depvar(M, 1, "mpg")
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
: moptimize_init_eq_indepvars(M, 2, "")
```

```

: moptimize(M)
initial:      f(p) =      -<inf>  (could not be evaluated)
feasible:     f(p) = -12949.708
rescale:      f(p) = -243.04355
rescale eq:   f(p) = -236.58999
Iteration 0:   f(p) = -236.58999  (not concave)
Iteration 1:   f(p) = -227.46719
Iteration 2:   f(p) = -205.62547  (backed up)
Iteration 3:   f(p) = -195.58046
Iteration 4:   f(p) = -194.20252
Iteration 5:   f(p) = -194.18313
Iteration 6:   f(p) = -194.18306
Iteration 7:   f(p) = -194.18306
: moptimize_result_display(M)

```

Number of obs = 74

	mpg	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
eq1	weight	-.0065879	.0006241	-10.56	0.000	-.007811	-.0053647
	foreign	-1.650029	1.053958	-1.57	0.117	-3.715748	.4156903
	_cons	41.6797	2.121197	19.65	0.000	37.52223	45.83717
eq2	_cons	11.13746	1.830987	6.08	0.000	7.548789	14.72613

## The type II evaluator

Let's now interpret the code we wrote, which was

```

: function linregeval(transmorphic M, real rowvector b,
                      real colvector lnf)
{
    real colvector  p1, p2
    real colvector  y1
    p1 = moptimize_util_xb(M, b, 1)
    p2 = moptimize_util_xb(M, b, 2)
    y1 = moptimize_util_depvar(M, 1)
    lnf = ln(normalden(y1:-p1, 0, sqrt(p2)))
}

: M = moptimize_init()
: moptimize_init_evaluator(M, &linregeval())
: moptimize_init_depvar(M, 1, "mpg")
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
: moptimize_init_eq_indepvars(M, 2, "")
: moptimize(M)
: moptimize_result_display(M)

```

We first defined the function to evaluate our likelihood function—we named the function `linregeval()`. The name was of our choosing. After that, we began an optimization problem by typing `M = moptimize_init()`, described the problem with `moptimize_init_*`() functions, performed the optimization by typing `moptimize()`, and displayed results by using `moptimize_result_display()`.

Function `linregeval()` is an example of a type `lf` evaluator. There are several different evaluator types, including `d0`, `d1`, `d2`, through `q1`. Of all of them, type `lf` is the easiest to use and is the one `moptimize()` uses unless we tell it differently. What makes `lf` easy is that we need only calculate the likelihood function; we are not required to calculate its derivatives. A description of `lf` appears under the heading *Syntax of type lf evaluators* under *Syntax* above.

In the syntax diagrams, you will see that type `lf` evaluators receive three arguments,  $M$ ,  $b$ , and  $fv$ , although in `linregeval()`, we decided to call them `M`, `b`, and `lnf`. The first two arguments are inputs, and your evaluator is expected to fill in the third argument with observation-by-observation values of the log-likelihood function.

The input arguments are  $M$  and  $b$ .  $M$  is the problem handle, which we have not explained yet. Basically, all evaluators receive  $M$  as the first argument and are expected to pass  $M$  to any `moptimize*()` subroutines that they call.  $M$  in fact contains all the details of the optimization problem. The second argument,  $b$ , is the entire coefficient vector, which in the `linregeval()` case will be all the coefficients of our regression, the constant (intercept), and the variance of the residual. Those details are unimportant. Instead, your evaluator will pass  $M$  and  $b$  to `moptimize()` utility programs that will give you back what you need.

Using one of those utilities is the first action our `linregeval()` evaluator performs:

```
p1 = moptimize_util_xb(M, b, 1)
```

That returns observation-by-observation values of the first parameter, namely,  $X \times b_1 :+ c_1$ . `moptimize_util_xb(x, b, 1)` returns the first parameter because the last argument specified is 1. We obtained the second parameter similarly:

```
p2 = moptimize_util_xb(M, b, 2)
```

To evaluate the likelihood function, we also need the dependent variable. Another `moptimize*()` utility returns that to us:

```
y1 = moptimize_util_depvar(M, 1)
```

Having `p1`, `p2`, and `y1`, we are ready to fill in the log-likelihood values:

```
lnf = ln(normalden(y1:-p1, 0, sqrt(p2)))
```

For a type `lf` evaluator, you are to return observation-by-observation values of the log-likelihood function; `moptimize()` itself will sum them to obtain the overall log likelihood. That is exactly what the line `lnf = ln(normalden(y1:-p1, 0, sqrt(p2)))` did. Note that `y1` is  $N \times 1$ , `p1` is  $N \times 1$ , and `p2` is  $N \times 1$ , so the `lnf` result we calculate is also  $N \times 1$ . Some of the other evaluator types are expected to return a scalar equal to the overall value of the function.

With the evaluator defined, we can estimate a linear regression by typing

```
: M = moptimize_init()
: moptimize_init_evaluator(M, &linregeval())
: moptimize_init_depvar(M, 1, "mpg")
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
: moptimize_init_eq_indepvars(M, 2, "")
: moptimize(M)
: moptimize_result_display(M)
```

All estimation problems begin with

```
M = moptimize_init()
```

The returned value `M` is called a problem handle, and from that point on, you will pass `M` to every other `moptimize()` function you call. `M` contains the details of your problem. If you were to list `M`, you would see something like

```
: M
0x15369a
```

`0x15369a` is in fact the address where all those details are stored. Exactly how `M` works does not matter, but it is important that you understand what `M` is. `M` is your problem. In a more complicated problem, you might need to perform nested optimizations. You might have one optimization problem, and right in the middle of it, even right in the middle of evaluating its log-likelihood function, you might need to set up and solve another optimization problem. You can do that. The first problem you would set up as `M1 = moptimize_init()`. The second problem you would set up as `M2 = moptimize_init()`. `moptimize()` would not confuse the two problems because it would know to which problem you were referring by whether you used `M1` or `M2` as the argument of the `moptimize()` functions. As another example, you might have one optimization problem, `M = moptimize_init()`, and halfway through it, decide you want to try something wild. You could code `M2 = M`, thus making a copy of the problem, use the `moptimize*()` functions with `M2`, and all the while your original problem would remain undisturbed.

Having obtained a problem handle, that is, having coded `M = moptimize_init()`, you now need to fill in the details of your problem. You do that with the `moptimize_init_*()` functions. The order in which you do this does not matter. We first informed `moptimize()` of the identity of the evaluator function:

```
: moptimize_init_evaluator(M, &linregeval())
```

We must also inform `moptimize()` as to the type of evaluator function `linregeval()` is, which we could do by coding

```
: moptimize_init_evaluortype(M, "lf")
```

We did not bother, however, because type `lf` is the default.

After that, we need to inform `moptimize()` as to the identity of the dependent variables:

```
: moptimize_init_depvar(M, 1, "mpg")
```

Dependent variables play no special role in `moptimize()`; they are merely something that are remembered so that they can be passed to the evaluator function that we write. One problem might have no dependent variables and another might have lots of them. `moptimize_init_depvar(M, i, y)`'s second argument specifies which dependent variable is being set. There is no requirement that the number of dependent variables match the number of equations. In the linear regression case, we have one dependent variable and two equations.

Next we set the independent variables, or equivalently, the mapping of coefficients, into parameters. When we code

```
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
```

we are stating that there is a parameter,  $\mathbf{p}_1 = \mathbf{X}_1 \times \mathbf{b}_1 :+ c_1$ , and that  $\mathbf{X}_1 = (\text{weight}, \text{foreign})$ . Thus  $\mathbf{b}_1$  contains two coefficients, that is,  $\mathbf{p}_1 = (\text{weight}, \text{foreign}) \times (b_{11}, b_{12})' :+ c_1$ . Actually, we have not yet specified whether there is a constant,  $c_1$ , on the end, but if we do

not specify otherwise, the constant will be included. If we want to suppress the constant, after coding `moptimize_init_eq_indepvars(M, 1, "weight foreign")`, we would code `moptimize_init_eq_cons(M, 1, "off")`. The 1 says first equation, and the "off" says to turn the constant off.

As an aside, we coded `moptimize_init_eq_indepvars(M, 1, "weight foreign")` and so specified that the independent variables were the Stata variables `weight` and `foreign`, but the independent variables do not have to be in Stata. If we had a  $74 \times 2$  matrix named `data` in Mata that we wanted to use, we would have coded `moptimize_init_eq_indepvars(M, 1, data)`.

To define the second parameter, we code

```
: moptimize_init_eq_indepvars(M, 2, "")
```

Thus we are stating that there is a parameter,  $\mathbf{p}_2 = \mathbf{X}_2 \times \mathbf{b}_2 :+ c_2$ , and that  $\mathbf{X}_2$  does not exist, leaving  $\mathbf{p}_2 = c_2$ , meaning that the second parameter is a constant.

Our problem defined, we code

```
: moptimize(M)
```

to obtain the solution, and we code

```
: moptimize_result_display(M)
```

to see the results. There are many `moptimize_result_*`() functions for use after the solution is obtained.

## The type `d`, `lf`\*, `gf`, and `q` evaluators

Above we wrote our evaluator function in the style of type `lf`. `moptimize()` provides four other evaluator types—called types `d`, `lf`\*, `gf`, and `q`—and each have their uses.

Using type `lf` above, we were required to calculate the observation-by-observation log likelihoods and that was all. Using another type of evaluator, say, type `d`, we are required to calculate the overall log likelihood, and optionally, its first derivatives, and optionally, its second derivatives. The corresponding evaluator types are called `d0`, `d1`, and `d2`. Type `d` is better than type `lf` because if we do calculate the derivatives, then `moptimize()` can execute more quickly and it can produce a slightly more accurate result (more accurate because numerical derivatives are not involved). These speed and accuracy gains justify type `d1` and `d2`, but what about type `d0`? For many optimization problems, type `d0` is redundant and amounts to nothing more than a slight variation on type `lf`. In these cases, type `d0`'s justification is that if we want to write a type `d1` or type `d2` evaluator, then it is usually easiest to start by writing a type `d0` evaluator. Make that work, and then add to the code to convert our type `d0` evaluator into a type `d1` evaluator; make that work, and then, if we are going all the way to type `d2`, add the code to convert our type `d1` evaluator into a type `d2` evaluator.

For other optimization problems, however, there is a substantive reason for type `d0`'s existence. Type `lf` requires observation-by-observation values of the log-likelihood function, and for some likelihood functions, those simply do not exist. Think of a panel-data model. There may be observations within each of the panels, but there is no corresponding log-likelihood value for each of them. The log-likelihood function is defined only across the entire panel. Type `lf` cannot handle problems like that. Type `d0` can.

That makes type `d0` seem to be a strict improvement on type `lf`. Type `d0` can handle any problem that type `lf` can handle, and it can handle other problems to boot. Where both can handle the problem, the only extra work to use type `d0` is that we must sum the individual values we produce, and that is not difficult. Type `lf`, however, has other advantages. If you write a type `lf` evaluator, then without writing another line of code, you can obtain the robust estimates of variance, adjust for clustering, account for survey design effects, and more. `moptimize()` can do all that because it has the results of an observation-by-observation calculation. `moptimize()` can break into the assembly of those observation-by-observation results and modify how that is done. `moptimize()` cannot do that for `d0`.

So there are advantages both ways.

Another provided evaluator type is type `lf*`. Type `lf*` is a variation on type `lf`. It also comes in the subflavors `lf0`, `lf1`, and `lf2`. Type `lf*` allows you to make observation-level derivative calculations, which means that results can be obtained more quickly and more accurately. Type `lf*` is designed to always work where `lf` is appropriate, which means panel-data estimators are excluded. In return, it provides all the ancillary features provided by type `lf`, meaning that robust standard errors, clustering, and survey-data adjustments are available. You write the evaluator function in a slightly different style when you use type `lf*` rather than type `d`.

Type `gf` is a variation on type `lf*` that relaxes the requirement that the log-likelihood function be summable over the observations. Thus type `gf` can work with panel-data models and resurrect the features of robust standard errors, clustering, and survey-data adjustments. Type `gf` evaluators, however, are more difficult to write than type `lf*` evaluators.

Type `q` is for the special case of quadratic optimization. You either need it, and then only type `q` will do, or you do not.

## Example using type `d`

Let's return to our linear regression maximum-likelihood estimator. To remind you, this is a two-parameter model, and the log-likelihood function is

$$f(\mathbf{p}_1, \mathbf{p}_2; \mathbf{y}) = \sum \ln(\text{normalden}(\mathbf{y} - \mathbf{p}_1, 0, \text{sqrt}(\mathbf{p}_2)))$$

This time, however, we are going to parameterize the variance parameter  $\mathbf{p}_2$  as the log of the standard deviation, so we will write

$$f(\mathbf{p}_1, \mathbf{p}_2; \mathbf{y}) = \sum \ln(\text{normalden}(\mathbf{y} - \mathbf{p}_1, 0, \exp(\mathbf{p}_2)))$$

It does not make any conceptual difference which parameterization we use, but the log parameterization converges a little more quickly, and the derivatives are easier, too. We are going to implement a type `d2` evaluator for this function. To save you from pulling out pencil and paper, let's tell you the derivatives:

$$\begin{aligned} df/d\mathbf{p}_1 &= \mathbf{z}:/\mathbf{s} \\ df/d\mathbf{p}_2 &= \mathbf{z}:^2:-1 \\ d^2f/d\mathbf{p}_1^2 &= -1:/\mathbf{s}:^2 \\ d^2f/d\mathbf{p}_2^2 &= -2 \times \mathbf{z}:^2 \\ d^2f/d\mathbf{p}d\mathbf{p}_2 &= -2 \times \mathbf{z}:/\mathbf{s} \end{aligned}$$

where

$$\mathbf{z} = (\mathbf{y} : -\mathbf{p}_1) : /s$$

$$\mathbf{s} = \exp(\mathbf{p}_2)$$

The d2 evaluator function for this problem is

```
function linregevald2(transmorphic M, real scalar todo,
                      real rowvector b, fv, g, H)
{
    y1 = moptimize_calc_depvar(M, 1)
    p1 = moptimize_calc_xb(M, b, 1)
    p2 = moptimize_calc_xb(M, b, 2)
    s = exp(p2)
    z = (y1:-p1):/s
    fv = moptimize_util_sum(M, ln(normalden(y1:-p1, 0, s)))
    if (todo>=1) {
        s1 = z:/s
        s2 = z:^2 :- 1
        g1 = moptimize_util_vecsum(M, 1, s1, fv)
        g2 = moptimize_util_vecsum(M, 2, s2, fv)
        g = (g1, g2)
        if (todo==2) {
            h11 = -1:/s:^2
            h22 = -2*z:^2
            h12 = -2*z:/s
            H11 = moptimize_util_matsum(M, 1,1, h11, fv)
            H22 = moptimize_util_matsum(M, 2,2, h22, fv)
            H12 = moptimize_util_matsum(M, 1,2, h12, fv)
            H = (H11, H12 \ H12', H22)
        }
    }
}
```

The code to fit a model of mpg on weight and foreign reads

```
: M = moptimize_init()
: moptimize_init_evaluator(M, &linregevald2())
: moptimize_init_evaluatortype(M, "d2")
: moptimize_init_depvar(M, 1, "mpg")
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
: moptimize_init_eq_indepvars(M, 2, "")
: moptimize(M)
: moptimize_result_display(M)
```

By the way, function `linregevald2()` will work not only with type `d2`, but also with types `d1` and `d0`. Our function has the code to calculate first and second derivatives, but if we use type `d1`, `todo` will never be 2 and the second derivative part of our code will be ignored. If we use type `d0`, `todo` will never be 1 or 2 and so the first and second derivative parts of our code will be ignored. You could delete the unnecessary blocks if you wanted.

It is also worth trying the above code with types `d1debug` and `d2debug`. Type `d1debug` is like `d1`; the second derivative code will not be used. Also type `d1debug` will almost ignore the first



derivative code. Our program will be asked to make the calculation, but `moptimize()` will not use the results except to report a comparison of the derivatives we calculate with numerically calculated derivatives. That way, we can check that our program is right. Once we have done that, we move to type `d2debug`, which will check our second-derivative calculation.

## Example using type `lf*`

The `lf2` evaluator function for the linear-regression problem is almost identical to the type `d2` evaluator. It differs in that rather than return the summed log likelihood, we return the observation-level log likelihoods. And rather than return the gradient vector, we return the equation-level scores that when used with the chain-rule can be summed to produce the gradient. The conversion from `d2` to `lf2` was possible because of the observation-by-observation nature of the linear-regression problem; if the evaluator was not going to be implemented as `lf`, it always should have been implemented as `lf1` or `lf2` instead of `d1` or `d2`. In the `d2` evaluator above, we went to extra work—summing the scores—the result of which was to eliminate `moptimize()` features such as being able to automatically adjust for clusters and survey data. In a more appropriate type `d` problem—a problem for which a type `lf*` evaluator could not have been implemented—those scores never would have been available in the first place.

The `lf2` evaluator is

```
function linregevallf2(transmorphic M, real scalar todo,
                      real rowvector b, fv, S, H)
{
    y1 = moptimize_calc_depvar(M, 1)
    p1 = moptimize_calc_xb(M, b, 1)
    p2 = moptimize_calc_xb(M, b, 2)
    s   = exp(p2)
    z   = (y1:-p1):/s
    fv  = ln(normalden(y1:-p1, 0, s))
    if (todo>=1) {
        s1 = z:/s
        s2 = z:^2 :- 1
        S  = (s1, s2)
        if (todo==2) {
            h11 = -1:/s:^2
            h22 = -2*z:^2
            h12 = -2*z:/s
            mis = 0
            H11 = moptimize_util_matsum(M, 1,1, h11, mis)
            H22 = moptimize_util_matsum(M, 2,2, h22, mis)
            H12 = moptimize_util_matsum(M, 1,2, h12, mis)
            H    = (H11, H12 \ H12', H22)
        }
    }
}
```

The code to fit a model of mpg on weight and foreign reads nearly identically to the code we used in the [type lf2 case](#). We must specify the name of our type lf2 evaluator and specify that it is type lf2:

```
: M = moptimize_init()
: moptimize_init_evaluator(M, &linregevallf2())
: moptimize_init_evaluatortype(M, "lf2")
: moptimize_init_depvar(M, 1, "mpg")
: moptimize_init_eq_indepvars(M, 1, "weight foreign")
: moptimize_init_eq_indepvars(M, 2, "")
: moptimize(M)
: moptimize_result_display(M)
```

## Conformability

See [Syntax](#) above.

## Diagnostics

All functions abort with error when used incorrectly.

`moptimize()` aborts with error if it runs into numerical difficulties. `_moptimize()` does not; it instead returns a nonzero error code.

The `moptimize_result*()` functions abort with error if they run into numerical difficulties when called after `moptimize()` or `moptimize_evaluate()`. They do not abort when run after `_moptimize()` or `_moptimize_evaluate()`. They instead return a properly dimensioned missing result and set `moptimize_result_errorcode()` and `moptimize_result_errortext()`.

Ludwig Otto Hesse (1811–1874) was born in Königsberg, Prussia (now Kaliningrad, Russia) and studied mathematics and natural sciences at the university there, where Jacobi was one of his teachers and his doctoral supervisor. After short periods of school teaching, his career included various university posts at Königsberg, Halle, Heidelberg, and finally Munich, where he died. Hesse's main work was in algebra, including geometry, the use of determinants, and the calculus of variations. The Hessian matrix is named for him, as are Hesse's normal forms of linear and planar equations.

Donald Wesley Marquardt (1929–1997) was born in New York and obtained degrees in physics, mathematics, and statistics from Columbia and the University of Delaware. For 39 years, he worked at DuPont as a statistician and founding manager of the company's Quality Management and Technology Center. In retirement, Marquardt set up his own consultancy and remained an international leader in establishing standards for quality management and quality assurance. His work on nonlinear estimation is highly cited. Marquardt also made major contributions to ridge and generalized inverse regression, mixture designs, and analysis of unequally spaced time series.

## References

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- Haas, K. 1972. Ludwig Otto Hesse. In Vol. 6 of *Dictionary of Scientific Biography*, ed. C. C. Gillispie, 356–358. New York: Charles Scribner's Sons.
- Hahn, G. J. 1995. A conversation with Donald Marquardt. *Statistical Science* 10: 377–393.

## Also see

- [M-5] `optimize()` — Function optimization
- [M-4] `mathematical` — Important mathematical functions
- [M-4] `statistical` — Statistical functions

# Title

**[M-5] more()** — Create `--more--` condition

## Syntax

```
void          more()

real scalar  setmore()

void          setmore(real scalar onoff)

void          setmoreonexit(real scalar onoff)
```

## Description

`more()` displays `--more--` and waits for a key to be pressed. That is, `more()` does that if `more` is on, which it usually is, and it does nothing otherwise. `more` can be turned on and off by Stata's `set more` command (see [\[R\] more](#)) or by the functions below.

`setmore()` returns whether `more` is on or off, encoded as 1 and 0.

`setmore(onoff)` sets `more` on if *onoff*  $\neq$  0 and sets `more` off otherwise.

`setmoreonexit(onoff)` sets `more` on or off when the current execution ends. It has no effect on the current setting. The specified setting will take effect when control is passed back to the Mata prompt or to the calling ado-file or to Stata itself, and it will take effect regardless of whether execution ended because of a `return`, `exit()`, error, or abort. Only the first call to `setmoreonexit()` has that effect. Later calls have no effect whatsoever.

## Remarks

`setmoreonexit()` is used to ensure that the `more` setting is restored if a program wants to temporarily reset it:

```
setmoreonexit(setmore())
setmore(0)
```

Only the first invocation of `setmoreonexit()` has any effect. This way, a subroutine that is used in various contexts might also contain

```
setmoreonexit(setmore())
setmore(0)
```

and that will not cause the wrong `more` setting to be restored if an earlier routine had already done that and yet still cause the right setting to be restored if the subroutine is the first to issue `setmoreonexit()`.

## Conformability

`more()` takes no arguments and returns *void*.

`setmore()`:

*result:*       $1 \times 1$

`setmore(onoff), setmoreonexit(onoff):`

*onoff:*       $1 \times 1$

*result:*      *void*

## Diagnostics

None.

## Also see

[P] [more](#) — Pause until key is pressed

[M-4] [io](#) — I/O functions

# Title

[M-5] `_negate()` — Negate real matrix

## Syntax

```
void _negate(real matrix X)
```

## Description

`_negate(X)` speedily replaces  $X = -X$ .

## Remarks

Coding `_negate(X)` executes more quickly than coding  $X = -X$ .

However, coding

```
B = A
_negate(B)
```

does not execute more quickly than coding

```
B = -A
```

## Conformability

```
_negate(X):
      X:     $r \times c$ 
result:    void
```

## Diagnostics

None.  $X$  may be a view.

## Also see

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] norm() — Matrix and vector norms

## Syntax

*real scalar*   `norm(numeric matrix A)`

*real scalar*   `norm(numeric matrix A, real scalar p)`

## Description

`norm(A)` returns `norm(A, 2)`.

`norm(A, p)` returns the value of the norm of *A* for the specified *p*. The possible values and the meaning of *p* depend on whether *A* is a vector or a matrix.

When *A* is a vector, `norm(A, p)` returns

$$\begin{aligned} \text{sum}(\text{abs}(A)^{\wedge p})^{\wedge (1/p)} & \quad \text{if } 1 \leq p < \infty \\ \text{max}(\text{abs}(A)) & \quad \text{if } p \geq \infty \end{aligned}$$

When *A* is a matrix, returned is

<i>p</i>	<code>norm(A, p)</code>
0	<code>sqrt(trace(conj(A)'A))</code>
1	<code>max(colsum(abs(A)))</code>
2	<code>max(svdsv(A))</code>
∞	<code>max(rowsum(abs(A)))</code>

## Remarks

`norm(A)` and `norm(A, p)` calculate vector norms and matrix norms. *A* may be real or complex and need not be square when it is a matrix.

The formulas presented above are not the actual ones used in calculation. In the vector-norm case when  $1 \leq p < \infty$ , the formula is applied to `A:/max(abs(A))` and the result then multiplied by `max(abs(A))`. This prevents numerical overflow. A similar technique is used in calculating the matrix norm for  $p = 0$ , and that technique also avoids storage of `conj(A)'A`.

## Conformability

```
norm(A):
    A:      r × c
    result:  1 × 1

norm(A, p):
    A:      r × c
    p:      1 × 1
    result:  1 × 1
```

## Diagnostics

The `norm()` is defined to return 0 if  $A$  is void and missing if any element of  $A$  is missing.

`norm( $A$ ,  $p$ )` aborts with error if  $p$  is out of range. When  $A$  is a vector,  $p$  must be greater than or equal to 1. When  $A$  is a matrix,  $p$  must be 0, 1, 2, or . (missing).

`norm( $A$ )` and `norm( $A$ ,  $p$ )` return missing if the 2-norm is requested and the singular value decomposition does not converge, an event not expected to occur; see [M-5] `svd()`.

## Also see

[M-4] `matrix` — Matrix functions



## Syntax

*Gaussian normal*

```

f = normalden(z)
f = normalden(z, sd)
f = normalden(x, mean, sd)
p = normal(z)
z = invnormal(p)
ln(f) = lnnormalden(z)
ln(f) = lnnormalden(z, sd)
ln(f) = lnnormalden(x, mean, sd)
ln(p) = lnnormal(z)

```

*Binormal*

```

p = binormal(z1, z2, rho)

```

*Beta*

```

f = betaden(a, b, x)
p = ibeta(a, b, x)
q = ibetatail(a, b, x)
x = invibeta(a, b, p)
x = invibetatail(a, b, q)

```

*Binomial*

```

pk = binomialp(n, k, pi)
p = binomial(n, k, pi)
q = binomialtail(n, k, pi)
pi = invbinomial(n, k, p)
pi = invbinomialtail(n, k, q)

```

*Chi-squared*

```

p = chi2(n, x)
q = chi2tail(n, x)
x = invchi2(n, p)
x = invchi2tail(n, q)

```

*Dunnett's multiple range*

```

p = dunnettprob(k, df, x)
x = invdunnettprob(k, df, p)

```

*F*

```

f = Fden( $n_1$ ,  $n_2$ ,  $Fstat$ )
p = F( $n_1$ ,  $n_2$ ,  $Fstat$ )
q = Ftail( $n_1$ ,  $n_2$ ,  $Fstat$ )
 $Fstat$  = invF( $n_1$ ,  $n_2$ ,  $p$ )
 $Fstat$  = invFtail( $n_1$ ,  $n_2$ ,  $q$ )

```

*Gamma*

```

f = gammaden( $a$ ,  $b$ ,  $g$ ,  $x$ )
p = gammap( $a$ ,  $x$ )
q = gammaptail( $a$ ,  $x$ )
 $x$  = invgammap( $a$ ,  $p$ )
 $x$  = invgammaptail( $a$ ,  $q$ )
dg/da = dgammabda( $a$ ,  $x$ )
dg/dx = dgammabdx( $a$ ,  $x$ )
d2g/da2 = dgammabdada( $a$ ,  $x$ )
d2g/dadx = dgammabdadx( $a$ ,  $x$ )
d2g/dx2 = dgammabdxdx( $a$ ,  $x$ )

```

*Hypergeometric*

```

pk = hypergeometricp( $N$ ,  $K$ ,  $n$ ,  $k$ )
p = hypergeometric( $N$ ,  $K$ ,  $n$ ,  $k$ )

```

*Negative binomial*

```

pk = nbinomialp( $n$ ,  $k$ ,  $pi$ )
p = nbinomial( $n$ ,  $k$ ,  $pi$ )
q = nbinomialtail( $n$ ,  $k$ ,  $pi$ )
pi = invnbinomial( $n$ ,  $k$ ,  $p$ )
pi = invnbinomialtail( $n$ ,  $k$ ,  $q$ )

```

*Noncentral Beta*

```

f = nbetaden( $a$ ,  $b$ ,  $L$ ,  $x$ )
p = nibeta( $a$ ,  $b$ ,  $L$ ,  $x$ )
 $x$  = invnibeta( $a$ ,  $b$ ,  $L$ ,  $p$ )

```

*Noncentral chi-squared*

```

p = nchi2( $n$ ,  $L$ ,  $x$ )
 $x$  = invnchi2( $n$ ,  $L$ ,  $p$ )
 $L$  = npnchi2( $n$ ,  $x$ ,  $p$ )

```

*Noncentral F*

```
f = nFden( $n_1$ ,  $n_2$ ,  $L$ ,  $F$ )
q = nFtail( $n_1$ ,  $n_2$ ,  $L$ ,  $F$ )
F = invnFtail( $n_1$ ,  $n_2$ ,  $L$ ,  $q$ )
```

*Poisson*

```
pk = poissonp( $mean$ ,  $k$ )
p = poisson( $mean$ ,  $k$ )
q = poissontail( $mean$ ,  $k$ )
m = invpoisson( $k$ ,  $p$ )
m = invpoissontail( $k$ ,  $q$ )
```

*Student's t*

```
f = tden( $n$ ,  $t$ )
q = ttail( $n$ ,  $t$ )
t = invttail( $n$ ,  $q$ )
```

*Tukey's Studentized range*

```
p = tukeyprob( $k$ ,  $df$ ,  $x$ )
x = invtukeyprob( $k$ ,  $df$ ,  $p$ )
```

where

1. All functions return real and all arguments are real.
2. The left-hand-side notation is used to assist in interpreting the meaning of the returned value:

```
f = density value
pk = probability of discrete outcome  $K = \Pr(K = k)$ 
p = left cumulative
  =  $\Pr(-\text{infinity} < \text{statistic} \leq x)$  (continuous)
  =  $\Pr(0 \leq K \leq k)$  (discrete)
q = right cumulative
  =  $1 - p$  (continuous)
  =  $\Pr(K \geq k) = 1 - p + pk$  (discrete)
```

3. Hypergeometric distribution:

```
N = number of objects in the population
K = number of objects in the population with the characteristic of interest,
     $K < N$ 
n = sample size,  $n < N$ 
k = number of objects in the sample with the characteristic of interest,
     $\max(0, n - N + K) \leq k \leq \min(K, n)$ 
```

4. Negative binomial distribution:  $n > 0$  and may be nonintegral.

## Description

The above functions return density values, cumulatives, reverse cumulatives, and in one case, derivatives of the indicated probability density function. These functions mirror the Stata functions of the same name and in fact are the Stata functions.

See [Probability distributions and density functions](#) in [D] [functions](#) for function details. In the syntax diagram above, some arguments have been renamed in hope of aiding understanding, but the function arguments match one to one with the underlying Stata functions.

## Remarks

Remarks are presented under the following headings:

- [R-conformability](#)
- [A note concerning invbinomial\(\) and invbinomialtail\(\)](#)
- [A note concerning ibeta\(\)](#)
- [A note concerning gammap\(\)](#)

### R-conformability

The above functions are usually used with scalar arguments and then return a scalar result:

```
: x = chi2(10, 12)
: x
.7149434997
```

The arguments may, however, be vectors or matrices. For instance,

```
: x = chi2((10,11,12), 12)
: x
      1      2      3
1 | .7149434997  .6363567795  .5543203586
```

```
: x = chi2(10, (12,12.5,13))
: x
      1      2      3
1 | .7149434997  .7470146767  .7763281832
```

```
: x = chi2((10,11,12), (12,12.5,13))
: x
      1      2      3
1 | .7149434997  .6727441644  .6309593164
```

In the last example, the numbers correspond to `chi2(10,12)`, `chi2(11,12.5)`, and `chi2(12,13)`.

Arguments are required to be [r-conformable](#) (see [\[M-6\] Glossary](#)), and thus,

```
: x = chi2((10\11\12), (12,12.5,13))
: x
```

	1	2	3
1	.7149434997	.7470146767	.7763281832
2	.6363567795	.6727441644	.7066745906
3	.5543203586	.593595966	.6309593164

which corresponds to

	1	2	3
1	<code>chi2(10,12)</code>	<code>chi2(10,12.5)</code>	<code>chi2(10,13)</code>
2	<code>chi2(11,12)</code>	<code>chi2(11,12.5)</code>	<code>chi2(11,13)</code>
3	<code>chi2(12,12)</code>	<code>chi2(12,12.5)</code>	<code>chi2(12,13)</code>

## A note concerning `invbinomial()` and `invbinomialtail()`

`invbinomial( $n, k, pi$ )` `invbinomialtail( $n, k, q$ )` are useful for calculating confidence intervals for  $pi$ , the probability of a success. `invbinomial()` returns the probability  $pi$  such that the probability of observing  $k$  or fewer successes in  $n$  trials is  $p$ . `invbinomialtail()` returns the probability  $pi$  such that the probability of observing  $k$  or more successes in  $n$  trials is  $q$ .

## A note concerning `ibeta()`

`ibeta( $a, b, x$ )` is known as the cumulative beta distribution, and it is known as the incomplete beta function  $I_x(a, b)$ .

## A note concerning `gammap()`

`gammap( $a, x$ )` is known as the cumulative gamma distribution, and it is known as the incomplete gamma function  $P(a, x)$ .

## Conformability

All functions require that arguments be r-conformable; see [R-conformability](#) above. Returned is matrix of `max(argument rows)` rows and `max(argument columns)` columns containing element-by-element calculated results.

## Diagnostics

All functions return missing when arguments are out of range.

## Also see

[M-4] [statistical](#) — Statistical functions

## Syntax

`S = optimize_init()`

*(varies)* `optimize_init_which(S [ , { "max" | "min" } ])`  
*(varies)* `optimize_init_evaluator(S [ , &function() ])`  
*(varies)* `optimize_init_evaluatortype(S [ , evaluatortype ])`  
*(varies)* `optimize_init_negH(S, { "off" | "on" })`  
*(varies)* `optimize_init_params(S [ , real rowvector initialvalues ])`  
*(varies)* `optimize_init_nmsimplexdeltas(S [ , real rowvector delta ])`  
*(varies)* `optimize_init_argument(S, real scalar k [ , X ])`  
*(varies)* `optimize_init_narguments(S [ , real scalar K ])`  
*(varies)* `optimize_init_cluster(S, c)`  
*(varies)* `optimize_init_colstripe(S [ , stripe ])`  
*(varies)* `optimize_init_technique(S [ , technique ])`  
*(varies)* `optimize_init_singularHmethod(S [ , singularHmethod ])`  
*(varies)* `optimize_init_conv_maxiter(S [ , real scalar maxiter ])`  
*(varies)* `optimize_init_conv_warning(S, { "on" | "off" })`  
*(varies)* `optimize_init_conv_ptol(S [ , real scalar ptol ])`  
*(varies)* `optimize_init_conv_vtol(S [ , real scalar vtol ])`  
*(varies)* `optimize_init_conv_nrtol(S [ , real scalar nrtol ])`  
*(varies)* `optimize_init_conv_ignorenrtol(S, { "off" | "on" })`  
*(varies)* `optimize_init_iterid(S [ , string scalar id ])`  
*(varies)* `optimize_init_valueid(S [ , string scalar id ])`  
*(varies)* `optimize_init_tracelevel(S [ , tracelevel ])`

```

(varies)      optimize_init_trace_dots(S, { "off" | "on" })
(varies)      optimize_init_trace_value(S, { "on" | "off" })
(varies)      optimize_init_trace_tol(S, { "off" | "on" })
(varies)      optimize_init_trace_step(S, { "off" | "on" })
(varies)      optimize_init_trace_paramdiffs(S, { "off" | "on" })
(varies)      optimize_init_trace_params(S, { "off" | "on" })
(varies)      optimize_init_trace_gradient(S, { "off" | "on" })
(varies)      optimize_init_trace_Hessian(S, { "off" | "on" })
(varies)      optimize_init_evaluations(S, { "off" | "on" })
(varies)      optimize_init_constraints(S [, real matrix Cc])
(varies)      optimize_init_verbose(S [, real scalar verbose])

```

```
real rowvector optimize(S)
```

```
real scalar      _optimize(S)
```

```
void             optimize_evaluate(S)
```

```
real scalar      _optimize_evaluate(S)
```

```
real rowvector optimize_result_params(S)
```

```
real scalar      optimize_result_value(S)
```

```
real scalar      optimize_result_value0(S)
```

```
real rowvector optimize_result_gradient(S)
```

```
real matrix      optimize_result_scores(S)
```

```
real matrix      optimize_result_Hessian(S)
```

```
real matrix      optimize_result_V(S)
```

```
string scalar    optimize_result_Vtype(S)
```

```
real matrix      optimize_result_V_oim(S)
```

```
real matrix      optimize_result_V_opg(S)
```

```
real matrix      optimize_result_V_robust(S)
```

```
real scalar      optimize_result_iterations(S)
```

```
real scalar      optimize_result_converged(S)
```

```
real colvector   optimize_result_iterationlog(S)
```

```
real rowvector   optimize_result_evaluations(S)
```

```
real scalar      optimize_result_errorcode(S)
```

```
string scalar    optimize_result_errortext(S)
```

```
real scalar      optimize_result_returncode(S)
```



`void optimize_query(S)`

where *S*, if it is declared, should be declared

`transmorphic S`

and where *evaluator**type* optionally specified in `optimize_init_evaluator`*type*() is

<i>evaluator</i> <i>type</i>	Description
"d0"	<i>function</i> () returns <i>scalar</i> value
"d1"	same as "d0" and returns gradient <i>rowvector</i>
"d2"	same as "d1" and returns Hessian <i>matrix</i>
"d1debug"	same as "d1" but checks gradient
"d2debug"	same as "d2" but checks gradient and Hessian
"gf0"	<i>function</i> () returns <i>colvector</i> value
"gf1"	same as "gf0" and returns score <i>matrix</i>
"gf2"	same as "gf1" and returns Hessian <i>matrix</i>
"gf1debug"	same as "gf1" but checks gradient
"gf2debug"	same as "gf2" but checks gradient and Hessian

The default is "d0" if not set.

and where *technique* optionally specified in `optimize_init_technique`() is

<i>technique</i>	Description
"nr"	modified Newton–Raphson
"dfp"	Davidon–Fletcher–Powell
"bfgs"	Broyden–Fletcher–Goldfarb–Shanno
"bhhh"	Berndt–Hall–Hall–Hausman
"nm"	Nelder–Mead

The default is "nr".

and where *singularHmethod* optionally specified in `optimize_init_singularHmethod`() is

<i>singularHmethod</i>	Description
"m-marquardt"	modified Marquardt algorithm
"hybrid"	mixture of steepest descent and Newton

The default is "m-marquardt" if not set;

"hybrid" is equivalent to `ml`'s `difficult` option; see [R] [ml](#).

and where *tracelevel* optionally specified in `optimize_init_tracelevel()` is

<i>tracelevel</i>	To be displayed each iteration
"none"	nothing
"value"	function value
"tolerance"	previous + convergence values
"step"	previous + stepping information
"paramdiffs"	previous + parameter relative differences
"params"	previous + parameter values
"gradient"	previous + gradient vector
"hessian"	previous + Hessian matrix

The default is "value" if not set.

## Description

These functions find parameter vector or scalar  $p$  such that function  $f(p)$  is a maximum or a minimum.

`optimize_init()` begins the definition of a problem and returns  $S$ , a problem-description handle set to contain default values.

The `optimize_init_*( $S$ , ...)` functions then allow you to modify those defaults. You use these functions to describe your particular problem: to set whether you wish maximization or minimization, to set the identity of function  $f()$ , to set initial values, and the like.

`optimize( $S$ )` then performs the optimization. `optimize()` returns *real rowvector*  $p$  containing the values of the parameters that produce a maximum or minimum.

The `optimize_result_*( $S$ )` functions can then be used to access other values associated with the solution.

Usually you would stop there. In other cases, you could restart optimization by using the resulting parameter vector as new initial values, change the optimization technique, and restart the optimization:

```
optimize_init_param( $S$ , optimize_result_param( $S$ ))
optimize_init_technique( $S$ , "dfp")
optimize( $S$ )
```

Aside: The `optimize_init_*( $S$ , ...)` functions have two modes of operation. Each has an optional argument that you specify to set the value and that you omit to query the value. For instance, the full syntax of `optimize_init_params()` is

```
void optimize_init_params( $S$ , real rowvector initialvalues)
real rowvector optimize_init_params( $S$ )
```

The first syntax sets the initial values and returns nothing. The second syntax returns the previously set (or default, if not set) initial values.

All the `optimize_init_*( $S$ , ...)` functions work the same way.

## Remarks

Remarks are presented under the following headings:

*First example*

*Notation*

*Type d evaluators*

*Example of d0, d1, and d2*

*d1debug and d2debug*

*Type gf evaluators*

*Example of gf0, gf1, and gf2*

*Functions*

```
optimize_init()
optimize_init_which()
optimize_init_evaluator() and optimize_init_evaluortype()
optimize_init_negH()
optimize_init_params()
optimize_init_nmsimplexdeltas()
optimize_init_argument() and optimize_init_narguments()
optimize_init_cluster()
optimize_init_colstripe()
optimize_init_technique()
optimize_init_singularHmethod()
optimize_init_conv_maxiter()
optimize_init_conv_warning()
optimize_init_conv_ptol(), ... _vtol(), ... _nrtol()
optimize_init_conv_ignorenrtol()
optimize_init_iterid()
optimize_init_valueid()
optimize_init_tracelevel()
optimize_init_trace_dots(), ... _value(), ... _tol(), ... _step(), ... _paramdiffs(),
... _params(), ... _gradient(), ... _Hessian()
optimize_init_evaluations()
optimize_init_constraints()
optimize_init_verbose()

optimize()
_optimize()
optimize_evaluate()
_optimize_evaluate()

optimize_result_params()
optimize_result_value() and optimize_result_value0()
optimize_result_gradient()
optimize_result_scores()
optimize_result_Hessian()
optimize_result_V() and optimize_result_Vtype()
optimize_result_V_oim(), ... _opg(), ... _robust()
optimize_result_iterations()
optimize_result_converged()
optimize_result_iterationlog()
optimize_result_evaluations()
optimize_result_errorcode(), ... _errortext(), and ... _returncode()
optimize_query()
```

## First example

The optimization functions may be used interactively.

Below we use the functions to find the value of  $x$  that maximizes  $y = \exp(-x^2 + x - 3)$ :

```

: void myeval(todo, x, y, g, H)
> {
>     y = exp(-x^2 + x - 3)
> }
note: argument todo unused
note: argument g unused
note: argument H unused
: S = optimize_init()
: optimize_init_evaluator(S, &myeval())
: optimize_init_params(S, 0)
: x = optimize(S)
Iteration 0: f(p) = .04978707
Iteration 1: f(p) = .04978708
Iteration 2: f(p) = .06381186
Iteration 3: f(p) = .06392786
Iteration 4: f(p) = .06392786
: x
.5

```

## Notation

We wrote the above in the way that mathematicians think, that is, optimizing  $y = f(x)$ . Statisticians, on the other hand, think of optimizing  $s = f(b)$ . To avoid favoritism, we will write  $v = f(p)$  and write the general problem with the following notation:

Maximize or minimize  $v = f(p)$ ,

$v$ : a scalar

$p$ :  $1 \times np$

subject to the constraint  $Cp' = c$ ,

$C$ :  $nc \times np$  ( $nc = 0$  if no constraints)

$c$ :  $nc \times 1$

where  $g$ , the gradient vector, is  $g = f'(p) = df/dp$ ,

$g$ :  $1 \times np$

and  $H$ , the Hessian matrix, is  $H = f''(p) = d^2f/dpdp'$

$H$ :  $np \times np$

## Type d evaluators

You must write an evaluator function to calculate  $f()$  before you can use the optimization functions. The example we showed above was of what is called a type d evaluator. Let's stay with that.

The evaluator function we wrote was

```
void myeval(todo, x, y, g, H)
{
    y = exp(-x^2 + x - 3)
}
```

All type d evaluators open the same way,

```
void evaluator(todo, x, y, g, H)
```

although what you name the arguments is up to you. We named the arguments the way that mathematicians think, although we could just as well have named them the way that statisticians think:

```
void evaluator(todo, b, s, g, H)
```

To avoid favoritism, we will write them as

```
void evaluator(todo, p, v, g, H)
```

that is, we will think in terms of optimizing  $v = f(p)$ .

Here is the full definition of a type d evaluator:

---

```
void evaluator(real scalar todo, real rowvector p, v, g, H)
```

where  $v$ ,  $g$ , and  $H$  are values to be returned:

```
v:    real scalar
g:    real rowvector
H:    real matrix
```

*evaluator()* is to fill in  $v$  given the values in  $p$  and optionally to fill in  $g$  and  $H$ , depending on the value of *todo*:

---

<i>todo</i>	Required action by <i>evaluator()</i>
-------------	---------------------------------------

---

0	calculate $v = f(p)$ and store in $v$
---	---------------------------------------

1	calculate $v = f(p)$ and $g = f'(p)$ and store in $v$ and $g$
---	---

2	calculate $v = f(p)$ , $g = f'(p)$ , and $H = f''(p)$ and store in $v$ , $g$ , and $H$
---	--

---

*evaluator()* may return  $v = .$  if  $f()$  cannot be evaluated at  $p$ . Then  $g$  and  $H$  need not be filled in even if requested.

---

An evaluator does not have to be able to do all of this. In the first example, *myeval()* could handle only *todo* = 0. There are three types of type d evaluators:

d type	Capabilities expected of <i>evaluator()</i>
d0	can calculate $v = f(p)$
d1	can calculate $v = f(p)$ and $g = f'(p)$
d2	can calculate $v = f(p)$ and $g = f'(p)$ and $H = f''(p)$

---

`myeval()` is a type `d0` evaluator. Type `d0` evaluators are never asked to calculate  $g$  or  $H$ . Type `d0` is the default type but, if we were worried that it was not, we could set the evaluator type before invoking `optimize()` by coding

```
optimize_init_evaluatoretype(S, "d0")
```

Here are the code outlines of the three types of evaluators:

---

```
void d0_evaluator(todo, p, v, g, H)
{
    v = ...
}
```

---



---

```
void d1_evaluator(todo, p, v, g, H)
{
    v = ...
    if (todo>=1) {
        g = ...
    }
}
```

---



---

```
void d2_evaluator(todo, p, v, g, H)
{
    v = ...
    if (todo>=1) {
        g = ...
        if (todo==2) {
            H = ...
        }
    }
}
```

---

Here is the code outline where there are three additional user arguments to the evaluator:

---

```
void d0_user3_eval(todo, p, u1, u2, u3, v, g, H)
{
    v = ...
}
```

---

## Example of `d0`, `d1`, and `d2`

We wish to find the  $p_1$  and  $p_2$  corresponding to the maximum of

$$v = \exp(-p_1^2 - p_2^2 - p_1 p_2 + p_1 - p_2 - 3)$$

A d0 solution to the problem would be

```
: void eval0(todo, p, v, g, H)
> {
>     v = exp(-p[1]^2 - p[2]^2 - p[1]*p[2] + p[1] - p[2] - 3)
> }
note: argument todo unused
note: argument g unused
note: argument h unused
: S = optimize_init()
: optimize_init_evaluator(S, &eval0())
: optimize_init_params(S, (0,0))
: p = optimize(S)
Iteration 0: f(p) = .04978707 (not concave)
Iteration 1: f(p) = .12513024
Iteration 2: f(p) = .13495886
Iteration 3: f(p) = .13533527
Iteration 4: f(p) = .13533528
: p
      1      2
1  

|   |    |
|---|----|
| 1 | -1 |
|---|----|


```

A d1 solution to the problem would be

```
: void eval1(todo, p, v, g, H)
> {
>     v = exp(-p[1]^2 - p[2]^2 - p[1]*p[2] + p[1] - p[2] - 3)
>     if (todo==1) {
>         g[1] = (-2*p[1] - p[2] + 1)*v
>         g[2] = (-2*p[2] - p[1] - 1)*v
>     }
> }
note: argument H unused
: S = optimize_init()
: optimize_init_evaluator(S, &eval1())
: optimize_init_evaluortype(S, "d1") ← important
: optimize_init_params(S, (0,0))
: p = optimize(S)
Iteration 0: f(p) = .04978707 (not concave)
Iteration 1: f(p) = .12513026
Iteration 2: f(p) = .13496887
Iteration 3: f(p) = .13533527
Iteration 4: f(p) = .13533528
: p
      1      2
1  

|   |    |
|---|----|
| 1 | -1 |
|---|----|


```

The d1 solution is better than the d0 solution because it runs faster and usually is more accurate. Type d1 evaluators require more code, however, and deriving analytic derivatives is not always possible.

A d2 solution to the problem would be

```

: void eval2(todo, p, v, g, H)
> {
>     v = exp(-p[1]^2 - p[2]^2 - p[1]*p[2] + p[1] - p[2] - 3)
>     if (todo>=1) {
>         g[1] = (-2*p[1] - p[2] + 1)*v
>         g[2] = (-2*p[2] - p[1] - 1)*v
>         if (todo==2) {
>             H[1,1] = -2*v + (-2*p[1]-p[2]+1)*g[1]
>             H[2,1] = -1*v + (-2*p[2]-p[1]-1)*g[1]
>             H[2,2] = -2*v + (-2*p[2]-p[1]-1)*g[2]
>             _makesymmetric(H)
>         }
>     }
> }

: S = optimize_init()
: optimize_init_evaluator(S, &eval2())
: optimize_init_evaluortype(S, "d2")           ← important
: optimize_init_params(S, (0,0))

: p = optimize(S)
Iteration 0:  f(p) = .04978707  (not concave)
Iteration 1:  f(p) = .12513026
Iteration 2:  f(p) = .13496887
Iteration 3:  f(p) = .13533527
Iteration 4:  f(p) = .13533528

: p
      1      2
1  

|   |    |
|---|----|
| 1 | -1 |
|---|----|


```

A d2 solution is best because it runs fastest and usually is the most accurate. Type d2 evaluators require the most code, and deriving analytic derivatives is not always possible.

In the d2 evaluator `eval2()`, note our use of `_makesymmetric()`. Type d2 evaluators are required to return  $H$  as a symmetric matrix; filling in just the lower or upper triangle is not sufficient. The easiest way to do that is to fill in the lower triangle and then use `_makesymmetric()` to reflect the lower off-diagonal elements; see [M-5] [makesymmetric\(\)](#).

## d1debug and d2debug

In addition to evaluator types "d0", "d1", and "d2" that are specified in `optimize_init_evaluortype(S, evaluortype)`, there are two more: "d1debug" and "d2debug". They assist in coding d1 and d2 evaluators.

In [Example of d0, d1, and d2](#) above, we admit that we did not correctly code the functions `eval1()` and `eval2()` at the outset, before you saw them. In both cases, that was because we had taken the derivatives incorrectly. The problem was not with our code but with our math. `d1debug` and `d2debug` helped us find the problem.

`d1debug` is an alternative to `d1`. When you code `optimize_init_evaluortype(S, "d1debug")`, the derivatives you calculate are not taken seriously. Instead, `optimize()` calculates its own numerical derivatives and uses those. Each time `optimize()` does that, however, it compares your derivatives to the ones it calculated and gives you a report on how they differ. If you have coded correctly, they should not differ by much.



d2debug does the same thing, but for d2 evaluators. When you code `optimize_init_evaluortype(S, "d2debug")`, `optimize()` uses numerical derivatives but, each time, `optimize()` gives you a report on how much your results for the gradient and for the Hessian differ from the numerical calculations.

For each comparison, `optimize()` reports just one number: the `mrldif()` (see [M-5] `reldif()`) between your results and the numerical ones. When you have done things right, gradient vectors will differ by approximately  $1e-12$  or less and Hessians will differ by  $1e-7$  or less.

When differences are large, you will want to see not only the summary comparison but also the full vectors and matrices so that you can compare your results element by element with those calculated numerically. Sometimes the error is in one element and not the others. To do this, set the trace level with `optimize_init_tracelevel(S, tracelevel)` before issuing `optimize()`. Code `optimize_init_tracelevel(S, "gradient")` to get a full report on the gradient comparison, or set `optimize_init_tracelevel(S, "hessian")` to get a full report on the gradient comparison and the Hessian comparison.

## Type gf evaluators

In some statistical applications, you will find `gf0`, `gf1`, and `gf2` more convenient to code than `d0`, `d1`, and `d2`. The *gf* stands for general form.

In statistical applications, one tends to think of a dataset of values arranged in matrix  $X$ , the rows of which are observations. A function  $h(p, X[i, :])$  can be calculated for each row separately, and it is the sum of those resulting values that forms the function  $f(p)$  that is to be maximized or minimized.

The `gf0`, `gf1`, and `gf2` methods are for such cases.

In a type `d0` evaluator, you return scalar  $v = f(p)$ .

In a type `gf0` evaluator, you return a column vector  $v$  such that `colsum(v) = f(p)`.

In a type `d1` evaluator, you return  $v = f(p)$  and you return a row vector  $g = f'(p)$ .

In a type `gf1` evaluator, you return  $v$  such that `colsum(v) = f(p)` and you return matrix  $g$  such that `colsum(g) = f'(p)`.

In a type `d2` evaluator, you return  $v = f(p)$ ,  $g = f'(p)$ , and you return  $H = f''(p)$ .

In a type `gf2` evaluator, you return  $v$  such that `colsum(v) = f(p)`,  $g$  such that `colsum(g) = f'(p)`, and you return  $H = f''(p)$ . This is the same  $H$  returned for `d2`.

The code outline for type *gf* evaluators is the same as those for *d* evaluators. For instance, the outline for a `gf2` evaluator is

---

```
void gf2_evaluator(todo, p, v, g, H)
{
    v = ...
    if (todo>=1) {
        g = ...
        if (todo==2) {
            H = ...
        }
    }
}
```

---

The above is the same as the outline for d2 evaluators. All that differs is that  $v$  and  $g$ , which were *real scalar* and *real rowvector* in the d2 case, are now *real colvector* and *real matrix* in the gf2 case. The same applies to gf1 and gf0.

The type gf evaluators arise in statistical applications and, in such applications, there are data; that is, just knowing  $p$  is not sufficient to calculate  $v$ ,  $g$ , and  $H$ . Actually, that same problem can arise when coding type d evaluators as well.

You can pass extra arguments to evaluators, whether they be d0, d1, or d2 or gf0, gf1, or gf2. The first line of all evaluators, regardless of style, is

```
void evaluator(todo, p, v, g, H)
```

If you code

```
optimize_init_argument(S, 1, X)
```

the first line becomes

```
void evaluator(todo, p, X, v, g, H)
```

If you code

```
optimize_init_argument(S, 1, X)
optimize_init_argument(S, 2, Y)
```

the first line becomes

```
void evaluator(todo, p, X, Y, v, g, H)
```

and so on, up to nine extra arguments. That is, you can specify extra arguments to be passed to your function. These extra arguments should be placed right after the parameter vector.

### Example of gf0, gf1, and gf2

You have the following data:

: x	
	1
1	.35
2	.29
3	.3
4	.3
5	.65
6	.56
7	.37
8	.16
9	.26
10	.19

You believe that the data are the result of a beta distribution process with fixed parameters alpha and beta and you wish to obtain the maximum likelihood estimates of alpha and beta ( $a$  and  $b$  in what follows). The formula for the density of the beta distribution is

$$\text{density}(x) = \frac{\Gamma(a+b)}{\Gamma(a)\Gamma(b)} x^{a-1} (1-x)^{b-1}$$

The gf0 solution to this problem is

```

: void lnbtaden0(todo, p, x, lnf, S, H)
> {
>     a  = p[1]
>     b  = p[2]
>     lnf = lngamma(a+b) :- lngamma(a) :- lngamma(b) :+
>           (a-1)*log(x) :+ (b-1)*log(1:-x)
> }
note: argument todo unused
note: argument S unused
note: argument H unused
: S = optimize_init()
: optimize_init_evaluator(S, &lnbtaden0())
: optimize_init_evaluatortype(S, "gf0")
: optimize_init_params(S, (1,1))
: optimize_init_argument(S, 1, x)           ← important
: p = optimize(S)
Iteration 0:  f(p) =           0
Iteration 1:  f(p) = 5.7294728
Iteration 2:  f(p) = 5.7646641
Iteration 3:  f(p) = 5.7647122
Iteration 4:  f(p) = 5.7647122
: p
      1          2
1  3.714209592  7.014926315

```

Note the following:

1. Rather than calling the returned value `v`, we called it `lnf`. You can name the arguments as you please.
2. We arranged for an extra argument to be passed by coding `optimize_init_argument(S, 1, x)`. The extra argument is the vector `x`, which we listed previously for you. In our function, we received the argument as `x`, but we could have used a different name, just as we used `lnf` rather than `v`.
3. We set the evaluator type to "gf0".

This being a statistical problem, we should be interested not only in the estimates `p` but also in their variance. We can get this from the inverse of the negative Hessian, which is the observed information matrix:

```

: optimize_result_V_oim(S)
[symmetric]
      1          2
1  2.556301184
2  4.498194785  9.716647065

```

The gf1 solution to this problem is

```

: void lnbetaden1(todo, p, x, lnf, S, H)
> {
>     a = p[1]
>     b = p[2]
>     lnf = lngamma(a+b) :- lngamma(a) :- lngamma(b) :+
>         (a-1)*log(x) :+ (b-1)*log(1:-x)
>     if (todo >= 1) {
>         S = J(rows(x), 2, .)
>         S[:,1] = log(x) :+ digamma(a+b) :- digamma(a)
>         S[:,2] = log(1:-x) :+ digamma(a+b) :- digamma(b)
>     }
> }
note: argument H unused
: S = optimize_init()
: optimize_init_evaluator(S, &lnbetaden1())
: optimize_init_evaluortype(S, "gf1")
: optimize_init_params(S, (1,1))
: optimize_init_argument(S, 1, x)
: p = optimize(S)
Iteration 0: f(p) = 0
Iteration 1: f(p) = 5.7297061
Iteration 2: f(p) = 5.7641349
Iteration 3: f(p) = 5.7647121
Iteration 4: f(p) = 5.7647122
: p
      1          2
1  [ 3.714209343   7.014925751 ]

: optimize_result_V_oim(S)
[symmetric]
      1          2
1  [ 2.556299425
2  [ 4.49819212   9.716643068 ]

```

Note the following:

1. We called the next-to-last argument of `lnbetaden1()` `S` rather than `g` in accordance with standard statistical jargon. What is being returned is in fact the observation-level scores, which sum to the gradient vector.
2. We called the next-to-last argument `S` even though that name conflicted with `S` outside the program, where `S` is the problem handle. Perhaps we should have renamed the outside `S`, but there is no confusion on Mata's part.
3. In our program, we allocated `S` for ourselves: `S = J(rows(x), 2, .)`. It is worth comparing this with the example of `d1` in [Example of d0, d1, and d2](#), where we did not need to allocate `g`. In `d1`, `optimize()` preallocates `g` for us. In `gf1`, `optimize()` cannot do this because it has no idea how many "observations" we have.

The gf2 solution to this problem is

```

: void lnbetaden2(todo, p, x, lnf, S, H)
> {
>     a  = p[1]
>     b  = p[2]
>     lnf = lngamma(a+b) :- lngamma(a) :- lngamma(b) :+
>         (a-1)*log(x) :+ (b-1)*log(1:-x)
>     if (todo >= 1) {
>         S      = J(rows(x), 2, .)
>         S[.,1] = log(x) :+ digamma(a+b) :- digamma(a)
>         S[.,2] = log(1:-x) :+ digamma(a+b) :- digamma(b)
>         if (todo==2) {
>             n = rows(x)
>             H[1,1] = n*(trigamma(a+b) - trigamma(a))
>             H[2,1] = n*(trigamma(a+b))
>             H[2,2] = n*(trigamma(a+b) - trigamma(b))
>             _makesymmetric(H)
>         }
>     }
> }
: S = optimize_init()
: optimize_init_evaluator(S, &lnbetaden2())
: optimize_init_evaluortype(S, "gf2")
: optimize_init_params(S, (1,1))
: optimize_init_argument(S, 1, x)
: p = optimize(S)
Iteration 0: f(p) = 0
Iteration 1: f(p) = 5.7297061
Iteration 2: f(p) = 5.7641349
Iteration 3: f(p) = 5.7647121
Iteration 4: f(p) = 5.7647122
: p
      1          2
1  [ 3.714209343   7.014925751 ]

: optimize_result_V_oim(S)
[symmetric]
      1          2
1  [ 2.556299574
2  [ 4.498192412   9.716643651 ]

```

## Functions

### optimize\_init()

*transmorphic* optimize\_init()

optimize\_init() is used to begin an optimization problem. Store the returned result in a variable name of your choosing; we have used *S* in this documentation. You pass *S* as the first argument to the other optimize\*() functions.

optimize\_init() sets all optimize\_init\_\*() values to their defaults. You may use the query form of the optimize\_init\_\*() to determine an individual default, or you can use optimize\_query() to see them all.

The query form of `optimize_init_*`() can be used before or after optimization performed by `optimize()`.

### optimize\_init\_which()

```
void          optimize_init_which(S, {"max" | "min"})
string scalar optimize_init_which(S)
```

`optimize_init_which(S, which)` specifies whether `optimize()` is to perform maximization or minimization. The default is maximization if you do not invoke this function.

`optimize_init_which(S)` returns "max" or "min" according to which is currently set.

### optimize\_init\_evaluator() and optimize\_init\_evaluortype()

```
void optimize_init_evaluator(S, pointer(real function) scalar fptr)
void optimize_init_evaluortype(S, evaluortype)

pointer(real function) scalar optimize_init_evaluator(S)
string scalar           optimize_init_evaluortype(S)
```

`optimize_init_evaluator(S, fptr)` specifies the function to be called to evaluate  $f(p)$ . Use of this function is required. If your function is named `myfcn()`, you code `optimize_init_evaluator(S, &myfcn())`.

`optimize_init_evaluortype(S, evaluortype)` specifies the capabilities of the function that has been set using `optimize_init_evaluator()`. Alternatives for *evaluortype* are "d0", "d1", "d2", "d1debug", "d2debug", "gf0", "gf1", "gf2", "gf1debug", and "gf2debug". The default is "d0" if you do not invoke this function.

`optimize_init_evaluator(S)` returns a pointer to the function that has been set.

`optimize_init_evaluortype(S)` returns the *evaluortype* currently set.

### optimize\_init\_negH()

`optimize_init_negH(S, {"off" | "on"})` sets whether the evaluator you have written returns  $H$  or  $-H$ , the Hessian or the negative of the Hessian, if it returns a Hessian at all. This is for backward compatibility with prior versions of Stata's `ml` command (see [R] `ml`). Modern evaluators return  $H$ . The default is "off".

### optimize\_init\_params()

```
void          optimize_init_params(S, real rowvector initialvalues)
real rowvector optimize_init_params(S)
```

`optimize_init_params(S, initialvalues)` sets the values of  $p$  to be used at the start of the first iteration. Use of this function is required.

`optimize_init_params(S)` returns the initial values that will be (or were) used.

**optimize\_init\_nmsimplexdeltas()**

*void*                    optimize\_init\_nmsimplexdeltas(*S*, *real rowvector delta*)  
*real rowvector* optimize\_init\_nmsimplexdeltas(*S*)

optimize\_init\_nmsimplexdeltas(*S*, *delta*) sets the values of *delta* to be used, along with the initial parameters, to build the simplex required by technique "nm" (Nelder–Mead). Use of this function is required only in the Nelder–Mead case. The values in *delta* must be at least 10 times larger than *ptol*, which is set by `optimize_init_conv_ptol()`. The initial simplex will be  $\{p, p + (d_1, 0), \dots, 0, p + (0, d_2, 0, \dots, 0), \dots, p + (0, 0, \dots, 0, d_k)\}$ .

optimize\_init\_nmsimplexdeltas(*S*) returns the deltas that will be (or were) used.

**optimize\_init\_argument() and optimize\_init\_narguments()**

*void*                    optimize\_init\_argument(*S*, *real scalar k*, *X*)  
*void*                    optimize\_init\_narguments(*S*, *real scalar K*)  
*pointer scalar* optimize\_init\_argument(*S*, *real scalar k*)  
*real scalar*    optimize\_init\_narguments(*S*)

optimize\_init\_argument(*S*, *k*, *X*) sets the *k*th extra argument of the evaluator function to be *X*, where *k* can only 1, 2, 3, ..., 9. *X* can be anything, including a view matrix or even a pointer to a function. No copy of *X* is made; it is a pointer to *X* that is stored, so any changes you make to *X* between setting it and *X* being used will be reflected in what is passed to the evaluator function.

optimize\_init\_narguments(*S*, *K*) sets the number of extra arguments to be passed to the evaluator function. This function is useless and included only for completeness. The number of extra arguments is automatically set as you use `optimize_init_argument()`.

optimize\_init\_argument(*S*) returns a pointer to the object that was previously set.

optimize\_init\_nargs(*S*) returns the number of extra arguments that are passed to the evaluator function.

**optimize\_init\_cluster()**

optimize\_init\_cluster(*S*, *c*) specifies a cluster variable. *c* may be a string scalar containing a Stata variable name, or *c* may be real colvector directly containing the cluster values. The default is "", meaning no clustering. If clustering is specified, the default *vcetype* becomes "robust".

**optimize\_init\_colstripe()**

optimize\_init\_colstripe(*S* [, *stripe*]) sets the string matrix to be associated with the parameter vector. See matrix colnames in [P] [matrix rownames](#).

optimize\_init\_technique()

void            optimize\_init\_technique(*S*, *string scalar technique*)

*string scalar* optimize\_init\_technique(*S*)

optimize\_init\_technique(*S*, *technique*) sets the optimization technique to be used. Current choices are

<i>technique</i>	Description
"nr"	modified Newton–Raphson
"dfp"	Davidon–Fletcher–Powell
"bfgs"	Broyden–Fletcher–Goldfarb–Shanno
"bhhh"	Berndt–Hall–Hall–Hausman
"nm"	Nelder–Mead

The default is "nr".

optimize\_init\_technique(*S*) returns the technique currently set.

Aside: All techniques require `optimize_init_params()` be set. Technique "nm" also requires that `optimize_init_nmsimplexdeltas()` be set. Parameters (and delta) can be set before or after the technique is set.

You can switch between "nr", "dfp", "bfgs", and "bhhh" by specifying two or more of them in a space-separated list. By default, `optimize()` will use an algorithm for five iterations before switching to the next algorithm. To specify a different number of iterations, include the number after the technique. For example, specifying `optimize_init_technique(M, "bhhh 10 nr 1000")` requests that `optimize()` perform 10 iterations using the Berndt–Hall–Hall–Hausman algorithm, followed by 1,000 iterations using the modified Newton–Raphson algorithm, and then switch back to Berndt–Hall–Hall–Hausman for 10 iterations, and so on. The process continues until `convergence` or until `maxiter` is exceeded.

optimize\_init\_singularHmethod()

void            optimize\_init\_singularHmethod(*S*, *string scalar singularHmethod*)

*string scalar* optimize\_init\_singularHmethod(*S*)

optimize\_init\_singularHmethod(*S*, *singularHmethod*) specifies what the optimizer should do when, at an iteration step, it finds that *H* is singular. Current choices are

<i>singularHmethod</i>	Description
"m-marquardt"	modified Marquardt algorithm
"hybrid"	mixture of steepest descent and Newton

The default is "m-marquardt" if not set;  
"hybrid" is equivalent to `ml`'s `difficult` option; see [R] `ml`.

optimize\_init\_technique(*S*) returns the *singularHmethod* currently set.



**optimize\_init\_conv\_maxiter()**

```
void      optimize_init_conv_maxiter(S, real scalar maxiter)
real scalar optimize_init_conv_maxiter(S)
```

`optimize_init_conv_maxiter(S, maxiter)` sets the maximum number of iterations to be performed before `optimize()` is stopped; results are posted to `optimize_result_*`(*S*) just as if convergence were achieved, but `optimize_result_converged()` is set to 0. The default *maxiter* if not set is `c(maxiter)`, which is probably 16,000; type `return list` in Stata to determine the current default value.

`optimize_init_conv_maxiter(S)` returns the *maxiter* currently set.

**optimize\_init\_conv\_warning()**

`optimize_init_conv_warning(S, {"on"|"off"})` specifies whether the warning message “convergence not achieved” is to be displayed when this stopping rule is invoked. The default is “on”.

**optimize\_init\_conv\_ptol(), ...\_vtol(), ...\_nrtol()**

```
void      optimize_init_conv_ptol(S, real scalar ptol)
void      optimize_init_conv_vtol(S, real scalar vtol)
void      optimize_init_conv_nrtol(S, real scalar nrtol)

real scalar optimize_init_conv_ptol(S)
real scalar optimize_init_conv_vtol(S)
real scalar optimize_init_conv_nrtol(S)
```

The two-argument form of these functions set the tolerances that control `optimize()`’s convergence criterion. `optimize()` performs iterations until the convergence criterion is met or until the number of iterations exceeds `optimize_init_conv_maxiter()`. When the convergence criterion is met, `optimize_result_converged()` is set to 1. The default values of *ptol*, *vtol*, and *nrtol* are  $1e-6$ ,  $1e-7$ , and  $1e-5$ , respectively.

The single-argument form of these functions return the current values of *ptol*, *vtol*, and *nrtol*.

*Optimization criterion:* In all cases except `optimize_init_technique(S)=="nm"`, that is, in all cases except Nelder–Mead, that is, in all cases of derivative-based maximization, the optimization criterion is defined as follows:

Define

```
C_ptol:       $\text{mreldif}(p, p_{\text{prior}}) < ptol$ 
C_vtol:       $\text{reldif}(v, v_{\text{prior}}) < vtol$ 
C_nrtol:      $g * \text{invsym}(-H) * g' < nrtol$ 
C_concave:   $-H$  is positive semidefinite
```

The above definitions apply for maximization. For minimization, think of it as maximization of  $-f(p)$ . `optimize()` declares convergence when

$$(C\_ptol \mid C\_vtol) \ \& \ C\_concave \ \& \ C\_nrtol$$

For `optimize_init_technique(S)=="nm"` (Nelder–Mead), the criterion is defined as follows:

Let  $R$  be the minimum and maximum values on the simplex and define

$$C\_ptol: \text{mreldif}(\text{vertices of } R) < ptol$$
$$C\_vtol: \text{reldif}(R) < vtol$$

`optimize()` declares successful convergence when

$$C\_ptol \mid C\_vtol$$

### **optimize\_init\_conv\_ignorenrtol()**

`optimize_init_conv_ignorenrtol(S, {"off"|"on"})` sets whether  $C\_nrtol$  should simply be treated as true in all cases, which in effects removes the  $nrtol$  criterion from the convergence rule. The default is "off".

### **optimize\_init\_iterid()**

*void*            `optimize_init_iterid(S, string scalar id)`

*string scalar* `optimize_init_iterid(S)`

By default, `optimize()` shows an iteration log, a line of which looks like

Iteration 1: f(p) = 5.7641349

See `optimize_init_tracelevel()` below.

`optimize_init_iterid(S, id)` sets the string used to label the iteration in the iteration log. The default is "Iteration".

`optimize_init_iterid(S)` returns the  $id$  currently in use.

### **optimize\_init\_valueid()**

*void*            `optimize_init_valueid(S, string scalar id)`

*string scalar* `optimize_init_valueid(S)`

By default, `optimize()` shows an iteration log, a line of which looks like

Iteration 1: f(p) = 5.7641349

See `optimize_init_tracelevel()` below.

`optimize_init_valueid(S, id)` sets the string used to identify the value. The default is "f(p)".

`optimize_init_valueid(S)` returns the  $id$  currently in use.

### **optimize\_init\_tracelevel()**

*void*            `optimize_init_tracelevel(S, string scalar tracelevel)`

*string scalar* `optimize_init_tracelevel(S)`

`optimize_init_tracelevel(S, tracelevel)` sets what is displayed in the iteration log. Allowed values of *tracelevel* are

<i>tracelevel</i>	To be displayed each iteration
"none"	nothing (suppress the log)
"value"	function value
"tolerance"	previous + convergence values
"step"	previous + stepping information
"paramdiffs"	previous + parameter relative differences
"params"	previous + parameter values
"gradient"	previous + gradient vector
"hessian"	previous + Hessian matrix

The default is "value" if not reset.

`optimize_init_tracelevel(S)` returns the value of *tracelevel* currently set.

**`optimize_init_trace_dots(), ..._value(), ..._tol(), ..._step(), ..._paramdiffs(), ..._params(), ..._gradient(), ..._Hessian()`**

`optimize_init_trace_dots(S, { "off" | "on" })` displays a dot each time your evaluator is called. The default is "off".

`optimize_init_trace_value(S, { "on" | "off" })` displays the function value at the start of each iteration. The default is "on".

`optimize_init_trace_tol(S, { "off" | "on" })` displays the value of the calculated result that is compared to the effective [convergence](#) criterion at the end of each iteration. The default is "off".

`optimize_init_trace_step(S, { "off" | "on" })` displays the steps within iteration. Listed are the value of objective function along with the word forward or backward. The default is "off".

`optimize_init_trace_paramdiffs(S, { "off" | "on" })` displays the parameter relative differences from the previous iteration that are greater than the parameter tolerance *ptol*. The default is "off".

`optimize_init_trace_params(S, { "off" | "on" })` displays the parameters at the start of each iteration. The default is "off".

`optimize_init_trace_gradient(S, { "off" | "on" })` displays the gradient vector at the start of each iteration. The default is "off".

`optimize_init_trace_Hessian(S, { "off" | "on" })` displays the Hessian matrix at the start of each iteration. The default is "off".

**`optimize_init_evaluations()`**

`optimize_init_evaluations(S, { "off" | "on" })` specifies whether the system is to count the number of times the [evaluator](#) is called. The default is "off".

**optimize\_init\_constraints()**

```
void          optimize_init_constraints(S, real matrix Cc)
real matrix  optimize_init_constraints(S)
```

$nc$  linear constraints may be imposed on the  $np$  parameters in  $p$  according to  $Cp' = c$ ,  $C$ :  $nc \times np$  and  $c$ :  $nc \times 1$ . For instance, if there are four parameters and you wish to impose the single constraint  $p_1 = p_2$ , then  $C = (1, -1, 0, 0)$  and  $c = (0)$ . If you wish to add the constraint  $p_4 = 2$ , then  $C = (1, -1, 0, 0, 0, 0, 0, 1)$  and  $c = (0 \setminus 2)$ .

`optimize_init_constraints(S, Cc)` allows you to impose such constraints where  $Cc = (C, c)$ . Use of this function is optional. If no constraints have been set, then  $Cc$  is  $0 \times (np + 1)$ .

`optimize_init_constraints(S)` returns the current  $Cc$  matrix.

**optimize\_init\_verbose()**

```
void          optimize_init_verbose(S, real scalar verbose)
real scalar  optimize_init_verbose(S)
```

`optimize_init_verbose(S, verbose)` sets whether error messages that arise during the execution of `optimize()` or `_optimize()` are to be displayed. *verbose*=1 means that they are; 0 means that they are not. The default is 1. Setting *verbose* to 0 is of interest only to users of `_optimize()`. If you wish to suppress all output, code

```
optimize_init_verbose(S, 0)
optimize_init_tracelevel(S, "none")
```

`optimize_init_verbose(S)` returns the current value of *verbose*.

**optimize()**

```
real rowvector optimize(S)
```

`optimize(S)` invokes the optimization process and returns the resulting parameter vector. If something goes wrong, `optimize()` aborts with error.

Before you can invoke `optimize()`, you must have defined your evaluator function `evaluator()` and you must have set initial values:

```
S = optimize_init()
optimize_init_evaluator(S, &evaluator())
optimize_init_params(S, (...))
```

The above assumes that your evaluator function is `d0`. Often you will also have coded

```
optimize_init_evaluatoretype(S, "...")
```

and you may have coded other `optimize_init_*`() functions as well.

Once `optimize()` completes, you may use the `optimize_result_*`() functions. You may also continue to use the `optimize_init_*`() functions to access initial settings, and you may use them to change settings and restart optimization (that is, invoke `optimize()` again) if you wish. If you do that, you will usually want to use the resulting parameter values from the first round of optimization as initial values for the second. If so, do not forget to code

```
optimize_init_params(S, optimize_result_params(S))
```

## `_optimize()`

*real scalar* `_optimize(S)`

`_optimize(S)` performs the same actions as `optimize(S)` except that, rather than returning the resulting parameter vector, `_optimize()` returns a real scalar and, rather than aborting if numerical issues arise, `_optimize()` returns a nonzero value. `_optimize()` returns 0 if all went well. The returned value is called an error code.

`optimize()` returns the resulting parameter vector  $p$ . It can work that way because optimization must have gone well. Had it not, `optimize()` would have aborted execution.

`_optimize()` returns an error code. If it is 0, optimization went well and you can obtain the parameter vector by using `optimize_result_params()`. If optimization did not go well, you can use the error code to diagnose what went wrong and take the appropriate action.

Thus `_optimize(S)` is an alternative to `optimize(S)`. Both functions do the same thing. The difference is what happens when there are numerical difficulties.

`optimize()` and `_optimize()` work around most numerical difficulties. For instance, the evaluator function you write is allowed to return  $v$  equal to missing if it cannot calculate the  $f()$  at the current values of  $p$ . If that happens during optimization, `optimize()` and `_optimize()` will back up to the last value that worked and choose a different direction. `optimize()`, however, cannot tolerate that happening with the initial values of the parameters because `optimize()` has no value to back up to. `optimize()` issues an error message and aborts, meaning that execution is stopped. There can be advantages in that. The calling program need not include complicated code for such instances, figuring that stopping is good enough because a human will know to address the problem.

`_optimize()`, however, does not stop execution. Rather than aborting, `_optimize()` returns a nonzero value to the caller, identifying what went wrong.

Programmers implementing advanced systems will want to use `_optimize()` instead of `optimize()`. Everybody else should use `optimize()`.

Programmers using `_optimize()` will also be interested in the functions

```
optimize_init_verbose()
optimize_result_errorcode()
optimize_result_errortext()
optimize_result_returncode()
```

If you perform optimization by using `_optimize()`, the behavior of all `optimize_result_*`() functions is altered. The usual behavior is that, if calculation is required and numerical problems arise, the functions abort with error. After `_optimize()`, however, a properly dimensioned missing result is returned and `optimize_result_errorcode()` and `optimize_result_errortext()` are set appropriately.

The error codes returned by `_optimize()` are listed under the heading [\*optimize\\_result\\_errorcode\(\)\*](#) below.

### `optimize_evaluate()`

```
void optimize_evaluate(S)
```

`optimize_evaluate(S)` evaluates  $f()$  at `optimize_init_params()` and posts results to `optimize_result_*`() just as if optimization had been performed, meaning that all `optimize_result_*`() functions are available for use. `optimize_result_converged()` is set to 1.

The setup for running this function is the same as for running `optimize()`:

```
S = optimize_init()
optimize_init_evaluator(S, &evaluator())
optimize_init_params(S, (...))
```

Usually, you will have also coded

```
optimize_init_evaluatoretype(S, ...))
```

The other `optimize_init_*`() settings do not matter.

### `_optimize_evaluate()`

```
real scalar _optimize_evaluate(S)
```

The relationship between `_optimize_evaluate()` and `optimize_evaluate()` is the same as that between `_optimize()` and `optimize()`; see [\*\\_optimize\(\)\*](#).

`_optimize_evaluate()` returns an error code.

### `optimize_result_params()`

```
real rowvector optimize_result_params(S)
```

`optimize_result_params(S)` returns the resulting parameter values. These are the same values that were returned by `optimize()` itself. There is no computational cost to accessing the results, so rather than coding

```
p = optimize(S)
```

if you find it more convenient to code

```
(void) optimize(S)
...
p = optimize_result_params(S)
```

then do so.

**optimize\_result\_value() and optimize\_result\_value0()**

*real scalar* optimize\_result\_value(*S*)

*real scalar* optimize\_result\_value0(*S*)

optimize\_result\_value(*S*) returns the value of  $f()$  evaluated at  $p$  equal to [optimize\\_result\\_params\(\)](#).

optimize\_result\_value0(*S*) returns the value of  $f()$  evaluated at  $p$  equal to [optimize\\_init\\_params\(\)](#).

These functions may be called regardless of the evaluator or technique used.

**optimize\_result\_gradient()**

*real rowvector* optimize\_result\_gradient(*S*)

optimize\_result\_gradient(*S*) returns the value of the gradient vector evaluated at  $p$  equal to [optimize\\_result\\_params\(\)](#). This function may be called regardless of the evaluator or technique used.

**optimize\_result\_scores()**

*real matrix* optimize\_result\_scores(*S*)

optimize\_result\_scores(*S*) returns the value of the scores evaluated at  $p$  equal to [optimize\\_result\\_params\(\)](#). This function may be called only if a type gf evaluator is used, but regardless of the technique used.

**optimize\_result\_Hessian()**

*real matrix* optimize\_result\_Hessian(*S*)

optimize\_result\_Hessian(*S*) returns the value of the Hessian matrix evaluated at  $p$  equal to [optimize\\_result\\_params\(\)](#). This function may be called regardless of the evaluator or technique used.

**optimize\_result\_V() and optimize\_result\_Vtype()**

*real matrix* optimize\_result\_V(*S*)

*string scalar* optimize\_result\_Vtype(*S*)

optimize\_result\_V(*S*) returns [optimize\\_result\\_V\\_oim\(S\)](#) or [optimize\\_result\\_V\\_opg\(S\)](#), depending on which is the natural conjugate for the optimization technique used. If there is no natural conjugate, [optimize\\_result\\_V\\_oim\(S\)](#) is returned.

optimize\_result\_Vtype(*S*) returns "oim" or "opg".

**optimize\_result\_V\_oim(), ...\_opg(), ...\_robust()**

*real matrix* optimize\_result\_V\_oim( $S$ )  
*real matrix* optimize\_result\_V\_opg( $S$ )  
*real matrix* optimize\_result\_V\_robust( $S$ )

These functions return the variance matrix of  $p$  evaluated at  $p$  equal to `optimize_result_params()`. These functions are relevant only for maximization of log-likelihood functions but may be called in any context, including minimization.

`optimize_result_V_oim( $S$ )` returns `invsym( $-H$ )`, which is the variance matrix obtained from the observed information matrix. For minimization, returned is `invsym( $H$ )`.

`optimize_result_V_opg( $S$ )` returns `invsym( $S'S$ )`, where  $S$  is the  $N \times np$  matrix of scores. This is known as the variance matrix obtained from the outer product of the gradients. `optimize_result_V_opg()` is available only when the evaluator function is type `gf`, but regardless of the technique used.

`optimize_result_V_robust( $S$ )` returns  $H * \text{invsym}(S'S) * H$ , which is the robust estimate of variance, also known as the sandwich estimator of variance. `optimize_result_V_robust()` is available only when the evaluator function is type `gf`, but regardless of the technique used.

**optimize\_result\_iterations()**

*real scalar* optimize\_result\_iterations( $S$ )

`optimize_result_iterations( $S$ )` returns the number of iterations used in obtaining results.

**optimize\_result\_converged()**

*real scalar* optimize\_result\_converged( $S$ )

`optimize_result_converged( $S$ )` returns 1 if results converged and 0 otherwise. See `optimize_init_conv_ptol()` for the definition of convergence.

**optimize\_result\_iterationlog()**

*real colvector* optimize\_result\_iterationlog( $S$ )

`optimize_result_iterationlog( $S$ )` returns a column vector of the values of  $f()$  at the start of the final 20 iterations, or, if there were fewer, however many iterations there were. Returned vector is  $\min(\text{optimize\_result\_iterations}(), 20) \times 1$ .

**optimize\_result\_evaluations()**

`optimize_result_evaluations( $S$ )` returns a  $1 \times 3$  real rowvector containing the number of times the evaluator was called, assuming `optimize_init_evaluations()` was set on. Contents are the number of times called for the purposes of 1) calculating the objective function, 2) calculating the objective function and its first derivative, and 3) calculating the objective function and its first and second derivatives. If `optimize_init_evaluations()` was set to off, returned is  $(0, 0, 0)$ .



**optimize\_result\_errorcode(), ...\_errortext(), and ...\_returncode()**

*real scalar*    optimize\_result\_errorcode(*S*)

*string scalar*    optimize\_result\_errortext(*S*)

*real scalar*    optimize\_result\_returncode(*S*)

These functions are for use after `_optimize()`.

`optimize_result_errorcode(S)` returns the error code of `_optimize()`, `_optimize_evaluate()`, or the last `optimize_result_*`() run after either of the first two functions. The value will be zero if there were no errors. The error codes are listed directly [below](#).

`optimize_result_errortext(S)` returns a string containing the error message corresponding to the error code. If the error code is zero, the string will be "".

`optimize_result_returncode(S)` returns the Stata return code corresponding to the error code. The mapping is listed directly [below](#).

In advanced code, these functions might be used as

```
(void) _optimize(S)
...
if (ec = optimize_result_code(S)) {
    errprintf("{p}\n")
    errprintf("%s\n", optimize_result_errortext(S))
    errprintf("{p_end}\n")
    exit(optimize_result_returncode(S))
    /*NOTREACHED*/
}
```

The error codes and their corresponding Stata return codes are

Error code	Return code	Error text
1	1400	initial values not feasible
2	412	redundant or inconsistent constraints
3	430	missing values returned by evaluator
4	430	Hessian is not positive semidefinite or Hessian is not negative semidefinite
5	430	could not calculate numerical derivatives—discontinuous region with missing values encountered
6	430	could not calculate numerical derivatives—flat or discontinuous region encountered
7	430	could not calculate improvement—discontinuous region encountered
8	430	could not calculate improvement—flat region encountered
9	430	Hessian could not be updated—Hessian is unstable
10	111	technique unknown
11	111	incompatible combination of techniques
12	111	singular H method unknown
13	198	matrix stripe invalid for parameter vector
14	198	negative convergence tolerance values are not allowed
15	503	invalid starting values
16	111	<code>optimize()</code> subroutine not found
17	111	simplex delta required
18	3499	simplex delta not conformable with parameter vector
19	198	simplex delta value too small (must be greater than $10 \times \text{ptol}$ in absolute value)
20	198	evaluator type requires the <code>nr</code> technique
21	198	evaluator type not allowed with specified technique
22	111	<code>optimize()</code> subroutine not found
23	198	evaluator type not allowed with <code>bhhh</code> technique
24	111	evaluator functions required
25	198	starting values for parameters required
26	198	missing parameter values not allowed
27	198	invalid evaluator type

NOTES: (1) Error 1 can occur only when evaluating  $f()$  at initial parameters.

(2) Error 2 can occur only if constraints are specified.

(3) Error 3 can occur only if the technique is `"nm"`.

(4) Error 9 can occur only if technique is `"bfgs"` or `"dfp"`.

**optimize\_query()**

```
void optimize_query(S)
```

optimize\_query(*S*) displays a report on all optimize\_init\_\*() and optimize\_result\_\*() values. optimize\_query() may be used before or after optimize() and is useful when using optimize() interactively or when debugging a program that calls optimize() or \_optimize().

**Conformability**

All functions have  $1 \times 1$  inputs and have  $1 \times 1$  or *void* outputs except the following:

```
optimize_init_params(S, initialvalues):
```

```
      S:      transmorphic
initialvalues:   $1 \times np$ 
      result:  void
```

```
optimize_init_params(S):
```

```
      S:      transmorphic
      result:   $1 \times np$ 
```

```
optimize_init_argument(S, k, X):
```

```
      S:      transmorphic
      k:       $1 \times 1$ 
      X:      anything
      result:  void
```

```
optimize_init_nmsimplexdeltas(S, delta):
```

```
      S:      transmorphic
      delta:   $1 \times np$ 
      result:  void
```

```
optimize_init_nmsimplexdeltas(S):
```

```
      S:      transmorphic
      result:   $1 \times np$ 
```

```
optimize_init_constraints(S, Cc):
```

```
      S:      transmorphic
      Cc:      $nc \times (np + 1)$ 
      result:  void
```

```
optimize_init_constraints(S):
```

```
      S:      transmorphic
      result:   $nc \times (np + 1)$ 
```

```
optimize(S):
```

```
      S:      transmorphic
      result:   $1 \times np$ 
```

```
optimize_result_params(S):
```

```
      S:      transmorphic
      result:   $1 \times np$ 
```

optimize\_result\_gradient( $S$ ), optimize\_result\_evaluations( $S$ ):

$S$ : *transmorphic*  
result:  $1 \times np$

optimize\_result\_scores( $S$ ):

$S$ : *transmorphic*  
result:  $N \times np$

optimize\_result\_Hessian( $S$ ):

$S$ : *transmorphic*  
result:  $np \times np$

optimize\_result\_V( $S$ ), optimize\_result\_V\_oim( $S$ ), optimize\_result\_V\_opg( $S$ ),  
optimize\_result\_V\_robust( $S$ ):

$S$ : *transmorphic*  
result:  $np \times np$

optimize\_result\_iterationlog( $S$ ):

$S$ : *transmorphic*  
result:  $L \times 1, L \leq 20$

For optimize\_init\_cluster( $S, c$ ) and optimize\_init\_colstripe( $S$ ), see [Syntax](#) above.

## Diagnostics

All functions abort with error when used incorrectly.

optimize() aborts with error if it runs into numerical difficulties. \_optimize() does not; it instead returns a nonzero error code.

optimize\_evaluate() aborts with error if it runs into numerical difficulties.  
\_optimize\_evaluate() does not; it instead returns a nonzero error code.

The optimize\_result\_\*( $S$ ) functions abort with error if they run into numerical difficulties when called after optimize() or optimize\_evaluate(). They do not abort when run after \_optimize() or \_optimize\_evaluate(). They instead return a properly dimensioned missing result and set optimize\_result\_errorcode() and optimize\_result\_errortext().

The formula  $x_{i+1} = x_i - f(x_i)/f'(x_i)$  and its generalizations for solving  $f(x) = 0$  (and its generalizations) are known variously as Newton's method or the Newton–Raphson method. The real history is more complicated than these names imply and has roots in the earlier work of Arabic algebraists and François Viète.

Newton's first formulation dating from about 1669 refers only to solution of polynomial equations and does not use calculus. In his *Philosophiae Naturalis Principia Mathematica*, first published in 1687, the method is used, but not obviously, to solve a nonpolynomial equation. Raphson's work, first published in 1690, also concerns polynomial equations, and proceeds algebraically without using calculus, but lays more stress on iterative calculation and so is closer to present ideas. It was not until 1740 that Thomas Simpson published a more general version explicitly formulated in calculus terms that was applied to both polynomial and nonpolynomial equations and to both single equations and systems of equations. Simpson's work was in turn overlooked in influential later accounts by Lagrange and Fourier, but his contribution also deserves recognition.

Isaac Newton (1643–1727) was an English mathematician, astronomer, physicist, natural philosopher, alchemist, theologian, biblical scholar, historian, politician and civil servant. He was born in Lincolnshire and later studied there and at the University of Cambridge, where he was a fellow of Trinity College and elected Lucasian Professor in 1669. Newton demonstrated the generalized binomial theorem, did major work on power series, and deserves credit with Gottfried Leibniz for the development of calculus. They entered a longstanding priority dispute in 1711, which lasted until Leibniz died in 1716.

Newton described universal gravitation and the laws of motion central to classical mechanics and showed that the motions of objects on Earth and beyond are subject to the same laws. Newton invented the reflecting telescope and developed a theory of color that was based on the fact that a prism splits white light into a visible spectrum. He also studied cooling and the speed of sound and proposed a theory of the origin of stars. Much of his later life was spent in London, including brief spells as member of Parliament and longer periods as master of the Mint and president of the Royal Society. He was knighted in 1705. Although undoubtedly one of the greatest mathematical and scientific geniuses of all time, Newton was also outstandingly contradictory, secretive, and quarrelsome.

Joseph Raphson (1648–1715) was an English or possibly Irish mathematician. No exact dates are known for his birth or death years. He appears to have been largely self-taught and was awarded a degree by the University of Cambridge after the publication of his most notable work, *Analysis Aequationum Universalis* (1690), and his election as a fellow of the Royal Society.

Thomas Simpson (1710–1761) was born in Market Bosworth, Leicestershire, England. Although he lacked formal education, he managed to teach himself mathematics. Simpson moved to London and worked as a teacher in London coffee houses (as did De Moivre) and then at the Royal Military Academy at Woolwich. He published texts on calculus, astronomy, and probability. His legacy includes work on interpolation and numerical methods of integration; namely Simpson's Rule, which Simpson learned from Newton. Simpson was also a fellow of the Royal Society.

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## Also see

[M-5] **`moptimize()`** — Model optimization

[M-4] **mathematical** — Important mathematical functions

[M-4] **statistical** — Statistical functions

## Syntax

```

        info = panelsetup(V, idcol)
        info = panelsetup(V, idcol, minobs)
        info = panelsetup(V, idcol, minobs, maxobs)

real rowvector  panelstats(info)
real matrix     panelsubmatrix(V, i, info)
void            panelsubview(SV, V, i, info)
    
```

where,

<i>V</i> :	<i>real</i> or <i>string</i> matrix, possibly a view
<i>idcol</i> :	<i>real</i> scalar
<i>minobs</i> :	<i>real</i> scalar
<i>maxobs</i> :	<i>real</i> scalar
<i>info</i> :	<i>real</i> matrix
<i>i</i> :	<i>real</i> scalar
<i>SV</i> :	matrix to be created, possibly as view

## Description

These functions assist with the processing of panel data. The idea is to make it easy and fast to write loops like

```

for (i=1; i<=number_of_panels; i++) {
    X = matrix corresponding to panel i
    ...
    ... (calculations using X) ...
    ...
}
    
```

Using these functions, this loop could become

```

st_view(Vid, ., "idvar",      "touse")
st_view(V,   ., ("x1", "x2"), "touse")
info = panelsetup(Vid, 1)
for (i=1; i<=rows(info); i++) {
    X = panelsubmatrix(V, i, info)
    ...
    ... (calculations using X) ...
    ...
}
    
```

`panelsetup(V, idcol, ...)` sets up panel processing. It returns a matrix (*info*) that is passed to other panel-processing functions.

`panelstats(info)` returns a row vector containing the number of panels, number of observations, minimum number of observations per panel, and maximum number of observations per panel.

`panelsubmatrix(V, i, info)` returns a matrix containing the contents of *V* for panel *i*.

`panelsubview(SV, V, i, info)` does nearly the same thing. Rather than returning a matrix, however, it places the matrix in *SV*. If *V* is a view, then the matrix placed in *SV* will be a view.

## Remarks

Remarks are presented under the following headings:

- Definition of panel data
- Definition of problem
- Preparation
- Use of `panelsetup()`
- Using `panelstats()`
- Using `panelsubmatrix()`
- Using `panelsubview()`

## Definition of panel data

Panel data include multiple observations on subjects, countries, etc.:

Subject ID	Time ID	x1	x2
1	1	4.2	3.7
1	2	3.2	3.7
1	3	9.2	4.2
2	1	1.7	4.0
2	2	1.9	5.0
3	1	9.5	1.3
⋮	⋮	⋮	⋮

In the above dataset, there are three observations for subject 1, two for subject 2, etc. We labeled the identifier within subject to be time, but that is only suggestive, and in any case, the secondary identifier will play no role in what follows.

If we speak about the first panel, we are discussing the first 3 observations of this dataset. If we speak about the second, that corresponds to observations 4 and 5.

It is common to refer to panel numbers with the letter *i*. It is common to refer to the number of observations in the *i*th panel as  $T_i$  even when the data within panel have nothing to do with repeated observations over time.



## Definition of problem

We want to calculate some statistic on panel data. The calculation amounts to

$$\sum_{i=1}^K f(X_i)$$

where the sum is performed across panels, and  $X_i$  is the data matrix for panel  $i$ . For instance, given the example in the previous section

$$X_1 = \begin{bmatrix} 4.2 & 3.7 \\ 3.2 & 3.7 \\ 9.2 & 4.2 \end{bmatrix}$$

and  $X_2$  is a similarly constructed  $2 \times 2$  matrix.

Depending on the nature of the calculation, there will be problems for which

1. we want to use all the panels,
2. we want to use only panels for which there are two or more observations, and
3. we want to use the same number of observations in all the panels (balanced panels).

In addition to simple problems of the sort,

$$\sum_{i=1}^K f(X_i)$$

you may also need to deal with problems of the form,

$$\sum_{i=1}^K f(X_i, Y_i, \dots)$$

That is, you may need to deal with problems where there are multiple matrices per subject.

We use the sum operator purely for illustration, although it is the most common. Your problem might be

$$F(X_1, Y_1, \dots, X_2, Y_2, \dots)$$

## Preparation

Before using the functions documented here, create a matrix or matrices containing the data. For illustration, it will be sufficient to create  $V$  containing all the data in our example problem:

$$V = \begin{bmatrix} 1 & 1 & 4.2 & 3.7 \\ 1 & 2 & 3.2 & 3.7 \\ 1 & 3 & 9.2 & 4.2 \\ 2 & 1 & 1.7 & 4.0 \\ 2 & 2 & 1.9 & 5.0 \\ 3 & 1 & 9.5 & 1.3 \\ \vdots & \vdots & \vdots & \vdots \end{bmatrix}$$

But you will probably find it more convenient (and we recommend) if you create at least two matrices, one containing the subject identifier and the other containing the  $x$  variables (and omit the within-subject "time" identifier altogether):

$$V1 = \begin{bmatrix} 1 \\ 1 \\ 1 \\ 2 \\ 2 \\ 3 \\ \vdots \end{bmatrix} \qquad V2 = \begin{bmatrix} 4.2 & 3.7 \\ 3.2 & 3.7 \\ 9.2 & 4.2 \\ 1.7 & 4.0 \\ 1.9 & 5.0 \\ 9.5 & 1.3 \\ \vdots & \vdots \end{bmatrix}$$

In the above, matrix  $V1$  contains the subject identifier, and matrix  $V2$  contains the data for all the  $X_i$  matrices in

$$\sum_{i=1}^K f(X_i)$$

If your calculation is

$$\sum_{i=1}^K f(X_i, Y_i, \dots)$$

create additional  $V$  matrices,  $V3$  corresponding to  $Y_i$ , and so on.

To create these matrices, use [\[M-5\] st\\_view\(\)](#)

```
st_view(V1, ., "idvar", "touse")
st_view(V2, ., ("x1", "x2"), "touse")
```

although you could use [\[M-5\] st\\_data\(\)](#) if you preferred. Using `st_view()` will save memory. You can also construct  $V1$ ,  $V2$ , ..., however you wish; they are just matrices. Be sure that the matrices align, for example, that row 4 of one matrix corresponds to row 4 of another. We did that above by assuming a *touse* variable had been included (or constructed) in the dataset.

## Use of panelsetup()

`panelsetup(V, idcol, ...)` sets up panel processing, returning a  $K \times 2$  matrix that contains a row for each panel. The row records the first and last observation numbers (row numbers in  $V$ ) that correspond to the panel.

For instance, with our example, `panelsetup()` will return

$$\begin{bmatrix} 1 & 3 \\ 4 & 5 \\ 6 & 9 \\ \vdots & \vdots \end{bmatrix}$$

The first panel is recorded in observations 1 to 3; it contains  $3 - 1 + 1 = 3$  observations. The second panel is recorded in observations 4 to 5 and it contains  $5 - 4 + 1 = 2$  observations, and so on. We recorded the third panel as being observations 6 to 9, although we did not show you enough of the original data for you to know that 9 was the last observation with ID 3.

`panelsetup()` has many more capabilities in constructing this result, but it is important to appreciate that returning this observation-number matrix is all that `panelsetup()` does. This matrix is all that other panel functions need to know. They work with the information produced by `panelsetup()`, but they will equally well work with any two-column matrix that contains observation numbers. Correspondingly, `panelsetup()` engages in no secret behavior that ties up memory, puts you in a mode, or anything else. `panelsetup()` merely produces this matrix.

The number of rows of the matrix `panelsetup()` returns equals  $K$ , the number of panels.

The syntax of `panelsetup()` is

```
info = panelsetup(V, idcol, minobs, maxobs)
```

The last two arguments are optional.

The required argument  $V$  specifies a matrix containing at least the panel identification numbers and required argument  $idcol$  specifies the column of  $V$  that contains that ID. Here we will use the matrix  $VI$ , which contains only the identification number:

```
info = panelsetup(VI, 1)
```

The two optional arguments are *minobs* and *maxobs*. *minobs* specifies the minimum number of observations within panel that we are willing to tolerate; if a panel has fewer observations, we want to omit it entirely. For instance, were we to specify

```
info = panelsetup(VI, 1, 3)
```

then the matrix `panelsetup()` would contain fewer rows. In our example, the returned *info* matrix would contain

$$\begin{bmatrix} 1 & 3 \\ 6 & 9 \\ \vdots & \vdots \end{bmatrix}$$

Observations 4 and 5 are now omitted because they correspond to a two-observation panel, and we said only panels with three or more observations should be included.

We chose three as a demonstration. In fact, it is most common to code

```
info = panelsetup(VI, 1, 2)
```

because that eliminates the singletons (panels with one observation).

The final optional argument is *maxobs*. For example,

```
info = panelsetup(VI, 1, 2, 5)
```

means to include only up to five observations per panel. Any observations beyond five are to be trimmed. If we code

```
info = panelsetup(VI, 1, 3, 3)
```

then all the panels contained in *info* would have three observations. If a panel had fewer than three observations, it would be omitted entirely. If a panel had more than three observations, only the first three would be included.

Panel datasets with the same number of observations per panel are said to be balanced. `panelsetup()` also provides panel-balancing capabilities. If you specify *maxobs* as 0, then

1. `panelsetup()` first calculates the  $\min(T_i)$  among the panels with *minobs* observations or more. Call that number *m*.
2. `panelsetup()` then returns `panelsetup(VI, idcol, m, m)`, thus creating balanced panels of size *m* and producing a dataset that has the maximum number of within-panel observations given it has the maximum number of panels.

If we coded

```
info = panelsetup(VI, 1, 2, 0)
```

then `panelsetup()` would create the maximum number of panels with the maximum number of within-panel observations subject to the constraint of no singletons and the panels being balanced.

## Using `panelstats()`

`panelstats(info)` can be used on any two-column matrix that contains observation numbers. `panelstats()` returns a row vector containing

```
panelstats()[1] = number of panels (same as rows(info))
panelstats()[2] = number of observations
panelstats()[3] =  $\min(T_i)$ 
panelstats()[4] =  $\max(T_i)$ 
```

## Using `panelsubmatrix()`

Having created an *info* matrix using `panelsetup()`, you can obtain the matrix corresponding to the *i*th panel using

```
X = panelsubmatrix(V, i, info)
```

It is not necessary that `panelsubmatrix()` be used with the same matrix that was used to produce *info*. We created matrix *VI* containing the ID numbers, and we created matrix *V2* containing the *x* variables

```
st_view(VI, ., "idvar", "touse")
st_view(V2, ., ("x1", "x2"), "touse")
```

and we create *info* using *VI*:

```
info = panelsetup(VI, 1)
```

We can now create the corresponding  $X$  matrix by coding

```
X = panelsubmatrix(V2, i, info)
```

and, had we created a  $V3$  matrix corresponding to  $Y_i$ , we could also code

```
Y = panelsubmatrix(V3, i, info)
```

and so on.

## Using panelsubview()

`panelsubview()` works much like `panelsubmatrix()`. The difference is that rather than coding

```
X = panelsubmatrix(V, i, info)
```

you code

```
panelsubview(X, V, i, info)
```

The matrix to be defined becomes the first argument of `panelsubview()`. That is because `panelsubview()` is designed especially to work with views. `panelsubmatrix()` will work with views, but `panelsubview()` does something special. Rather than returning an ordinary matrix (an array, in the jargon), if  $V$  is a view, `panelsubview()` returns a view in its first argument. Views save memory.

Views can save much memory, so it would seem that you would always want to use `panelsubview()` in place of `panelsubmatrix()`. What is not always appreciated, however, is that it takes Mata longer to access the data recorded in views, and so there is a tradeoff.

If the panels are likely to be large, you want to use `panelsubview()`. Conserving memory trumps all other considerations.

In fact, the panels that occur in most datasets are not that large, even when the dataset itself is. If you are going to make many calculations on  $X$ , you may wish to use `panelsubmatrix()`.

Both `panelsubmatrix()` and `panelsubview()` work with view and nonview matrices. `panelsubview()` produces a regular matrix when the base matrix  $V$  is not a view, just as does `panelsubmatrix()`. The difference is that `panelsubview()` will produce a view when  $V$  is a view, whereas `panelsubmatrix()` always produces a nonview matrix.

## Conformability

```
panelsetup(V, idcol, minobs, maxobs):
```

<i>V</i> :	$r \times c$	
<i>idcol</i> :	$1 \times 1$	
<i>minobs</i> :	$1 \times 1$	(optional)
<i>maxobs</i> :	$1 \times 1$	(optional)
<i>result</i> :	$K \times 2$ ,	$K = \text{number of panels}$

```
panelstats(info):
```

<i>info</i> :	$K \times 2$
<i>result</i> :	$1 \times 4$

```

panelsubmatrix(V, i, info):
    V:       $r \times c$ 
    i:       $1 \times 1, \quad 1 \leq i \leq \text{rows}(info)$ 
    info:    $K \times 2$ 
    result:  $t \times c, \quad t = \text{number of obs. in panel}$ 

panelsubview(SV, V, i, info):
    input:
        SV:      irrelevant
        V:       $r \times c$ 
        i:       $1 \times 1, \quad 1 \leq i \leq \text{rows}(info)$ 
        info:    $K \times 2$ 
        result:  $t \times c, \quad t = \text{number of obs. in panel}$ 
    output:
        SV:       $t \times c, \quad t = \text{number of obs. in panel}$ 

```

## Diagnostics

`panelsubmatrix(V, i, info)` and `panelsubview(SV, V, i, info)` abort with error if  $i < 1$  or  $i > \text{rows}(info)$ .

`panelsetup()` can return a  $0 \times 2$  result.

## Also see

[\[M-4\] utility](#) — Matrix utility functions

## Syntax

<i>string scalar</i>	<code>pathjoin(string scalar path1, string scalar path2)</code>
<i>void</i>	<code>pathsplrit(string scalar path, path1, path2)</code>
<i>string scalar</i>	<code>pathbasename(string scalar path)</code>
<i>string scalar</i>	<code>pathsuffix(string scalar path)</code>
<i>string scalar</i>	<code>pathrmsuffix(string scalar path)</code>
<i>real scalar</i>	<code>pathisurl(string scalar path)</code>
<i>real scalar</i>	<code>pathisabs(string scalar path)</code>
<i>real scalar</i>	<code>pathasciisuffix(string scalar path)</code>
<i>real scalar</i>	<code>pathstata suffix(string scalar path)</code>
<i>string rowvector</i>	<code>pathlist(string scalar dirlist)</code>
<i>string rowvector</i>	<code>pathlist()</code>
<i>string rowvector</i>	<code>pathsubsysdir(string rowvector pathlist)</code>
<i>string rowvector</i>	<code>pathsearchlist(string scalar fn)</code>

## Description

`pathjoin(path1, path2)` forms, logically speaking, *path1/path2*, but does so in the appropriate style. For instance, *path1* might be a URL and *path2* a Windows *dirname\filename*, and the two paths will, even so, be joined correctly. All issues of whether *path1* ends with a directory separator, *path2* begins with one, etc., are handled automatically.

`pathsplrit(path, path1, path2)` performs the inverse operation, removing the last element of the path (which is typically a filename) and storing it in *path2* and storing the rest in *path1*.

`pathbasename(path)` returns the last element of *path*.

`pathsuffix(path)` returns the file suffix, with leading dot, if there is one, and returns "" otherwise. For instance, `pathsuffix("this\that.ado")` returns ".ado".

`pathrmsuffix(path)` returns *path* with the suffix removed, if there was one. For instance, `pathrmsuffix("this\that.ado")` returns "this\that".

`pathisurl(path)` returns 1 if *path* is a URL and 0 otherwise.

`pathisabs(path)` returns 1 if *path* is absolute and 0 if relative. `c:\this` is an absolute path. `this\that` is a relative path. URLs are considered to be absolute.

`pathasciisuffix(path)` and `pathstatastatusuffix(path)` are more for StataCorp use than anything else. `pathasciisuffix()` returns 1 if the file is known to be ASCII, based on its file suffix. StataCorp uses this function in Stata's `net` command to decide whether end-of-line characters, which differ across operating systems, should be modified during downloads. `pathstatastatusuffix()` is the function used by Stata's `net` and `update` commands to decide whether a file belongs in the official directories. `pathstatastatusuffix("example.ado")` is true, but `pathstatastatusuffix("example.do")` is false because do-files do not go in system directories.

`pathlist(dirlist)` returns a row vector, each element of which contains an element of a semicolon-separated path list *dirlist*. For instance, `pathlist("a;b;c")` returns ("a", "b", "c").

`pathlist()` without arguments returns `pathlist(c("adopath"))`, the broken-out elements of the official Stata ado-path.

`pathsubsysdir(pathlist)` returns *pathlist* with any elements that are Stata system directories' shorthands, such as `UPDATES`, `PLUS`, `PERSONAL`, substituted with the actual directory names. For instance, the right way to obtain the official directories over which Stata searches for files is `pathsubsysdir(pathlist())`.

`pathsearchlist(fn)` returns a row vector. The elements are full paths/filenames specifying all the locations, in order, where Stata would look for *fn* along the official Stata ado-path.

## Remarks

Using these functions, you are more likely to produce code that works correctly regardless of operating system.



## Conformability

`pathjoin(path1, path2):`

*path1:*       $1 \times 1$

*path2:*       $1 \times 1$

*result:*       $1 \times 1$

`pathsplit(path, path1, path2):`

*input:*

*path:*       $1 \times 1$

*output:*

*path1:*       $1 \times 1$

*path2:*       $1 \times 1$

`pathbasename(path), pathsuffix(path), pathrmsuffix(path):`

*path:*       $1 \times 1$

*result:*       $1 \times 1$

`pathisurl(path), pathisabs(path), pathasciisuffix(path), pathstata suffix(path):`

*path:*       $1 \times 1$

*result:*       $1 \times 1$

`pathlist(dirlist):`

*dirlist:*       $1 \times 1$  (optional)

*result:*       $1 \times k$

`pathsubsysdir(pathlist):`

*pathlist:*       $1 \times k$

*result:*       $1 \times k$

`pathsearchlist(fn):`

*fn:*       $1 \times 1$

*result:*       $1 \times k$

## Diagnostics

All routines abort with error if the path is too long for the operating system; nothing else causes abort with error.

## Also see

[M-4] [io](#) — I/O functions

# Title

[M-5] `pinv()` — Moore–Penrose pseudoinverse

## Syntax

<i>numeric matrix</i>	<code>pinv(numeric matrix A)</code>
<i>numeric matrix</i>	<code>pinv(numeric matrix A, rank)</code>
<i>numeric matrix</i>	<code>pinv(numeric matrix A, rank, real scalar tol)</code>
<i>real scalar</i>	<code>_pinv(numeric matrix A)</code>
<i>real scalar</i>	<code>_pinv(numeric matrix A, real scalar tol)</code>

where the type of *rank* is irrelevant; the rank of *A* is returned there.

To obtain a generalized inverse of a symmetric matrix with a different normalization, see [M-5] `invsym()`.

## Description

`pinv(A)` returns the unique Moore–Penrose pseudoinverse of real or complex, symmetric or non-symmetric, square or nonsquare matrix *A*.

`pinv(A, rank)` does the same thing, and it returns in *rank* the rank of *A*.

`pinv(A, rank, tol)` does the same thing, and it allows you to specify the tolerance used to determine the rank of *A*, which is also used in the calculation of the pseudoinverse. See [M-5] `svsolve()` and [M-1] **tolerance** for information on the optional *tol* argument.

`_pinv(A)` and `_pinv(A, tol)` do the same thing as `pinv()`, except that *A* is replaced with its inverse and the rank is returned.

## Remarks

The Moore–Penrose pseudoinverse is also known as the Moore–Penrose inverse and as the generalized inverse. Whatever you call it, the pseudoinverse  $A^*$  of *A* satisfies four conditions,

$$\begin{aligned}A(A^*)A &= A \\(A^*)A(A^*) &= A^* \\(A(A^*))' &= A(A^*) \\((A^*)A)' &= (A^*)A\end{aligned}$$

where the transpose operator  $'$  is understood to mean the conjugate transpose when *A* is complex. Also, if *A* is of full rank, then

$$A^* = A^{-1}$$

`pinv(A)` is logically equivalent to `svsolve(A, I(rows(A)))`; see [M-5] `svsolve()` for details and for use of the optional *tol* argument.

## Conformability

`pinv(A, rank, tol)`:

*input*:

*A*:  $r \times c$

*tol*:  $1 \times 1$  (optional)

*output*:

*rank*:  $1 \times 1$  (optional)

*result*:  $c \times r$

`_pinv(A, tol)`:

*input*:

*A*:  $r \times c$

*tol*:  $1 \times 1$  (optional)

*output*:

*A*:  $c \times r$

*result*:  $1 \times 1$  (containing rank)

## Diagnostics

The inverse returned by these functions is real if *A* is real and is complex if *A* is complex.

`pinv(A, rank, tol)` and `_pinv(A, tol)` return missing results if *A* contains missing values.

`pinv()` and `_pinv()` also return missing values if the algorithm for computing the SVD, [M-5] `svd()`, fails to converge. This is a near zero-probability event. Here *rank* also is returned as missing.

See [M-5] `svsolve()` and [M-1] **tolerance** for information on the optional *tol* argument.

## References

- James, I. 2002. *Remarkable Mathematicians: From Euler to von Neumann*. Cambridge: Cambridge University Press.
- Moore, E. H. 1920. On the reciprocal of the general algebraic matrix. *Bulletin of the American Mathematical Society* 26: 394–395.
- Penrose, R. 1955. A generalized inverse for matrices. *Mathematical Proceedings of the Cambridge Philosophical Society* 51: 406–413.

## Also see

[M-5] `invsym()` — Symmetric real matrix inversion

[M-5] `cholinv()` — Symmetric, positive-definite matrix inversion

[M-5] `luinv()` — Square matrix inversion

[M-5] `qrinv()` — Generalized inverse of matrix via QR decomposition

[M-5] `svd()` — Singular value decomposition

[M-5] `fullsvd()` — Full singular value decomposition

[M-4] `matrix` — Matrix functions

[M-4] `solvers` — Functions to solve  $AX=B$  and to obtain  $A$  inverse

Eliakim Hastings Moore (1862–1932) was born in Marietta, Ohio. He studied mathematics and astronomy at Yale and was awarded a Ph.D. for a thesis on  $n$ -dimensional geometry. After a year studying in Germany and teaching posts at Northwestern and Yale, he settled at the University of Chicago in 1892. Moore worked on algebra, including fields and groups, the foundations of geometry and the foundations of analysis, algebraic geometry, number theory, and integral equations. He was an inspiring teacher and a great organizer in American mathematics, playing an important part in the early years of the American Mathematical Society.

Roger Penrose (1931– ) was born in Colchester in England. His father was a statistically minded medical geneticist and his mother was a doctor. Penrose studied mathematics at University College London and Cambridge and published an article on generalized matrix inverses in 1955. He taught and researched at several universities in Great Britain and the United States before being appointed Rouse Ball Professor of Mathematics at Oxford in 1973. Penrose is perhaps best known for papers ranging from cosmology and general relativity (including work with Stephen Hawking) to pure mathematics (including results on tilings of the plane) and for semipopular and wide-ranging books making controversial connections between physics, computers, mind, and consciousness. He was knighted in 1994.

## Syntax

<i>numeric vector</i>	<code>polyeval(numeric rowvector <i>c</i>, numeric vector <i>x</i>)</code>
<i>numeric rowvector</i>	<code>polysolve(numeric vector <i>y</i>, numeric vector <i>x</i>)</code>
<i>numeric rowvector</i>	<code>polytrim(numeric vector <i>c</i>)</code>
<i>numeric rowvector</i>	<code>polyderiv(numeric rowvector <i>c</i>, real scalar <i>i</i>)</code>
<i>numeric rowvector</i>	<code>polyinteg(numeric rowvector <i>c</i>, real scalar <i>i</i>)</code>
<i>numeric rowvector</i>	<code>polyadd(numeric rowvector <i>c</i><sub>1</sub>, numeric rowvector <i>c</i><sub>2</sub>)</code>
<i>numeric rowvector</i>	<code>polymult(numeric rowvector <i>c</i><sub>1</sub>, numeric rowvector <i>c</i><sub>2</sub>)</code>
<i>void</i>	<code>polydiv(numeric rowvector <i>c</i><sub>1</sub>, numeric rowvector <i>c</i><sub>2</sub>, <i>c</i><sub>q</sub>, <i>c</i><sub>r</sub>)</code>
<i>complex rowvector</i>	<code>polyroots(numeric rowvector <i>c</i>)</code>

In the above, row vector *c* contains the coefficients for a `cols(c) - 1` degree polynomial. For instance,

$$c = (4, 2, 1)$$

records the polynomial

$$4 + 2x + x^2$$

## Description

`polyeval(c, x)` evaluates polynomial *c* at each value recorded in *x*, returning the results in a p-conformable-with-*x* vector. For instance, `polyeval((4,2,1), (3\5))` returns `(4+2*3+3^2 \ 4+2*5+5^2) = (19\39)`.

`polysolve(y, x)` returns the minimal-degree polynomial *c* fitting *y* = `polyeval(c, x)`. Solution is via Lagrange's interpolation formula.

`polytrim(c)` returns polynomial *c* with trailing zeros removed. For instance, `polytrim((1,2,3,0))` returns `(1,2,3)`. `polytrim((0,0,0,0))` returns `(0)`. Thus if *n* = `cols(polytrim(c))`, then *c* records an (*n* - 1)th degree polynomial.

`polyderiv(c, i)` returns the polynomial that is the *i*th derivative of polynomial *c*. For instance, `polyderiv((4,2,1), 1)` returns `(2,2)` (the derivative of  $4 + 2x + x^2$  is  $2 + 2x$ ). The value of the first derivative of polynomial *c* at *x* is `polyeval(polyderiv(c,1), x)`.

`polyinteg(c, i)` returns the polynomial that is the *i*th integral of polynomial *c*. For instance, `polyinteg((4,2,1), 1)` returns `(0,4,1,.3333)` (the integral of  $4 + 2x + x^2$  is  $0 + 4x + x^2 + .3333x^3$ ). The value of the integral of polynomial *c* at *x* is `polyeval(polyinteg(c,1), x)`.

`polyadd( $c_1$ ,  $c_2$ )` returns the polynomial that is the sum of the polynomials  $c_1$  and  $c_2$ . For instance, `polyadd((2,1), (3,5,1))` is (5,6,1) (the sum of  $2 + x$  and  $3 + 5x + x^2$  is  $5 + 6x + x^2$ ).

`polymult( $c_1$ ,  $c_2$ )` returns the polynomial that is the product of the polynomials  $c_1$  and  $c_2$ . For instance, `polymult((2,1), (3,5,1))` is (6,13,7,1) (the product of  $2 + x$  and  $3 + 5x + x^2$  is  $6 + 13x + 7x^2 + x^3$ ).

`polydiv( $c_1$ ,  $c_2$ ,  $c_q$ ,  $c_r$ )` calculates polynomial  $c_1/c_2$ , storing the quotient polynomial in  $c_q$  and the remainder polynomial in  $c_r$ . For instance, `polydiv((3,5,1), (2,1),  $c_q$ ,  $c_r$ )` returns  $c_q=(3,1)$  and  $c_r=(-3)$ ; that is,

$$\frac{3 + 5x + x^2}{2 + x} = 3 + x \text{ with a remainder of } -3$$

or

$$3 + 5x + x^2 = (3 + x)(2 + x) - 3$$

`polyroots( $c$ )` find the roots of polynomial  $c$  and returns them in complex row vector (complex even if  $c$  is real). For instance, `polyroots((3,5,1))` returns (-4.303+0i, -.697+0i) (the roots of  $3 + 5x + x^2$  are -4.303 and -.697).

## Remarks

Given the real or complex coefficients  $c$  that define an  $n - 1$  degree polynomial in  $x$ , `polyroots( $c$ )` returns the  $n - 1$  roots for which

$$0 = c_1 + c_2x^1 + c_3x^2 + \cdots + c_nx^{n-1}$$

`polyroots( $c$ )` obtains the roots by calculating the eigenvalues of the companion matrix. The  $(n - 1) \times (n - 1)$  companion matrix for the polynomial defined by  $c$  is

$$C = \begin{bmatrix} -c_{n-1}s & -c_{n-2}s & \cdots & -c_2s & -c_1s \\ 1 & 0 & \cdots & 0 & 0 \\ 0 & 1 & \cdots & 0 & 0 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 0 & 0 & \cdots & 1 & 0 \\ 0 & 0 & \cdots & 0 & 1 \end{bmatrix}$$

where  $s = 1/c_n$  if  $c$  is real and

$$s = C \left( \frac{\operatorname{Re}(c_n)}{\operatorname{Re}(c_n)^2 + \operatorname{Im}(c_n)^2}, \frac{-\operatorname{Im}(c_n)}{\operatorname{Re}(c_n)^2 + \operatorname{Im}(c_n)^2} \right)$$

otherwise.

As in all nonsymmetric eigenvalue problems, the returned roots are complex and sorted from largest to smallest, see [M-5] **eigensystem()**.

## Conformability

`polyeval(c, x):`

*c:*  $1 \times n, n > 0$   
*x:*  $r \times 1$  or  $1 \times c$   
*result:*  $r \times 1$  or  $1 \times c$

`polysolve(y, x):`

*y:*  $n \times 1$  or  $1 \times n, n \geq 1$   
*x:*  $n \times 1$  or  $1 \times n$   
*result:*  $1 \times k, 1 \leq k \leq n$

`polytrim(c):`

*c:*  $1 \times n$   
*result:*  $1 \times k, 1 \leq k \leq n$

`polyderiv(c, i):`

*c:*  $1 \times n, n > 0$   
*i:*  $1 \times 1, i$  may be negative  
*result:*  $1 \times \max(1, n - i)$

`polyinteg(c, i):`

*c:*  $1 \times n, n > 0$   
*i:*  $1 \times 1, i$  may be negative  
*result:*  $1 \times \max(1, n + i)$

`polyadd(c1, c2):`

*c<sub>1</sub>:*  $1 \times n_1, n_1 > 0$   
*c<sub>2</sub>:*  $1 \times n_2, n_2 > 0$   
*result:*  $1 \times \max(n_1, n_2)$

`polymult(c1, c2):`

*c<sub>1</sub>:*  $1 \times n_1, n_1 > 0$   
*c<sub>2</sub>:*  $1 \times n_2, n_2 > 0$   
*result:*  $1 \times n_1 + n_2 - 1$

`polydiv(c1, c2, cq, cr):`

*input:*  
*c<sub>1</sub>:*  $1 \times n_1, n_1 > 0$   
*c<sub>2</sub>:*  $1 \times n_2, n_2 > 0$   
*output:*  
*c<sub>q</sub>:*  $1 \times k_1, 1 \leq k_1 \leq \max(n_1 - n_2 + 1, 1)$   
*c<sub>r</sub>:*  $1 \times k_2, 1 \leq k_2 \leq \max(n_1 - n_2, 1)$

`polyroots(c):`

*c:*  $1 \times n_i, n > 0$   
*result:*  $1 \times k - 1, k = \text{cols}(\text{polytrim}(c))$

## Diagnostics

All functions abort with an error if a polynomial coefficient row vector is void, but they do not necessarily give indicative error messages as to the problem. Polynomial coefficient row vectors may contain missing values.

`polyderiv( $c$ ,  $i$ )` returns  $c$  when  $i = 0$ . It returns `polyinteg( $c$ ,  $-i$ )` when  $i < 0$ . It returns `(0)` when  $i$  is missing (think of missing as positive infinity).

`polyinteg( $c$ ,  $i$ )` returns  $c$  when  $i = 0$ . It returns `polyderiv( $c$ ,  $-i$ )` when  $i < 0$ . It aborts with error if  $i$  is missing (think of missing as positive infinity).

`polyroots( $c$ )` returns a vector of missing values if any element of  $c$  equals missing.

## Also see

[\[M-4\] mathematical](#) — Important mathematical functions



Syntax

```
void          printf(string scalar fmt, r1, r2, ..., rN)

string scalar sprintf(string scalar fmt, r1, r2, ..., rN)
```

where *fmt* may contain a mix of text and *%fmts*, such as

```
printf("The result is %9.2f, adjusted\n", result)
printf("%s = %9.0g\n", name, value)
```

There must be a one-to-one correspondence between the *%fmts* in *fmt* and the number of results to be displayed.

Along with the usual *%fmts* that Stata provides (see [\[D\] format](#)), also provided are

Format	Meaning
%f	%11.0g, compressed
%g	%11.0g, compressed
%e	%11.8e, compressed
%s	%#s, # = whatever necessary

Compressed means that, after the indicated format is applied, all leading and trailing blanks are removed.

C programmers, be warned: %d is Stata’s (old) calendar date format (equivalent to modern Stata’s %td format) and not an integer format; use %f for formatting integers.

The following character sequences are given a special meaning when contained within *fmt*:

Character sequence	Meaning
%%	one %
\n	newline
\r	carriage return
\t	tab
\\	one \

Description

printf() displays output at the terminal.

printf() returns a string that can then be displayed at the terminal, written to a file, or used in any other way a string might be used.

## Remarks

Remarks are presented under the following headings:

*printf()*  
*sprintf()*

### printf()

printf() displays output at the terminal. A program might contain the line

```
printf("the result is %f\n", result)
```

and display the output

```
the result is 5.213
```

or it might contain the lines

```
printf("{txt}{space 13}{c |}      Coef.      Std. Err.\n")
printf("{hline 13}{c +}{hline 24}\n")
printf("{txt}%12s {c |} {res}%10.0g  %10.0g\n",
       varname[i], coef[i], se[i])
```

and so display the output

	Coef.	Std. Err.
mpg	-.0059541	.0005921

Do not forget to include \n at the end of lines. When \n is not included, the line continues. For instance, the code

```
printf("{txt}{space 13}{c |}      Coef.      Std. Err.\n")
printf("{hline 13}{c +}{hline 24}\n")
printf("{txt}%12s {c |} {res}", varname[i])
printf("%10.0g", coef[i])
printf(" ")
printf("%10.0g", se[i])
printf("\n")
```

produces the same output as shown above.

Although users are unaware of it, Stata buffers output. This makes Stata faster. A side effect of the buffering, however, is that output may not appear when you want it to appear. Consider the code fragment

```
for (n=1; !converged(b, b0); n++) {
    printf("iteration %f:  diff = %12.0g\n", n, b-b0)
    b0 = b
    ... new calculation of b ...
}
```

One of the purposes of the iteration output is to keep the user informed that the code is indeed working, yet as the above code is written, the user probably will not see the iteration messages as they occur. Instead, nothing will appear for a while, and then, unexpectedly, many iteration messages will appear as Stata, buffers full, decides to send to the terminal the waiting output.

To force output to be displayed, use [M-5] **displayflush()**:

```
for (n=1; !converged(b, b0); n++) {
    printf("iteration %f: diff = %12.0g\n", n, b-b0)
    displayflush()
    b0 = b
    ... new calculation of b ...
}
```

It is only in situations like the above that use of `displayflush()` is necessary. In other cases, it is better to let Stata decide when output buffers should be flushed. (Ado-file programmers: you have never had to worry about this because, at the ado-level, all output is flushed as it is created. Mata, however, is designed to be fast and so `printf()` does not force output to be flushed until it is efficient to do so.)

## sprintf()

The difference between `sprintf()` and `printf()` is that, whereas `printf()` sends the resulting string to the terminal, `sprintf()` returns it. Since Mata displays the results of expressions that are not assigned to variables, `sprintf()` used by itself also displays output:

```
: sprintf("the result is %f\n", result)
the result is 5.2130a
```

The outcome is a little different from that produced by `printf()` because the output-the-unassigned-expression routine indents results by 2 and displays all the characters in the string (the `0a` at the end is the `\n` newline character). Also, the output-the-unassigned-expression routine does not honor SMCL, electing instead to display the codes:

```
: sprintf("{txt}the result is {res}%f", result)
{txt}the result is {res}5.213
```

The purpose of `sprintf()` is to create strings that will then be used with `printf()`, with [M-5] **display()**, with `fput()` (see [M-5] **fopen()**), or with some other function.

Pretend that we are creating a dynamically formatted table. One of the columns in the table contains integers, and we want to create a `%fmt` that is exactly the width required. That is, if the integers to appear in the table are 2, 9, and 20, we want to create a `%2.0f` format for the column. We assume the integers are in the column vector `dof` in what follows:

```
max = 0
for (i=1; i<=rows(dof); i++) {
    len = strlen(sprintf("%f", dof[i]))
    if (len>max) max = len
}
fmt = sprintf("%%%f.0f", max)
```

We used `sprintf()` twice in the above. We first used `sprintf()` to produce the string representation of the integer `dof[i]`, and we used the `%f` format so that the length would be whatever was necessary, and no more. We obtained in `max` the maximum length. If `dof` contained 2, 9, and 20, by the end of our loop, `max` will contain 2. We finally used `sprintf()` to create the `%.0f` format that we wanted: `%2.0f`.

The format string `%%f.0f` in the final `sprintf()` is a little difficult to read. The first two percent signs amount to one real percent sign, so in the output we now have `%` and we are left with `%f.0f`. The `%f` is a format—it is how we are to format `max`—and so in the output we now have `%2`, and we are left with `.0f`. `.0f` is just a string, so the final output is `%2.0f`.

## Conformability

```
printf(fmt, r1, r2, ..., rN)
```

```
  fmt:      1 × 1
    r1:      1 × 1
    r2:      1 × 1
    ...
    rN:      1 × 1
  result:    void
```

```
sprintf(fmt, r1, r2, ..., rN)
```

```
  fmt:      1 × 1
    r1:      1 × 1
    r2:      1 × 1
    ...
    rN:      1 × 1
  result:    1 × 1
```

## Diagnostics

`printf()` and `sprintf()` abort with error if a `%fmt` is misspecified, if a numeric `%fmt` corresponds to a string result or a string `%fmt` to a numeric result, or there are too few or too many `%fmts` in `fmt` relative to the number of `results` specified.

## Also see

[M-5] [displayas\(\)](#) — Set display level

[M-5] [displayflush\(\)](#) — Flush terminal-output buffer

[M-4] [io](#) — I/O functions

## Syntax

<i>void</i>	<code>qrd(</code> <i>numeric matrix A, Q, R</i> <code>)</code>
<i>void</i>	<code>hqrd(</code> <i>numeric matrix A, H, tau, R<sub>1</sub></i> <code>)</code>
<i>void</i>	<code>_hqrd(</code> <i>numeric matrix A, tau, R<sub>1</sub></i> <code>)</code>
<i>numeric matrix</i>	<code>hqrdmultq(</code> <i>numeric matrix H, rowvector tau,</i> <i>numeric matrix X, real scalar transpose</i> <code>)</code>
<i>numeric matrix</i>	<code>hqrdmultq1t(</code> <i>numeric matrix H, rowvector tau,</i> <i>numeric matrix X</i> <code>)</code>
<i>numeric matrix</i>	<code>hqrdq(</code> <i>numeric matrix H, numeric matrix tau</i> <code>)</code>
<i>numeric matrix</i>	<code>hqrdq1(</code> <i>numeric matrix H, numeric matrix tau</i> <code>)</code>
<i>numeric matrix</i>	<code>hqrdr(</code> <i>numeric matrix H</i> <code>)</code>
<i>numeric matrix</i>	<code>hqrdr1(</code> <i>numeric matrix H</i> <code>)</code>
<i>void</i>	<code>qrdp(</code> <i>numeric matrix A, Q, R, real rowvector p</i> <code>)</code>
<i>void</i>	<code>hqrdp(</code> <i>numeric matrix A, H, tau, R<sub>1</sub>, real rowvector p</i> <code>)</code>
<i>void</i>	<code>_hqrdp(</code> <i>numeric matrix A, tau, R<sub>1</sub>, real rowvector p</i> <code>)</code>
<i>void</i>	<code>_hqrdp_la(</code> <i>numeric matrix A, tau, real rowvector p</i> <code>)</code>

## Description

`qrd(A, Q, R)` calculates the QR decomposition of  $A$ :  $m \times n$ ,  $m \geq n$ , returning results in  $Q$  and  $R$ .

`hqrd(A, H, tau, R1)` calculates the QR decomposition of  $A$ :  $m \times n$ ,  $m \geq n$ , but rather than returning  $Q$  and  $R$ , returns the Householder vectors in  $H$  and the scale factors  $tau$ —from which  $Q$  can be formed—and returns an upper-triangular matrix in  $R_1$  that is a submatrix of  $R$ ; see [Remarks](#) below for its definition. Doing this saves calculation and memory, and other routines allow you to manipulate these matrices:

1. `hqrdmultq(H, tau, X, transpose)` returns  $QX$  or  $Q'X$  on the basis of the  $Q$  implied by  $H$  and  $tau$ .  $QX$  is returned if  $transpose = 0$ , and  $Q'X$  is returned otherwise.
2. `hqrdmultq1t(H, tau, X)` returns  $Q'_1X$  on the basis of the  $Q_1$  implied by  $H$  and  $tau$ .
3. `hqrdq(H, tau)` returns the  $Q$  matrix implied by  $H$  and  $tau$ . This function is rarely used.
4. `hqrdq1(H, tau)` returns the  $Q_1$  matrix implied by  $H$  and  $tau$ . This function is rarely used.

5. `hqrdr(H)` returns the full  $R$  matrix. This function is rarely used. (It may surprise you that `hqrdr()` is a function of  $H$  and not  $R_1$ .  $R_1$  also happens to be stored in  $H$ , and there is other useful information there, as well.)
6. `hqrdr1(H)` returns the  $R_1$  matrix. This function is rarely used.

`_hqrd(A, tau, R1)` does the same thing as `hqrd(A, H, tau, R1)`, except that it overwrites  $H$  into  $A$  and so conserves even more memory.

`qrdp(A, Q, R, p)` is similar to `qrd(A, Q, R)`: it returns the QR decomposition of  $A$  in  $Q$  and  $R$ . The difference is that this routine allows for pivoting. New argument  $p$  specifies whether a column is available for pivoting and, on output,  $p$  is overwritten with a permutation vector that records the pivoting actually performed. On input,  $p$  can be specified as `.` (missing)—meaning all columns are available for pivoting—or  $p$  can be specified as an  $n \times 1$  column vector containing 0s and 1s, with 1 meaning the column is fixed and so may not be pivoted.

`hqrdp(A, H, tau, R1, p)` is a generalization of `hqrd(A, H, tau, R1)` just as `qrdp()` is a generalization of `qrd()`.

`_hqrdp(A, tau, R1, p)` does the same thing as `hqrdp(A, H, tau, R1, p)`, except that `_hqrdp()` overwrites  $H$  into  $A$ .

`_hqrdp_la()` is the interface into the [M-1] LAPACK routine that performs the QR calculation; it is used by all the above routines. Direct use of `_hqrdp_la()` is not recommended.

## Remarks

Remarks are presented under the following headings:

- [QR decomposition](#)
- [Avoiding calculation of Q](#)
- [Pivoting](#)
- [Least-squares solutions with dropped columns](#)

## QR decomposition

The decomposition of square or nonsquare matrix  $A$  can be written

$$A = QR$$

(1)

where  $Q$  is an orthogonal matrix ( $Q'Q = I$ ), and  $R$  is upper triangular. `qrd(A, Q, R)` will make this calculation:

```
: A
      1      2
1      7      4
2      9      6
3      9      6
4      7      2
5      3      1

: Q = R = .
: qrd(A, Q, R)
: Ahat = Q*R
: mreldif(Ahat, A)
3.55271e-16
```

## Avoiding calculation of $Q$

In fact, you probably do not want to use `qrd()`. Calculating the necessary ingredients for  $Q$  is not too difficult, but going from those necessary ingredients to form  $Q$  is devilish. The necessary ingredients are usually all you need, which are the Householder vectors and their scale factors, known as  $H$  and  $\tau$ . For instance, one can write down a mathematical function  $f(H, \tau, X)$  that will calculate  $QX$  or  $Q'X$  for some matrix  $X$ .

Also, QR decomposition is often carried out on violently nonsquare matrices  $A: m \times n, m \gg n$ . We can write

$$A_{m \times n} = \begin{bmatrix} Q_1 & Q_2 \\ m \times n & m \times m-n \end{bmatrix} \begin{bmatrix} R_1 \\ R_2 \\ m-n \times n \end{bmatrix} = \begin{matrix} Q_1 R_1 \\ m \times n & m \times n \end{matrix} + \begin{matrix} Q_2 R_2 \\ m \times n & m \times n \end{matrix}$$

$R_2$  is zero, and thus

$$A_{m \times n} = \begin{bmatrix} Q_1 & Q_2 \\ m \times n & m \times m-n \end{bmatrix} \begin{bmatrix} R_1 \\ n \times n \\ 0 \\ m-n \times n \end{bmatrix} = \begin{matrix} Q_1 R_1 \\ m \times n \end{matrix}$$

Thus it is enough to know  $Q_1$  and  $R_1$ . Rather than defining QR decomposition as

$$A = QR, \quad Q: m \times m, \quad R: m \times n \quad (1)$$

it is better to define it as

$$A = Q_1 R_1 \quad Q_1: m \times n \quad R_1: n \times n \quad (1')$$

To appreciate the savings, consider the reasonable case where  $m = 4,000$  and  $n = 3$ :

$$A = QR, \quad Q: 4,000 \times 4,000, \quad R: 4,000 \times 3$$

versus,

$$A = Q_1 R_1 \quad Q_1: 4,000 \times 3 \quad R_1: 3 \times 3$$

Memory consumption is reduced from 125,094 kilobytes to 94 kilobytes, a 99.92% saving!

Combining the arguments, we need not save  $Q$  because  $Q_1$  is sufficient, we need not calculate  $Q_1$  because  $H$  and  $\tau$  are sufficient, and we need not store  $R$  because  $R_1$  is sufficient.

That is what `hqrd(A, H, tau, R1)` does. Having used `hqrd()`, if you need to multiply the full  $Q$  by some matrix  $X$ , you can use `hqrdmultq()`. Having used `hqrd()`, if you need the full  $Q$ , you can use `hqrdq()` to obtain it, but by that point you will be making the devilish calculation you sought to avoid and so you might as well have used `qrd()` to begin with. If you want  $Q_1$ , you can use `hqrdq1()`. Finally, having used `hqrd()`, if you need  $R$  or  $R_1$ , you can use `hqrdr()` and `hqrdr1()`:

```

: A
      1  2
      ┌───┐
1     │ 7  4 │
2     │ 9  6 │
3     │ 9  6 │
4     │ 7  2 │
5     │ 3  1 │
      └───┘

: H = tau = R1 = .
: hqrd(A, H, tau, R1)
: Ahat = hqrdq1(H, tau) * R1           // i.e., Q1*R1
: mreldif(Ahat, A)
3.55271e-16

```

Pivoting

The QR decomposition with column pivoting solves

$$AP = QR \tag{2}$$

or, if you prefer,

$$AP = Q_1R_1 \tag{2'}$$

where  $P$  is a permutation matrix; see [M-1] [permutation](#). We can rewrite this as

$$A = QRP' \tag{3}$$

and

$$A = Q_1R_1P' \tag{3'}$$

Column pivoting can improve the numerical accuracy. The functions `qrdp(A, Q, R, p)` and `hqrdp(A, H, tau, R1, p)` perform pivoting and return the permutation matrix  $P$  in permutation vector form:

```

: A
      1  2
      ┌───┐
1     │ 7  4 │
2     │ 9  6 │
3     │ 9  6 │
4     │ 7  2 │
5     │ 3  1 │
      └───┘

: Q = R = p = .
: qrdp(A, Q, R, p)
: Ahat = (Q*R)[., invorder(p)]         // i.e., QRP'
: mreldif(Ahat, A)
3.55271e-16

: H = tau = R1 = p = .
: hqrdp(A, H, tau, R1, p)
: Ahat = (hqrdq1(H, tau)*R1)[., invorder(p)] // i.e., Q1*R1*P'
: mreldif(Ahat, A)
3.55271e-16

```



Before calling `qrdp()` or `hqrdp()`, we set  $p$  equal to `missing`, specifying that all columns could be pivoted. We could just as well have set  $p$  equal to  $(0, 0)$ , which would have stated that both columns were eligible for pivoting.

When pivoting is disallowed, and when  $A$  is not of full column rank, the order in which columns appear affects the kind of generalized solution produced; later columns are, in effect, dropped. When pivoting is allowed, the columns are reordered based on numerical accuracy considerations. In the rank-deficient case, you no longer know ahead of time which columns will be dropped, because you do not know in what order the columns will appear. Generally, you do not care, but there are occasions when you do.

In such cases, you can specify which columns are eligible for pivoting and which are not—you specify  $p$  as a vector and if  $p_i == 1$ , the  $i$ th column may not be pivoted. The  $p_i == 1$  columns are (conceptually) moved to appear first in the matrix, and the remaining columns are ordered optimally after that. The permutation vector that is returned in  $p$  accounts for all of this.

## Least-squares solutions with dropped columns

Least-square solutions are one popular use of QR decomposition. We wish to solve for  $x$

$$Ax = b \quad (A : m \times n, \quad m \geq n) \quad (4)$$

The problem is that there is no solution to (4) when  $m > n$  because we have more equations than unknowns. Then we want to find  $x$  such that  $(Ax - b)'(Ax - b)$  is minimized.

If  $A$  is of full column rank then it is well known that the least-squares solution for  $x$  is given by `solveupper( $R_1$ ,  $Q_1'b$ )` where `solveupper()` is an upper-triangular solver; see [M-5] `solve_lower()`.

If  $A$  is of less than full column rank and we do not care which columns are dropped, then we can use the same solution: `solveupper( $R_1$ ,  $Q_1'b$ )`.

Adding pivoting to the above hardly complicates the issue; the solution becomes `solveupper( $R_1$ ,  $Q_1'b$ ) [invorder( $p$ )]`.

For both cases, the full details are

```
: A
      1   2   3
1  

|   |   |   |
|---|---|---|
| 3 | 9 | 1 |
| 3 | 8 | 1 |
| 3 | 7 | 1 |
| 3 | 6 | 1 |


2
3
4

: b
      1
1  

|    |
|----|
| 7  |
| 3  |
| 12 |
| 0  |


2
3
4

: H = tau = R1 = p = .
: hqrdp(A, H, tau, R1, p)
: q1b = hqrdmultq1t(H, tau, b)           // i.e., Q1'b
: xhat = solveupper(R1, q1b)[invorder(p)]
```

```
: xhat
      1
1  -1.166666667
2      1.2
3      0
```

The  $A$  matrix in the above example has less than full column rank; the first column contains a variable with no variation and the third column contains the data for the intercept. The solution above is correct, but we might prefer a solution that included the intercept. To do that, we need to specify that the third column cannot be pivoted:

```
: p = (0, 0, 1)
: H = tau = R1 = .
: hqrdp(A, H, tau, R1, p)
: q1b = hqrdmultq1t(H, tau, b)
: xhat = solveupper(R1, q1b)[invorder(p)]
: xhat
      1
1      0
2      1.2
3     -3.5
```

## Conformability

`qrd(A, Q, R):`

*input:*

$A:$        $m \times n, \quad m \geq n$

*output:*

$Q:$        $m \times m$

$R:$        $m \times n$

`hqrd(A, H, tau, R1):`

*input:*

$A:$        $m \times n, \quad m \geq n$

*output:*

$H:$        $m \times n$

$tau:$       $1 \times n$

$R_1:$       $n \times n$

`_hqrd(A, tau, R1):`

*input:*

$A:$        $m \times n, \quad m \geq n$

*output:*

$A:$        $m \times n$     (contains  $H$ )

$tau:$       $1 \times n$

$R_1:$       $n \times n$

hqrddmultq( $H$ ,  $\tau$ ,  $X$ ,  $\text{transpose}$ ):

$H$ :  $m \times n$   
 $\tau$ :  $1 \times n$   
 $X$ :  $m \times c$   
 $\text{transpose}$ :  $1 \times 1$   
 $\text{result}$ :  $m \times c$

hqrddmultq1t( $H$ ,  $\tau$ ,  $X$ ):

$H$ :  $m \times n$   
 $\tau$ :  $1 \times n$   
 $X$ :  $m \times c$   
 $\text{result}$ :  $n \times c$

hqrddq( $H$ ,  $\tau$ ):

$H$ :  $m \times n$   
 $\tau$ :  $1 \times n$   
 $\text{result}$ :  $m \times m$

hqrddq1( $H$ ,  $\tau$ ):

$H$ :  $m \times n$   
 $\tau$ :  $1 \times n$   
 $\text{result}$ :  $m \times n$

hqrddr( $H$ ):

$H$ :  $m \times n$   
 $\text{result}$ :  $m \times n$

hqrddr1( $H$ ):

$H$ :  $m \times n$   
 $\text{result}$ :  $n \times n$

qrdp( $A$ ,  $Q$ ,  $R$ ,  $p$ ):

*input*:

$A$ :  $m \times n$ ,  $m \geq n$   
 $p$ :  $1 \times 1$  or  $1 \times n$

*output*:

$Q$ :  $m \times m$   
 $R$ :  $m \times n$   
 $p$ :  $1 \times n$

hqrddp( $A$ ,  $H$ ,  $\tau$ ,  $R_1$ ,  $p$ ):

*input*:

$A$ :  $m \times n$ ,  $m \geq n$   
 $p$ :  $1 \times 1$  or  $1 \times n$

*output*:

$H$ :  $m \times n$   
 $\tau$ :  $1 \times n$   
 $R_1$ :  $n \times n$   
 $p$ :  $1 \times n$

`_hqrqp(A, tau, R1, p)`:

*input*:

$A$ :  $m \times n$ ,  $m \geq n$   
 $p$ :  $1 \times 1$  or  $1 \times n$

*output*:

$A$ :  $m \times n$  (contains  $H$ )  
 $tau$ :  $1 \times n$   
 $R_1$ :  $n \times n$   
 $p$ :  $1 \times n$

`_hqrqp_la(A, tau, p)`:

*input*:

$A$ :  $m \times n$ ,  $m \geq n$   
 $p$ :  $1 \times 1$  or  $1 \times n$

*output*:

$A$ :  $m \times n$  (contains  $H$ )  
 $tau$ :  $1 \times n$   
 $p$ :  $1 \times n$

## Diagnostics

`qrd(A, ...)`, `hqrqp(A, ...)`, `_hqrqp(A, ...)`, `qrdp(A, ...)`, `hqrqp(A, ...)`, and `_hqrqp(A, ...)` return missing results if  $A$  contains missing values. That is,  $Q$  will contain all missing values.  $R$  will contain missing values on and above the diagonal.  $p$  will contain the integers 1, 2, ....

`_hqrqp(A, ...)` and `_hqrqp(A, ...)` abort with error if  $A$  is a view.

`hqrqp_multq(H, tau, X, transpose)` and `hqrqp_multq1t(H, tau, X)` return missing results if  $X$  contains missing values.

Alston Scott Householder (1904–1993) was born in Rockford, Illinois, and grew up in Alabama. He studied philosophy at Northwestern and Cornell, and then mathematics, earning a doctorate in the calculus of variations from the University of Chicago. Householder worked on mathematical biology for several years at Chicago, but in 1946 he moved on to Oak Ridge National Laboratory where he became the founding director of the Mathematics Division in 1948. There he moved into numerical analysis, specializing in linear equations and eigensystems and helping to unify the field through reviews and symposia. His last post was at the University of Tennessee.

## Also see

[M-5] `qrsolve()` — Solve  $AX=B$  for  $X$  using QR decomposition

[M-5] `qrinva()` — Generalized inverse of matrix via QR decomposition

[M-4] `matrix` — Matrix functions

## Syntax

```

numeric matrix    qrinv(numeric matrix A)
numeric matrix    qrinv(numeric matrix A, rank)
numeric matrix    qrinv(numeric matrix A, rank, real scalar tol)

real scalar       _qrinv(numeric matrix A)
real scalar       _qrinv(numeric matrix A, real scalar tol)
    
```

where the type of *rank* is irrelevant; the rank of *A* is returned there.

## Description

`qrinv(A, ...)` returns the inverse or generalized inverse of real or complex matrix *A*:  $m \times n$ ,  $m \geq n$ . If optional argument *rank* is specified, the rank of *A* is returned there.

`_qrinv(A, ...)` does the same thing except that, rather than returning the result, it overwrites the original matrix *A* with the result. `_qrinv()` returns the rank of *A*.

In both cases, optional argument *tol* specifies the tolerance for determining singularity; see *Remarks* below.

## Remarks

`qrinv()` and `_qrinv()` are most often used on square and possibly rank-deficient matrices but may be used on nonsquare matrices that have more rows than columns. Also see [\[M-5\] pinv\(\)](#) for an alternative. See [\[M-5\] luinv\(\)](#) for a more efficient way to obtain the inverse of full-rank, square matrices, and see [\[M-5\] invsym\(\)](#) for inversion of real, symmetric matrices.

When *A* is of full rank, the inverse calculated by `qrinv()` is essentially the same as that computed by the faster `luinv()`. When *A* is singular, `qrinv()` and `_qrinv()` compute a generalized inverse,  $A^*$ , which satisfies

$$\begin{aligned}
 A(A^*)A &= A \\
 (A^*)A(A^*) &= A^*
 \end{aligned}$$

This generalized inverse is also calculated for nonsquare matrices that have more rows than columns and, then returned is a least-squares solution. If *A* is  $m \times n$ ,  $m \geq n$ , and if the rank of *A* is equal to *n*, then  $(A^*)A = I$ , ignoring roundoff error.

`qrinv(A)` is implemented as `qrsolve(A, I(rows(A)))`; see [\[M-5\] qrsolve\(\)](#) for details and for use of the optional *tol* argument.

## Conformability

`qrinv(A, rank, tol):`

*input:*

*A:*         $m \times n$ ,     $m \geq n$   
*tol:*       $1 \times 1$     (optional)

*output:*

*rank:*      $1 \times 1$     (optional)  
*result:*     $n \times m$

`_qrinv(A, tol):`

*input:*

*A:*         $m \times n$ ,     $m \geq n$   
*tol:*       $1 \times 1$     (optional)

*output:*

*A:*         $n \times m$   
*result:*     $1 \times 1$     (containing rank)

## Diagnostics

The inverse returned by these functions is real if  $A$  is real and is complex if  $A$  is complex.

`qrinv(A, ...)` and `_qrinv(A, ...)` return a result containing missing values if  $A$  contains missing values.

`_qrinv(A, ...)` aborts with error if  $A$  is a view.

See [M-5] `qrsolve()` and [M-1] **tolerance** for information on the optional *tol* argument.

## Also see

[M-5] `invsym()` — Symmetric real matrix inversion

[M-5] `cholinv()` — Symmetric, positive-definite matrix inversion

[M-5] `luinv()` — Square matrix inversion

[M-5] `pinv()` — Moore–Penrose pseudoinverse

[M-5] `qrsolve()` — Solve  $AX=B$  for  $X$  using QR decomposition

[M-5] `solve_tol()` — Tolerance used by solvers and inverters

[M-4] **matrix** — Matrix functions

[M-4] **solvers** — Functions to solve  $AX=B$  and to obtain  $A$  inverse

# Title

[M-5] **qrsolve()** — Solve  $AX=B$  for  $X$  using QR decomposition

## Syntax

<i>numeric matrix</i>	<code>qrsolve(A, B)</code>
<i>numeric matrix</i>	<code>qrsolve(A, B, rank)</code>
<i>numeric matrix</i>	<code>qrsolve(A, B, rank, tol)</code>
<i>real scalar</i>	<code>_qrsolve(A, B)</code>
<i>real scalar</i>	<code>_qrsolve(A, B, tol)</code>

where

<i>A:</i>	<i>numeric matrix</i>
<i>B:</i>	<i>numeric matrix</i>
<i>rank:</i>	irrelevant; <i>real scalar</i> returned
<i>tol:</i>	<i>real scalar</i>

## Description

`qrsolve(A, B, ...)` uses QR decomposition to solve  $AX = B$  and returns  $X$ . When  $A$  is singular or nonsquare, `qrsolve()` computes a least-squares generalized solution. When *rank* is specified, in it is placed the rank of  $A$ .

`_qrsolve(A, B, ...)`, does the same thing, except that it destroys the contents of  $A$  and it overwrites  $B$  with the solution. Returned is the rank of  $A$ .

In both cases, *tol* specifies the tolerance for determining whether  $A$  is of full rank. *tol* is interpreted in the standard way—as a multiplier for the default if  $tol > 0$  is specified and as an absolute quantity to use in place of the default if  $tol \leq 0$  is specified; see [\[M-1\] tolerance](#).

## Remarks

`qrsolve(A, B, ...)` is suitable for use with square and possibly rank-deficient matrix  $A$ , or when  $A$  has more rows than columns. When  $A$  is square and full rank, `qrsolve()` returns the same solution as `lusolve()` (see [\[M-5\] lusolve\(\)](#)), up to roundoff error. When  $A$  is singular, `qrsolve()` returns a generalized (least-squares) solution.

Remarks are presented under the following headings:

- [Derivation](#)
- [Relationship to inversion](#)
- [Tolerance](#)

## Derivation

We wish to solve for  $X$

$$AX = B \tag{1}$$

Perform QR decomposition on  $A$  so that we have  $A = QRP'$ . Then (1) can be rewritten

$$QRP'X = B$$

Premultiplying by  $Q'$  and remembering that  $Q'Q = QQ' = I$ , we have

$$RP'X = Q'B \tag{2}$$

Define

$$Z = P'X \tag{3}$$

Then (2) can be rewritten

$$RZ = Q'B \tag{4}$$

It is easy to solve (4) for  $Z$  because  $R$  is upper triangular. Having  $Z$ , we can obtain  $X$  via (3), because  $Z = P'X$ , premultiplied by  $P$  (and if we remember that  $PP' = I$ ), yields

$$X = PZ$$

For more information on QR decomposition, see [M-5] `qrd()`.

## Relationship to inversion

For a general discussion, see *Relationship to inversion* in [M-5] `lusolve()`.

For an inverse based on QR decomposition, see [M-5] `qrinv()`. `qrinv(A)` amounts to `qrsolve(A, I(rows(A)))`, although it is not actually implemented that way.

## Tolerance

The default tolerance used is

$$eta = 1e-13 * trace(abs(R))/rows(R)$$

where  $R$  is the upper-triangular matrix of the QR decomposition; see *Derivation* above. When  $A$  is less than full rank, by, say,  $d$  degrees of freedom, then  $R$  is also rank deficient by  $d$  degrees of freedom and the bottom  $d$  rows of  $R$  are essentially zero. If the  $i$ th diagonal element of  $R$  is less than or equal to  $eta$ , then the  $i$ th row of  $Z$  is set to zero. Thus if the matrix is singular, `qrsolve()` provides a generalized solution.

If you specify  $tol > 0$ , the value you specify is used to multiply  $eta$ . You may instead specify  $tol \leq 0$ , and then the negative of the value you specify is used in place of  $eta$ ; see [M-1] *tolerance*.



## Conformability

`qrsolve(A, B, rank, tol):`

*input:*

$A:$       $m \times n, \quad m \geq n$   
 $B:$       $m \times k$   
 $tol:$      $1 \times 1$     (optional)

*output:*

$rank:$      $1 \times 1$     (optional)  
 $result:$     $n \times k$

`_qrsolve(A, B, tol):`

*input:*

$A:$       $m \times n, \quad m \geq n$   
 $B:$       $m \times k$   
 $tol:$      $1 \times 1$     (optional)

*output:*

$A:$       $0 \times 0$   
 $B:$       $n \times k$   
 $result:$     $1 \times 1$

## Diagnostics

`qrsolve(A, B, ...)` and `_qrsolve(A, B, ...)` return a result containing missing if  $A$  or  $B$  contain missing values.

`_qrsolve(A, B, ...)` aborts with error if  $A$  or  $B$  are views.

## Also see

[M-5] `qrinv()` — Generalized inverse of matrix via QR decomposition

[M-5] `qrd()` — QR decomposition

[M-5] `solvelower()` — Solve  $AX=B$  for  $X$ ,  $A$  triangular

[M-5] `cholsolve()` — Solve  $AX=B$  for  $X$  using Cholesky decomposition

[M-5] `lusolve()` — Solve  $AX=B$  for  $X$  using LU decomposition

[M-5] `svsolve()` — Solve  $AX=B$  for  $X$  using singular value decomposition

[M-5] `solve_tol()` — Tolerance used by solvers and inverters

[M-4] `matrix` — Matrix functions

[M-4] `solvers` — Functions to solve  $AX=B$  and to obtain  $A$  inverse

## Syntax

*real matrix*    `quadcross(X, Z)`

*real matrix*    `quadcross(X, w, Z)`

*real matrix*    `quadcross(X, xc, Z, zc)`

*real matrix*    `quadcross(X, xc, w, Z, zc)`

*real matrix*    `quadcrossdev(X, x, Z, z)`

*real matrix*    `quadcrossdev(X, x, w, Z, z)`

*real matrix*    `quadcrossdev(X, xc, x, Z, zc, z)`

*real matrix*    `quadcrossdev(X, xc, x, w, Z, zc, z)`

where

*X*:        *real matrix X*

*xc*:       *real scalar xc*

*x*:        *real rowvector x*

*w*:        *real vector w*

*Z*:        *real matrix Z*

*zc*:       *real scalar zc*

*z*:        *real rowvector z*

## Description

`quadcross()` makes calculations of the form

$X'X$

$X'Z$

$X'\text{diag}(w)X$

$X'\text{diag}(w)Z$

This function mirrors `cross()` (see [\[M-5\] `cross\(\)`](#)), the difference being that sums are formed in quad precision rather than in double precision, so `quadcross()` is more accurate.

`quadcrossdev()` makes calculations of the form

$$(X: -x)'(X: -x)$$

$$(X: -x)'(Z: -z)$$

$$(X: -x)'\text{diag}(w)(X: -x)$$

$$(X: -x)'\text{diag}(w)(Z: -z)$$

This function mirrors `crossdev()` (see [M-5] `crossdev()`), the difference being that sums are formed in quad precision rather than in double precision, so `quadcrossdev()` is more accurate.

## Remarks

The returned result is double precision, but the sum calculations made in creating that double-precision result were made in quad precision.

## Conformability

`quadcross()` has the same conformability requirements as `cross()`; see [M-5] `cross()`.

`quadcrossdev()` has the same conformability requirements as `crossdev()`; see [M-5] `crossdev()`.

## Diagnostics

See *Diagnostics* in [M-5] `cross()` and *Diagnostics* in [M-5] `crossdev()`.

## Also see

[M-5] `cross()` — Cross products

[M-5] `crossdev()` — Deviation cross products

[M-4] `statistical` — Statistical functions

[M-4] `utility` — Matrix utility functions

# Title

[M-5] **range()** — Vector over specified range

## Syntax

*numeric colvector*    **range**(*a*, *b*, *numeric scalar delta*)

*numeric colvector*    **rangen**(*a*, *b*, *real scalar n*)

where *a* and *b* are numeric scalars.

## Description

**range**(*a*, *b*, *delta*) returns a column vector going from *a* to *b* in steps of  $\text{abs}(\text{delta})$  ( $b \geq a$ ) or  $-\text{abs}(\text{delta})$  ( $b < a$ ).

**rangen**(*a*, *b*, *n*) returns a  $\text{round}(n) \times 1$  column vector going from *a* to *b* in  $\text{round}(n)-1$  steps. *a* may be less than, equal to, or greater than *b*.

## Remarks

**range**(0, 1, .25) returns (0 \ .25 \ .5 \ .75 \ 1). The sign of the third argument does not matter; **range**(0, 1, -.25) returns the same thing. **range**(1, 0, .25) and **range**(1, 0, -.25) return (1 \ .75 \ .5 \ .25 \ 0).

**rangen**(0, .5, 6) returns (0 \ .1 \ .2 \ .3 \ .4 \ .5). **rangen**(.5, 0, 6) returns (.5 \ .4 \ .3 \ .2 \ .1 \ 0).

**range**() and **rangen**() may be used with complex arguments. **range**(1, 1i, .4) returns (1 \ .75+.25i \ .5+.5i \ .25+.75i \ 1i). **rangen**(1, 1i, 5) returns the same thing. For **range**(), only the distance of *delta* from zero matters, so **range**(1, 1i, .4i) would produce the same result, as would **range**(1, 1i, .25+.312i).

## Conformability

**range**(*a*, *b*, *delta*):

<i>a</i> :	$1 \times 1$
<i>b</i> :	$1 \times 1$
<i>delta</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$ , if $a = b$
	$\max(1+\text{abs}(b-a)/\text{abs}(\text{delta}), 2) \times 1$ , otherwise

**rangen**(*a*, *b*, *n*):

<i>a</i> :	$1 \times 1$
<i>b</i> :	$1 \times 1$
<i>n</i> :	$n \times 1$

## Diagnostics

`range(a, b, delta)` aborts with error if  $a$ ,  $b$ , or  $delta$  contains missing, if  $\text{abs}(b-a)/\text{abs}(delta)$  results in overflow, or if  $1+\text{abs}(b-a)/\text{abs}(delta)$  results in a vector that is too big given the amount of memory available.

`range(a, b, delta)` returns a  $1 \times 1$  result when  $a = b$ . In all other cases, the result is  $2 \times 1$  or longer.

`rangen(a, b, n)` aborts with error if  $\text{round}(n)$  is less than 0 or missing.

## Also see

[\[M-4\] `standard`](#) — Functions to create standard matrices

# Title

[M-5] **rank()** — Rank of matrix

## Syntax

*real scalar*    **rank**(*numeric matrix A*)

*real scalar*    **rank**(*numeric matrix A*, *real scalar tol*)

## Description

**rank**(*A*) and **rank**(*A*, *tol*) return the rank of *A*:  $m \times n$ .

## Remarks

The row rank of a matrix *A*:  $m \times n$  is the number of rows of *A* that are linearly independent. The column rank is the number of columns that are linearly independent. The terms row rank and column rank, however, are used merely for emphasis. The ranks are equal, and the result is simply called the rank of *A*.

**rank**() calculates the rank by counting the number of nonzero singular values of the SVD of *A*, where nonzero is interpreted relative to a tolerance. **rank**() uses the same tolerance as **pinv**() (see [M-5] **pinv**()) and as **svsolve**() (see [M-5] **svsolve**()), and optional argument *tol* is specified in the same way as with those functions.

Thus if you were going to use **rank**() before calculating an inverse using **pinv**(), it would be better to skip **rank**() altogether and proceed to the **pinv**() step, because **pinv**() will return the rank, calculated as a by-product of calculating the inverse. Using **rank**() ahead of time, the SVD would be calculated twice.

**rank**() in general duplicates calculations; and, worse, if you are not planning on using **pinv**() or **svsolve**() but rather are planning on using some other function, the rank returned by **rank**() may disagree with the implied rank of whatever numerical method you subsequently use because each numerical method has its own precision and tolerances.

All that said, **rank**() is useful in interactive and pedagogical situations.

## Conformability

**rank**(*A*, *tol*):

<i>A</i> :	$m \times n$	
<i>tol</i> :	$1 \times 1$	(optional)
<i>result</i> :	$1 \times 1$	

## Diagnostics

**rank**(*A*) returns missing if *A* contains missing values.

## Also see

[M-5] [svd\(\)](#) — Singular value decomposition

[M-5] [fullsvd\(\)](#) — Full singular value decomposition

[M-5] [pinv\(\)](#) — Moore–Penrose pseudoinverse

[M-4] [matrix](#) — Matrix functions

# Title

[M-5] **Re()** — Extract real or imaginary part

## Syntax

*real matrix*    **Re**(*numeric matrix Z*)

*real matrix*    **Im**(*numeric matrix Z*)

## Description

**Re**(*Z*) returns a real matrix containing the real part of *Z*. *Z* may be real or complex.

**Im**(*Z*) returns a real matrix containing the imaginary part of *Z*. *Z* may be a real or complex. If *Z* is real, **Im**(*Z*) returns a matrix of zeros.

## Conformability

**Re**(*Z*), **Im**(*Z*):  
          *Z*:         $r \times c$   
          *result*:     $r \times c$

## Diagnostics

**Re**(*Z*), if *Z* is real, literally returns *Z* and not a copy of *Z*. This makes execution of **Re**() applied to real arguments instant.

## Also see

- [M-5] **C()** — Make complex
- [M-4] **scalar** — Scalar mathematical functions
- [M-4] **utility** — Matrix utility functions



## Syntax

*real matrix*    **reldif**(*numeric matrix X, numeric matrix Y*)

*real scalar*    **mrldif**(*numeric matrix X, numeric matrix Y*)

*real scalar*    **mrldifsym**(*numeric matrix X*)

*real scalar*    **mrldifre**(*numeric matrix X*)

## Description

**reldif**(*X, Y*) returns the relative difference defined by

$$r = \frac{|X - Y|}{|Y| + 1}$$

calculated element by element.

**mrldif**(*X, Y*) returns the maximum relative difference and is equivalent to **max**(**reldif**(*X, Y*)).

**mrldifsym**(*X*) is equivalent to **mrldif**(*X'*, *X*) and so is a measure of how far the matrix is from being symmetric (Hermitian).

**mrldifre**(*X*) is equivalent to **mrldif**(**Re**(*X*), *X*) and so is a measure of how far the matrix is from being real.

## Conformability

**reldif**(*X, Y*):  
    *X*:      $r \times c$   
    *Y*:      $r \times c$   
    *result*:  $r \times c$

**mrldif**(*X, Y*):  
    *X*:      $r \times c$   
    *Y*:      $r \times c$   
    *result*:  $1 \times 1$

**mrldifsym**(*X*):  
    *X*:      $n \times n$   
    *result*:  $1 \times 1$

**mrldifre**(*X*):  
    *X*:      $r \times c$   
    *result*:  $1 \times 1$

## Diagnostics

The relative difference function treats equal missing values as having a difference of 0 and different missing values as having a difference of missing (.):

```
reldif(., .) == reldif(.a, .a) == ... == reldif(.z, .z) == 0
reldif(., .a) == reldif(., .z) == ... == reldif(.y, .z) == .
```

## Also see

[\[M-4\] `utility`](#) — Matrix utility functions

# Title

[M-5] **rows()** — Number of rows and number of columns

## Syntax

*real scalar*    `rows(transmorphic matrix P)`

*real scalar*    `cols(transmorphic matrix P)`

*real scalar*    `length(transmorphic matrix P)`

## Description

`rows(P)` returns the number of rows of  $P$ .

`cols(P)` returns the number of columns of  $P$ .

`length(P)` returns `rows(P)*cols(P)`.

## Remarks

`length(P)` is typically used with vectors, as in

```
for (i=1; i<=length(x); i++) {  
    ... x[i] ...  
}
```

## Conformability

`rows(P)`, `cols(P)`, `length(P)`:

$P$ :	$r \times c$
<i>result</i> :	$1 \times 1$

## Diagnostics

`rows(P)`, `cols(P)`, and `length(P)` return a result that is greater than or equal to zero.

## Also see

[\[M-4\] utility](#) — Matrix utility functions

Syntax

*transmorphic matrix*    rowshape(*transmorphic matrix* *T*, *real scalar* *r*)

*transmorphic matrix*    colshape(*transmorphic matrix* *T*, *real scalar* *c*)

Description

rowshape(*T*, *r*) returns *T* transformed into a matrix with trunc(*r*) rows.

colshape(*T*, *c*) returns *T* having trunc(*c*) columns.

In both cases, elements are assigned sequentially with the column index varying more rapidly. See [M-5] **vec()** for a function that varies the row index more rapidly.

Remarks

Remarks are presented under the following headings:

*Example of rowshape()*  
*Example of colshape()*

Example of rowshape()

: A

	1	2	3	4
1	11	12	13	14
2	21	22	23	24
3	31	32	33	34
4	41	42	43	44

: rowshape(A,2)

	1	2	3	4	5	6	7	8
1	11	12	13	14	21	22	23	24
2	31	32	33	34	41	42	43	44

Example of colshape()

: colshape(A, 2)

	1	2
1	11	12
2	13	14
3	21	22
4	23	24
5	31	32
6	33	34
7	41	42
8	43	44

## Conformability

`rowshape(T, r):`

*T*:  $r_0 \times c_0$   
*r*:  $1 \times 1$   
*result*:  $r \times r_0 c_0 / r$

`colshape(T, c):`

*T*:  $r_0 \times c_0$   
*c*:  $1 \times 1$   
*result*:  $r_0 c_0 / c \times c$

## Diagnostics

Let  $r_0$  and  $c_0$  be the number of rows and columns of  $T$ .

`rowshape()` aborts with error if  $r_0 \times c_0$  is not evenly divisible by `trunc(r)`.

`colshape()` aborts with error if  $r_0 \times c_0$  is not evenly divisible by `trunc(c)`.

## Also see

[\[M-4\] manipulation](#) — Matrix manipulation

## Syntax

```

real matrix    runiform(real scalar r, real scalar c)

string scalar  rseed()
void           rseed(string scalar newseed)
void           rseed(real scalar newseed)

real matrix    rbeta(real scalar r, real scalar c, real matrix a, real matrix b)

real matrix    rbinomial(real scalar r, real scalar c, real matrix n, real matrix p)

real matrix    rchi2(real scalar r, real scalar c, real matrix df)

real matrix    rdiscrete(real scalar r, real scalar c, real colvector p)

real matrix    rgamma(real scalar r, real scalar c, real matrix a, real matrix b)

real matrix    rhypergeometric(real scalar r, real scalar c, real matrix N,
                               real matrix K, real matrix n)

real matrix    rnbinomial(real scalar r, real scalar c, real matrix n, real matrix p)

real matrix    rnormal(real scalar r, real scalar c, real matrix m, real matrix s)

real matrix    rpoisson(real scalar r, real scalar c, real matrix m)

real matrix    rt(real scalar r, real scalar c, real matrix df)

```

## Description

`runiform(r, c)` returns an  $r \times c$  real matrix containing uniformly distributed random variates on  $[0,1)$ . `runiform()` is the same function as Stata's [runiform\(\)](#) function.

`rseed()` returns the current random-variate seed in an encrypted string form. `rseed()` returns the same thing as Stata's `c(seed)`; see [\[R\] set seed](#) and [\[P\] creturn](#).

`rseed(newseed)` sets the seed: a string previously obtained from `rseed()` can be specified for the argument or an integer number can be specified. `rseed()` has the same effect as Stata's `set seed` command; see [\[R\] set seed](#).

`rbeta(r, c, a, b)` returns an  $ir \times jc$  real matrix containing beta random variates. The real-valued matrices *a* and *b* contain the beta shape parameters. The matrices *a* and *b* must be **r-conformable**, where  $i = \max(\text{rows}(a), \text{rows}(b))$  and  $j = \max(\text{cols}(a), \text{cols}(b))$ .

`rbinomial(r, c, n, p)` returns an  $ir \times jc$  real matrix containing binomial random variates. The real-valued matrices *n* and *p* contain the number of trials and the probability parameters. The matrices *n* and *p* must be **r-conformable**, where  $i = \max(\text{rows}(n), \text{rows}(p))$  and  $j = \max(\text{cols}(n), \text{cols}(p))$ .

`rchi2(r, c, df)` returns an  $ir \times jc$  real matrix containing chi-squared random variates. The real-valued matrix *df* contains the degrees of freedom parameters, where  $i = \text{rows}(df)$  and  $j = \text{cols}(df)$ .

`rdiscrete(r, c, p)` returns an  $r \times c$  real matrix containing random variates from the discrete distribution specified by the probabilities in the vector *p* of length *k*. The range of the discrete variates is 1, 2, ..., *k*. The alias method of [Walker \(1977\)](#) is used to sample from the discrete distribution.

`rgamma(r, c, a, b)` returns an  $ir \times jc$  real matrix containing gamma random variates. The real-valued matrices *a* and *b* contain the gamma shape and scale parameters, respectively. The matrices *a* and *b* must be **r-conformable**, where  $i = \max(\text{rows}(a), \text{rows}(b))$  and  $j = \max(\text{cols}(a), \text{cols}(b))$ .

`rhypergeometric(r, c, N, K, n)` returns an  $ir \times jc$  real matrix containing hypergeometric random variates. The integer-valued matrix *N* contains the population sizes, the integer-valued matrix *K* contains the number of elements in each population that have the attribute of interest, and the integer-valued matrix *n* contains the sample size. The matrices *N*, *K*, and *n* must be **r-conformable**, where  $i = \max(\text{rows}(N), \text{rows}(K), \text{rows}(n))$  and  $j = \max(\text{cols}(N), \text{cols}(K), \text{cols}(n))$ .

`rnbinomial(r, c, n, p)` returns an  $ir \times jc$  real matrix containing negative binomial random variates. When the elements of the matrix *n* are integer-valued, `rnbinomial()` returns the number of failures before the *n*th success, where the probability of success on a single draw is contained in the real-valued matrix *p*. The elements of *n* can also be nonintegral but must be positive. The matrices *n* and *p* must be **r-conformable**, where  $i = \max(\text{rows}(n), \text{rows}(p))$  and  $j = \max(\text{cols}(n), \text{cols}(p))$ .

`rnormal(r, c, m, s)` returns an  $ir \times jc$  real matrix containing normal (Gaussian) random variates. The real-valued matrices *m* and *s* contain the mean and standard deviation parameters, respectively. The matrices *m* and *s* must be **r-conformable**, where  $i = \max(\text{rows}(m), \text{rows}(s))$  and  $j = \max(\text{cols}(m), \text{cols}(s))$ .

`rpoisson(r, c, m)` returns an  $ir \times jc$  real matrix containing Poisson random variates. The real-valued matrix *m* contains the Poisson mean parameters, where  $i = \text{rows}(m)$  and  $j = \text{cols}(m)$ .

`rt(r, c, df)` returns an  $ir \times jc$  real matrix containing Student's *t* random variates. The real-valued matrix *df* contains the degrees-of-freedom parameters, where  $i = \text{rows}(df)$  and  $j = \text{cols}(df)$ .

## Remarks

The functions described here generate random variates. The parameter limits for each generator are the same as those documented for Stata's [random-number functions](#), except for `rdiscrete()`, which has no Stata equivalent.

In the example below, we generate and summarize 1,000 random normal deviates with a mean of 3 and standard deviation of 1.

```
: rseed(13579)
: x = rnormal(1000, 1, 3, 1)
: meanvariance(x)
      1
1  2.942981879
2  1.061992176
```

The next example uses a  $1 \times 3$  vector of gamma shape parameters to generate a  $1000 \times 3$  matrix of gamma random variates, X.

```
: a = (0.5,1.5,2.5)
: rseed(13579)
: X = rgamma(1000,1,a,1)
: mean(X)
      1          2          3
1  .5339343609  1.510028772  2.451447187
: diagonal(variance(X))'
      1          2          3
1  .6129729256  1.669457192  2.284915684
```

The first column of X contains gamma variates with shape parameter 0.5, the second column contains gamma variates with shape parameter 1.5, and the third column contains gamma variates with shape parameter 2.5.

Below we generate a  $4 \times 3$  matrix of beta variates where we demonstrate the use of two r-conformable parameter matrices, a and b.

```
: a = (0.5,1.5,2.5)
: b = (0.5,0.75,1.0\1.25,1.5,1.75)
: rseed(13579)
: rbeta(2,1,a,b)
      1          2          3
1  .668359305  .3238859912  .7785175363
2  .266459731  .7665943496  .634730294
3  .0373430126 .9246702534  .851879254
4  .1903514438 .5012842811  .8759050005
```

The  $4 \times 3$  shape-parameter matrices used to generate these beta variates are given below:

```
: J(2,1,J(rows(b),1,a))
      1          2          3
1  .5  1.5  2.5
2  .5  1.5  2.5
3  .5  1.5  2.5
4  .5  1.5  2.5
```



: J(2,1,b)

	1	2	3
1	.5	.75	1
2	1.25	1.5	1.75
3	.5	.75	1
4	1.25	1.5	1.75

## Conformability

runiform(*r*, *c*):*r*:  $1 \times 1$ *c*:  $1 \times 1$ *result*:  $r \times c$ 

rseed():

*result*:  $1 \times 1$ rseed(*newseed*):*newseed*:  $1 \times 1$ *result*: *void*rbeta(*r*, *c*, *a*, *b*):*r*:  $1 \times 1$ *c*:  $1 \times 1$ *a*:  $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ *b*:  $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ *result*:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ rbinomial(*r*, *c*, *n*, *p*):*r*:  $1 \times 1$ *c*:  $1 \times 1$ *n*:  $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ *p*:  $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ *result*:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ rchi2(*r*, *c*, *df*):*r*:  $1 \times 1$ *c*:  $1 \times 1$ *df*:  $i \times j$ *result*:  $ir \times jc$ rdiscrete(*r*,*c*,*p*):*r*:  $1 \times 1$ *c*:  $1 \times 1$ *p*:  $k \times 1$ *result*:  $r \times c$

```
rgamma(r, c, a, b):
  r:     $1 \times 1$ 
  c:     $1 \times 1$ 
  a:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
  b:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
result:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ 
```

```
rhypergeometric(r, c, N, K, n):
  r:     $1 \times 1$ 
  c:     $1 \times 1$ 
  N:     $1 \times 1$ 
  K:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
  n:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
result:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ 
```

```
rnbinomial(r, c, n, p):
  r:     $1 \times 1$ 
  c:     $1 \times 1$ 
  n:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
  p:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
result:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ 
```

```
rnormal(r, c, m, s):
  r:     $1 \times 1$ 
  c:     $1 \times 1$ 
  m:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
  s:     $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
result:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ 
```

```
rpoisson(r, c, m):
  r:     $1 \times 1$ 
  c:     $1 \times 1$ 
  m:     $i \times j$ 
result:  $ir \times jc$ 
```

```
rt(r, c, df):
  r:     $1 \times 1$ 
  c:     $1 \times 1$ 
  df:    $1 \times 1$  or  $i \times 1$  or  $1 \times j$  or  $i \times j$ 
result:  $r \times c$  or  $ir \times c$  or  $r \times jc$  or  $ir \times jc$ 
```

## Diagnostics

All random-variate generators abort with an error if  $r < 0$  or  $c < 0$ .

`rseed(seed)` aborts with error if a string seed is specified and it is malformed (was not obtained from `rseed()`).

`rnormal(r, c, m, s)`, `rbeta(r, c, a, b)`, `rbinomial(r, c, n, p)`, `rhypergeometric(r, c, N, K, n)`, and `rnbinomial(r, c, k, p)` abort with an error if the parameter matrices do not conform. See *r-conformability* in [M-6] **Glossary** for rules on matrix conformability.

`rdiscrete()` aborts with error if the probabilities in  $p$  are not in  $[0,1]$  or do not sum to 1.

## Reference

Walker, A. J. 1977. An efficient method for generating discrete random variables with general distributions. *ACM Transactions on Mathematical Software* 3: 253–256.

## Also see

[M-4] **standard** — Functions to create standard matrices

[M-4] **statistical** — Statistical functions

# Title

[M-5] **runningsum()** — Running sum of vector

## Syntax

```
numeric vector runningsum(numeric vector x [, missing])
numeric vector quadrunningsum(numeric vector x [, missing])
void _runningsum(y, numeric vector x [, missing])
void _quadrunningsum(y, numeric vector x [, missing])
```

where optional argument *missing* is a *real scalar* that determines how missing values in *x* are treated:

1. Specifying *missing* as 0 is equivalent to not specifying the argument; missing values in *x* are treated as contributing 0 to the sum.
2. Specifying *missing* as 1 specifies that missing values in *x* are to be treated as missing values and turn the sum to missing.

## Description

`runningsum(x)` returns a vector of the same dimension as *x* containing the running sum of *x*. Missing values are treated as contributing zero to the sum.

`runningsum(x, missing)` does the same but lets you specify how missing values are treated. `runningsum(x, 0)` is the same as `runningsum(x)`. `runningsum(x, 1)` specifies that missing values are to turn the sum to missing where they occur.

`quadrunningsum(x)` and `quadrunningsum(x, missing)` do the same but perform the accumulation in quad precision.

`_runningsum(y, x [, missing])` and `_quadrunningsum(y, x [, missing])` work the same way, except that rather than returning the running-sum vector, they store the result in *y*. This method is slightly more efficient when *y* is a view.

## Remarks

The running sum of (1, 2, 3) is (1, 3, 6).

All functions return the same type as the argument, real if argument is real, complex if complex.

## Conformability

`runningsum(x, missing)`, `quadrunchingsum(x, missing)`:

<i>x</i> :	$r \times 1$	or	$1 \times c$	
<i>missing</i> :	$1 \times 1$			(optional)
<i>result</i> :	$r \times 1$	or	$1 \times c$	

`_runningsum(y, x, missing)`, `_quadrunchingsum(y, x, missing)`:

*input*:

<i>x</i> :	$r \times 1$	or	$1 \times c$	
<i>y</i> :	$r \times 1$	or	$1 \times c$	(contents irrelevant)
<i>missing</i> :	$1 \times 1$			(optional)

*output*:

<i>y</i> :	$r \times 1$	or	$1 \times c$
------------	--------------	----	--------------

## Diagnostics

If *missing* = 0, missing values are treated as contributing zero to the sum; they do not turn the sum to missing. Otherwise, missing values turn the sum to missing.

`_runningsum(y, x, missing)` and `_quadrunchingsum(y, x, missing)` abort with error if *y* is not **p-conformable** with *x* and of the same **eltype**. The contents of *y* are irrelevant.

## Also see

[M-5] **sum()** — Sums

[M-4] **mathematical** — Important mathematical functions

[M-4] **utility** — Matrix utility functions

## Syntax

```

void          schurd(X, T, Q)
void          _schurd(X, Q)

void  schurddgroupby(X, f, T, Q, w, m)
void  _schurddgroupby(X, f,      Q, w, m)
    
```

where inputs are

*X*:    *numeric matrix*  
*f*:    *pointer scalar*    (points to a function used to group eigenvalues)

and outputs are

*T*:    *numeric matrix*    (Schur-form matrix)  
*Q*:    *numeric matrix*    (orthogonal or unitary)  
*w*:    *numeric vector* of eigenvalues  
*m*:    *real scalar*        (the number of eigenvalues satisfy the grouping condition)

## Description

`schurd(X, T, Q)` computes the Schur decomposition of a square, numeric matrix, *X*, returning the [Schur-form](#) matrix, *T*, and the matrix of Schur vectors, *Q*. *Q* is orthogonal if *X* is real and unitary if *X* is complex.

`_schurd(X, Q)` does the same thing as `schurd()`, except that it returns *T* in *X*.

`schurddgroupby(X, f, T, Q, w, m)` computes the Schur decomposition and the eigenvalues of a square, numeric matrix, *X*, and groups the results according to whether a condition on each eigenvalue is satisfied. `schurddgroupby()` returns the Schur-form matrix in *T*, the matrix of Schur vectors in *Q*, the eigenvalues in *w*, and the number of eigenvalues for which the condition is true in *m*. *f* is a pointer of the function that implements the condition on each eigenvalue, as discussed [below](#).

`_schurddgroupby(X, f, Q, w, m)` does the same thing as `schurddgroupby()` except that it returns *T* in *X*.

`_schurd_la()` and `_schurddgroupby_la()` are the interfaces into the LAPACK routines used to implement the above functions; see [\[M-1\] LAPACK](#). Their direct use is not recommended.

## Remarks

Remarks are presented under the following headings:

[Schur decomposition](#)  
[Grouping the results](#)

## Schur decomposition

Many algorithms begin by obtaining the Schur decomposition of a square matrix.

The Schur decomposition of matrix  $\mathbf{X}$  can be written as

$$\mathbf{Q}' \times \mathbf{X} \times \mathbf{Q} = \mathbf{T}$$

where  $\mathbf{T}$  is in Schur form,  $\mathbf{Q}$ , the matrix of Schur vectors, is orthogonal if  $\mathbf{X}$  is real or unitary if  $\mathbf{X}$  is complex.

A real, square matrix is in Schur form if it is block upper triangular with  $1 \times 1$  and  $2 \times 2$  diagonal blocks. Each  $2 \times 2$  diagonal block has equal diagonal elements and opposite sign off-diagonal elements. A complex, square matrix is in Schur form if it is upper triangular. The eigenvalues of  $\mathbf{X}$  are obtained from the Schur form by a few quick computations.

In the example below, we define  $\mathbf{X}$ , obtain the Schur decomposition, and list  $\mathbf{T}$ .

```
: X=(.31,.69,.13,.56\ .31,.5,.72,.42\ .68,.37,.71,.8\ .09,.16,.83,.9)
: schurd(X, T=., Q=.)
: T
```

	1	2	3	4
1	2.10742167	.1266712792	.0549744934	.3329112999
2	0	-.0766307549	.3470959084	.1042286546
3	0	-.4453774705	-.0766307549	.3000409803
4	0	0	0	.4658398402

## Grouping the results

In many applications, there is a stable solution if the modulus of an eigenvalue is less than one and an explosive solution if the modulus is greater than or equal to one. One frequently handles these cases differently and would group the Schur decomposition results into a block corresponding to stable solutions and a block corresponding to explosive solutions.

In the following example, we use `schurgroupby()` to put the stable solutions first. One of the arguments to `schurgroupby()` is a [pointer](#) to a function that accepts a complex scalar argument, an eigenvalue, and returns 1 to select the eigenvalue and 0 otherwise. Here `isstable()` returns 1 if the eigenvalue is less than 1:

```
: real scalar isstable(scalar p)
> {
>     return((abs(p)<1))
> }
```

Using this function to group the results, we see that the Schur-form matrix has been reordered.

```
: schurgroupby(X, &isstable(), T=., Q=., w=., m=.)
: T
```

	1	2	3	4
1	-.0766307549	.445046622	.3029641608	-.0341867415
2	-.3473539401	-.0766307549	-.1036266286	.0799058566
3	0	0	.4658398402	-.3475944606
4	0	0	0	2.10742167

Listing the moduli of the eigenvalues reveals that they are grouped into stable and explosive groups.

: abs(w)				
	1	2	3	4
1	.4005757984	.4005757984	.4658398402	2.10742167

m contains the number of stable solutions

: m  
3

## Conformability

schurd(*X*, *T*, *Q*):

*input:*  
*X*:  $n \times n$   
*output:*  
*T*:  $n \times n$   
*Q*:  $n \times n$

\_schurd(*X*, *Q*):

*input:*  
*X*:  $n \times n$   
*output:*  
*X*:  $n \times n$   
*Q*:  $n \times n$

schurddgroupby(*X*, *f*, *T*, *Q*, *w*, *m*):

*input:*  
*X*:  $n \times n$   
*f*:  $1 \times 1$   
*output:*  
*T*:  $n \times n$   
*Q*:  $n \times n$   
*w*:  $1 \times n$   
*m*:  $1 \times 1$

\_schurddgroupby(*X*, *f*, *Q*, *w*, *m*):

*input:*  
*X*:  $n \times n$   
*f*:  $1 \times 1$   
*output:*  
*X*:  $n \times n$   
*Q*:  $n \times n$   
*w*:  $1 \times n$   
*m*:  $1 \times 1$

## Diagnostics

\_schurd() and \_schurddgroupby() abort with error if *X* is a view.



`schurd()`, `_schurd()`, `schurdgrouby()`, and `_schurdgrouby()` return missing results if  $X$  contains missing values.

`schurdgrouby()` groups the results via a matrix transform. If the problem is very ill conditioned, applying this matrix transform can cause changes in the eigenvalues. In extreme cases, the grouped eigenvalues may no longer satisfy the condition used to perform the grouping.

Issai Schur (1875–1941) was born in Mogilev, which is now in Belarus. He studied mathematics and physics at the University of Berlin, where Frobenius was one of his teachers. Schur obtained his doctorate with a thesis on rational representations of the general linear group over the complex field. Thereafter, he taught and researched at Berlin University (with an interval in Bonn) until he was dismissed by the Nazis in 1935. He was a superb lecturer. Schur is now known mainly for his fundamental contributions to the representation theory of groups, but he also worked in other areas of algebra, including matrices, number theory, and analysis. In 1939, he emigrated to Palestine, where he later died in poverty.

## Reference

Ledermann, W. 1983. Issai Schur and his school in Berlin. *Bulletin of the London Mathematical Society* 15: 97–106.

## Also see

[M-1] [LAPACK](#) — The LAPACK linear-algebra routines

[M-5] [hessenbergd\(\)](#) — Hessenberg decomposition

[M-4] [matrix](#) — Matrix functions

# Title

[M-5] **select()** — Select rows or columns

## Syntax

*transmorphic matrix* **select**(*transmorphic matrix*  $X$ , *real vector*  $v$ )  
*void*                    **st\_select**( $A$ , *transmorphic matrix*  $X$ , *real vector*  $v$ )

## Description

**select**( $X$ ,  $v$ ) returns  $X$

1. omitting the rows for which  $v[i] == 0$  ( $v$  a column vector) or
2. omitting the columns for which  $v[j] == 0$  ( $v$  a row vector).

**st\_select**( $A$ ,  $X$ ,  $v$ ) does the same thing, except that the result is placed in  $A$  and, if  $X$  is a view,  $A$  will be a view.

## Remarks

Remarks are presented under the following headings:

*Examples*  
*Using st\_select()*

## Examples

1. To select rows 1, 2, and 4 of  $5 \times c$  matrix  $X$ ,

```
submat = select(X, (1\1\0\1\0))
```

See [M-2] **subscripts** for another solution, `submat = X[(1\2\4), .]`.

2. To select columns 1, 2, and 4 of  $r \times 5$  matrix  $X$ ,

```
submat = select(X, (1,1,0,1,0))
```

See [M-2] **subscripts** for another solution, `submat = X[:, (1,2,4)]`.

3. To select rows of  $X$  for which the first element is positive,

```
submat = select(X, X[:,1]:>0)
```

4. To select columns of  $X$  for which the first element is positive,

```
submat = select(X, X[1,:]:>0)
```

5. To select rows of  $X$  for which there are no missing values,

```
submat = select(X, rowmissing(X)==0)
```

6. To select rows and columns of square matrix  $X$  for which the diagonal elements are positive,

```
pos      = diagonal(X):>0
submat   = select(X, pos)
submat   = select(submat, pos')
```

or, equivalently,

```
pos      = diagonal(X):>0
submat   = select(select(X, pos), pos')
```

## Using st\_select()

Coding

```
st_select(submat, X, v) (1)
```

produces the same result as coding

```
submat = st_select(X, v) (2)
```

The difference is in how the result is stored. If  $X$  is a view (it need not be), then (1) will produce  $\text{submat}$  as a view or, if you will, a subview, whereas in (2),  $\text{submat}$  will always be a regular (nonview) matrix.

When  $X$  is a view, (1) executes more quickly than (2) and produces a result that consumes less memory.

See [M-5] [st\\_view\(\)](#) for a description of views.

## Conformability

`select(X, v):`

$X:$	$r_1 \times c_1$			
$v:$	$r_1 \times 1$	or	$1 \times c_1$	
<i>result:</i>	$r_2 \times c_1$	or	$r_1 \times c_2$ ,	$r_2 \leq r_1, c_2 \leq c_1$

`st_select(A, X, v):`

*input:*

$X:$	$r_1 \times c_1$		
$v:$	$r_1 \times 1$	or	$1 \times c_1$

*output:*

$A:$	$r_2 \times c_1$	or	$r_1 \times c_2$ ,	$r_2 \leq r_1, c_2 \leq c_1$
------	------------------	----	--------------------	------------------------------

## Diagnostics

None.

## Also see

[M-5] [st\\_subview\(\)](#) — Make view from view

[M-2] [op\\_colon](#) — Colon operators

[M-2] [subscripts](#) — Use of subscripts

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] **setbreakintr()** — Break-key processing

## Syntax

```
real scalar  setbreakintr(real scalar val)  
real scalar  querybreakintr()  
real scalar  breakkey()  
void         breakkeyreset()
```

## Description

`setbreakintr(val)` turns the break-key interrupt off (*val*==0) or on (*val*!=0) and returns the value of the previous break-key mode, 1, it was on, or 0, it was off.

`querybreakintr()` returns 1 if the break-key interrupt is on and 0 otherwise.

`breakkey()` (for use in `setbreakintr(0)` mode) returns 1 if the break key has been pressed since it was last reset.

`breakkeyreset()` (for use in `setbreakintr(0)` mode) resets the break key.

## Remarks

Remarks are presented under the following headings:

[\*Default break-key processing\*](#)  
[\*Suspending the break-key interrupt\*](#)  
[\*Break-key polling\*](#)

### Default break-key processing

By default, if the user presses *Break*, Mata stops execution and returns control to the console, setting the return code to 1.

To obtain this behavior, there is nothing you need do. You do not need to use these functions.

### Suspending the break-key interrupt

The default behavior is known as interrupt-on-break and is also known as `setbreakintr(1)` mode.

The alternative is break-key suspension, also known as `setbreakintr(0)` mode.

For instance, you have several steps that must be performed in their entirety or not at all. The way to do this is

```
val = setbreakintr(0)
...
... (critical code) ...
...
(void) setbreakintr(val)
```

The first line stores in *val* the current break-key processing mode and then sets the mode to break-key suspension. The critical code then runs. If the user presses *Break* during the execution of the critical code, that will be ignored. Finally, the code restores the previous break-key processing mode.

## Break-key polling

In coding large, interactive systems, you may wish to adopt the break-key polling style of coding rather than interrupt-on-break. In this alternative style of coding, you turn off interrupt-on-break:

```
val = setbreakintr(0)
```

and, from then on in your code, wherever you are willing to interrupt your code, you ask (poll whether) the break key has been pressed:

```
...
if (breakkey()) {
    ...
}
...
```

In this style of coding, you must decide where and when you are going to reset the break key, because once the break key has been pressed, `breakkey()` will continue to return 1 every time it is called. To reset the break key, code,

```
breakkeyreset()
```

You can also adopt a mixed style of coding, using interrupt-on-break in some places and polling in others. Function `querybreakintr()` can then be used to determine the current mode.

## Conformability

`setbreakintr(val):`

```
    val:      1 × 1
    result:    1 × 1
```

`querybreakintr(), breakkey():`

```
    result:    1 × 1
```

`breakkeyreset():`

```
    result:    void
```

## Diagnostics

`setbreakintr(1)` aborts with `break` if the break key has been pressed since the last `setbreakintr(0)` or `breakkeyreset()`. Code `breakkeyreset()` before `setbreakintr(1)` if you do not want this behavior.

After coding `setbreakintr(1)`, remember to restore `setbreakintr(0)` mode. It is not, however, necessary, to restore the original mode if `exit()` or `_error()` is about to be executed.

`breakkey()`, once the break key has been pressed, continues to return 1 until `breakkeyreset()` is executed.

There is absolutely no reason to use `breakkey()` in `setbreakintr(0)` mode, because the only value it could return is 0.

## Also see

[M-5] `error()` — Issue error message

[M-4] `programming` — Programming functions

Syntax

```
real matrix  sign(real matrix R)

real matrix  quadrant(complex matrix Z)
```

Description

`sign(R)` returns the elementwise sign of *R*. `sign()` is defined

Argument range	<code>sign(arg)</code>
$arg \geq .$	$.$
$arg < 0$	$-1$
$arg = 0$	$0$
$arg > 0$	$1$

`quadrant(Z)` returns a real matrix recording the quadrant of each complex entry in *Z*. `quadrant()` is defined

Argument range		<code>quadrant(arg)</code>
<code>Re(arg)</code>	<code>Im(arg)</code>	
$Re \geq .$		$.$
$Re = 0$	$Im = 0$	$.$
$Re > 0$	$Im \geq 0$	$1$
$Re \leq 0$	$Im > 0$	$2$
$Re < 0$	$Im \leq 0$	$3$
$Re \geq 0$	$Im < 0$	$4$

```
quadrant(1+0i)==1, quadrant(-1+0i)==3
quadrant(0+1i)==2, quadrant( 0-1i)==4
```

Conformability

```
sign(R):
    R:      r × c
    result: r × c

quadrant(Z):
    Z:      r × c
    result: r × c
```



## Diagnostics

`sign(R)` returns missing when *R* is missing.

`quadrant(Z)` returns missing when *Z* is missing.

## Also see

[M-5] `dsign()` — FORTRAN-like `DSIGN()` function

[M-4] `scalar` — Scalar mathematical functions

## Syntax

<i>numeric matrix</i>	<b>sin</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>cos</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>tan</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>asin</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>acos</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>atan</b> ( <i>numeric matrix Z</i> )
<i>real matrix</i>	<b>atan2</b> ( <i>real matrix X</i> , <i>real matrix Y</i> )
<i>real matrix</i>	<b>arg</b> ( <i>complex matrix Z</i> )
<i>numeric matrix</i>	<b>sinh</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>cosh</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>tanh</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>asinh</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>acosh</b> ( <i>numeric matrix Z</i> )
<i>numeric matrix</i>	<b>atanh</b> ( <i>numeric matrix Z</i> )
<i>real scalar</i>	<b>pi</b> ()

## Description

**sin**(*Z*), **cos**(*Z*), and **tan**(*Z*) return the appropriate trigonometric functions. Angles are measured in radians. All return real if the argument is real and complex if the argument is complex.

**sin**(*x*), *x* real, returns the sine of *x*. **sin**() returns a value between  $-1$  and  $1$ .

**sin**(*z*), *z* complex, returns the complex sine of *z*, mathematically defined as  $\{\exp(i * z) - \exp(-i * z)\} / 2i$ .

**cos**(*x*), *x* real, returns the cosine of *x*. **cos**() returns a value between  $-1$  and  $1$ .

**cos**(*z*), *z* complex, returns the complex cosine of *z*, mathematically defined as  $\{\exp(i * z) + \exp(-i * z)\} / 2$ .

**tan**(*x*), *x* real, returns the tangent of *x*.

**tan(*z*)**, *z* complex, returns the complex tangent of *z*, mathematically defined as  $\sin(z)/\cos(z)$ .

**asin(*Z*)**, **acos(*Z*)**, and **atan(*Z*)** return the appropriate inverse trigonometric functions. Returned results are in radians. All return real if the argument is real and complex if the argument is complex.

**asin(*x*)**, *x* real, returns arcsine in the range  $[-\pi/2, \pi/2]$ . If  $x < -1$  or  $x > 1$ , missing (.) is returned.

**asin(*z*)**, *z* complex, returns the complex arcsine, mathematically defined as  $-i * \ln\{i * z + \sqrt{1 - z * z}\}$ . **Re(asin())** is chosen to be in the interval  $[-\pi/2, \pi/2]$ .

**acos(*x*)**, *x* real, returns arccosine in the range  $[0, \pi]$ . If  $x < -1$  or  $x > 1$ , missing (.) is returned.

**acos(*z*)**, *z* complex, returns the complex arccosine, mathematically defined as  $-i * \ln\{z + \sqrt{z * z - 1}\}$ . **Re(acos())** is chosen to be in the interval  $[0, \pi]$ .

**atan(*x*)**, *x* real, returns arctangent in the range  $(-\pi/2, \pi/2)$ .

**atan(*z*)**, *z* complex, returns the complex arctangent, mathematically defined as  $\ln\{(1 + iz)/(1 - iz)\}/(2i)$ . **Re(atan())** is chosen to be in the interval  $[0, \pi]$ .

**atan2(*X*, *Y*)** returns the radian value in the range  $(-\pi, \pi]$  of the angle of the vector determined by (*X*,*Y*), the result being in the range  $[0, \pi]$  for quadrants 1 and 2 and  $[0, -\pi]$  for quadrants 4 and 3. *X* and *Y* must be real. **atan2(*X*, *Y*)** is equivalent to **arg(C(*X*, *Y*))**.

**arg(*Z*)** returns the arctangent of  $\text{Im}(Z)/\text{Re}(Z)$  in the correct quadrant, the result being in the range  $(-\pi, \pi]$ ;  $[0, \pi]$  in quadrants 1 and 2 and  $[0, -\pi]$  in quadrants 4 and 3. **arg(*Z*)** is equivalent to **atan2(Re(*Z*), Im(*Z*))**.

**sinh(*Z*)**, **cosh(*Z*)**, and **tanh(*Z*)** return the hyperbolic sine, cosine, and tangent, respectively. The returned value is real if the argument is real and complex if the argument is complex.

**sinh(*x*)**, *x* real, returns the inverse hyperbolic sine of *x*, mathematically defined as  $\{\exp(x) - \exp(-x)\}/2$ .

**sinh(*z*)**, *z* complex, returns the complex hyperbolic sine of *z*, mathematically defined as  $\{\exp(z) - \exp(-z)\}/2$ .

**cosh(*x*)**, *x* real, returns the inverse hyperbolic cosine of *x*, mathematically defined as  $\{\exp(x) + \exp(-x)\}/2$ .

**cosh(*z*)**, *z* complex, returns the complex hyperbolic cosine of *z*, mathematically defined as  $\{\exp(z) + \exp(-z)\}/2$ .

**tanh(*x*)**, *x* real, returns the inverse hyperbolic tangent of *x*, mathematically defined as  $\sinh(x)/\cosh(x)$ .

**tanh(*z*)**, *z* complex, returns the complex hyperbolic tangent of *z*, mathematically defined as  $\sinh(z)/\cosh(z)$ .

**asinh(*Z*)**, **acosh(*Z*)**, and **atanh(*Z*)** return the inverse hyperbolic sine, cosine, and tangent, respectively. The returned value is real if the argument is real and complex if the argument is complex.

**asinh(*x*)**, *x* real, returns the inverse hyperbolic sine.

`asinh(z)`,  $z$  complex, returns the complex inverse hyperbolic sine, mathematically defined as  $\ln\{z + \sqrt{z * z + 1}\}$ . `Im(asinh())` is chosen to be in the interval  $[-\pi/2, \pi/2]$ .

`acosh(x)`,  $x$  real, returns the inverse hyperbolic cosine. If  $x < 1$ , missing (.) is returned.

`acosh(z)`,  $z$  complex, returns the complex inverse hyperbolic cosine, mathematically defined as  $\ln\{z + \sqrt{z * z - 1}\}$ . `Im(acosh())` is chosen to be in the interval  $[-\pi, \pi]$ ; `Re(acosh())` is chosen to be nonnegative.

`atanh(x)`,  $x$  real, returns the inverse hyperbolic tangent. If  $|x| > 1$ , missing (.) is returned.

`atanh(z)`,  $z$  complex, returns the complex inverse hyperbolic tangent, mathematically defined as  $\ln\{(1 + z)/(1 - z)\}/2$ . `Im(atanh())` is chosen to be in the interval  $[-\pi/2, \pi/2]$ .

`pi()` returns the value of  $\pi$ .

## Conformability

```
atan2(X, Y):
  X:       $r_1 \times c_1$ 
  Y:       $r_2 \times c_2$ ,    $X$  and  $Y$  r-conformable
  result:  $\max(r_1, r_2) \times \max(c_1, c_2)$ 
```

`pi()` returns a  $1 \times 1$  scalar.

All other functions return a matrix of the same dimension as input containing element-by-element calculated results.

## Diagnostics

All functions return missing for real arguments when the result would be complex. For instance, `acos(2) = .`, whereas `acos(2+0i) = -1.317i`.

## Also see

[M-4] [scalar](#) — Scalar mathematical functions

# Title

[M-5] **sizeof()** — Number of bytes consumed by object

## Syntax

*real scalar*    `sizeof(transmorphic matrix A)`

## Description

`sizeof(A)` returns the number of bytes consumed by *A*.

## Remarks

`sizeof(A)` returns the same number as shown by `mata describe`; see [M-3] [mata describe](#).

A  $500 \times 5$  real matrix consumes 20,000 bytes:

```
: sizeof(mymatrix)
20000
```

A  $500 \times 5$  view matrix, however, consumes only 24 bytes:

```
: sizeof(myview)
24
```

To obtain the number of bytes consumed by a function, pass a dereferenced function pointer:

```
: sizeof(&myfcn())
320
```

## Conformability

`sizeof(A)`:

<i>A</i> :	$r \times c$
<i>result</i> :	$1 \times 1$

## Diagnostics

The number returned by `sizeof(A)` does not include any overhead, which usually amounts to 64 bytes, but can be less (as small as zero in the case of recently used scalars).

If *A* is a pointer matrix, the number returned reflects the amount of memory required to store *A* itself and does not include the memory consumed by its siblings.

## Also see

[M-4] [programming](#) — Programming functions

# Title

[M-5] solve\_tol() — Tolerance used by solvers and inverters

## Syntax

*real scalar*    solve\_tol(*numeric matrix Z, real scalar usertol*)

## Description

solve\_tol(*Z, usertol*) returns the tolerance used by many Mata solvers to solve  $AX = B$  and by many Mata inverters to obtain  $A^{-1}$ . *usertol* is the tolerance specified by the user or is missing value if the user did not specify a tolerance.

## Remarks

The tolerance used by many Mata solvers to solve  $AX = B$  and by many Mata inverters to obtain  $A^{-1}$  is

$$eta = s * \frac{\text{trace}(\text{abs}(Z))}{n} \tag{1}$$

where  $s = 1\text{e-}13$  or a value specified by the user,  $n$  is the `min(rows(Z), cols(Z))`, and  $Z$  is a matrix related to  $A$ , usually by some form of decomposition, but could be  $A$  itself (for instance, if  $A$  were triangular). See, for instance, [M-5] `solve_lower()` and [M-5] `cholsolve()`.

When *usertol* > 0 and *usertol* < . is specified, solve\_tol() returns *eta* calculated with  $s = \text{usertol}$ .

When *usertol* ≤ 0 is specified, solve\_tol() returns  $-\text{usertol}$ .

When *usertol* ≥ . is specified, solve\_tol() returns a default result, calculated as

1. If external real scalar `_solveterolerance` does not exist, as is usually the case, the value of *eta* is returned using  $s = 1\text{e-}13$ .
2. If external real scalar `_solveterolerance` does exist,
  - a. If `_solveterolerance` > 0, the value of *eta* is returned using  $s = \text{solveterolerance}$ .
  - b. If `_solveterolerance` ≤ 0,  $-\text{_solveterolerance}$  is returned.

## Conformability

solve\_tol(*Z, usertol*):

<i>Z</i> :	$r \times c$
<i>usertol</i> :	$1 \times 1$
<i>result</i> :	$1 \times 1$

## Diagnostics

`solve_tol(Z, usertol)` skips over missing values in  $Z$  in calculating (1);  $n$  is defined as the number of nonmissing elements on the diagonal.

## Also see

[M-4] [utility](#) — Matrix utility functions

## Syntax

*numeric matrix*     `solverlower(A, B [, rank [, tol [, d ]]])`

*numeric matrix*     `solveupper(A, B [, rank [, tol [, d ]]])`

*real scalar*         `_solverlower(A, B [, tol [, d ]])`

*real scalar*         `_solveupper(A, B [, tol [, d ]])`

where

<i>A</i> :	<i>numeric matrix</i>
<i>B</i> :	<i>numeric matrix</i>
<i>rank</i> :	irrelevant; <i>real scalar</i> returned
<i>tol</i> :	<i>real scalar</i>
<i>d</i> :	<i>numeric scalar</i>

## Description

These functions are used in the implementation of the other solve functions; see [\[M-5\] lusolve\(\)](#), [\[M-5\] qrsolve\(\)](#), and [\[M-5\] svsolve\(\)](#).

`solverlower(A, B, ...)` and `_solverlower(A, B, ...)` solve lower-triangular systems.

`solveupper(A, B, ...)` and `_solveupper(A, B, ...)` solve upper-triangular systems.

Functions without a leading underscore—`solverlower()` and `solveupper()`—return the solution;  $A$  and  $B$  are unchanged.

Functions with a leading underscore—`_solverlower()` and `_solveupper()`—return the solution in  $B$ .

All four functions produce a generalized solution if  $A$  is singular. The functions without an underscore place the rank of  $A$  in *rank*, if the argument is specified. The underscore functions return the rank.

Determination of singularity is made via *tol*. *tol* is interpreted in the standard way—as a multiplier for the default if  $tol > 0$  is specified and as an absolute quantity to use in place of the default if  $tol \leq 0$  is specified.

All four functions allow  $d$  to be optionally specified. Specifying  $d = .$  is equivalent to not specifying  $d$ .

If  $d \neq .$  is specified, that value is used as if it appeared on the diagonal of  $A$ . The four functions do not in fact require that  $A$  be triangular; they merely look at the lower or upper triangle and pretend that the opposite triangle contains zeros. This feature is useful when a decomposition utility has stored both the lower and upper triangles in one matrix, because one need not take apart the combined matrix. In such cases, it sometimes happens that the diagonal of the matrix corresponds to one matrix but not the other, and that for the other matrix, one merely knows that the diagonal elements are, say, 1. Then you can specify  $d = 1$ .



## Remarks

The triangular-solve functions documented here exploit the triangular structure in  $A$  and solve for  $X$  by recursive substitution.

When  $A$  is of full rank, these functions provide the same solution as the other solve functions, such as [M-5] `lusolve()`, [M-5] `qrsolve()`, and [M-5] `svsolve()`. The `solverlower()` and `solveupper()` functions, however, will produce the answer more quickly because of the large computational savings.

When  $A$  is singular, however, you may wish to consider whether you want to use these triangular-solve functions. The triangular-solve functions documented here reach a generalized solution by setting  $B_{ij} = 0$ , for all  $j$ , when  $A_{ij}$  is zero or too small (as determined by *tol*). The method produces a generalized inverse, but there are many generalized inverses, and this one may not have the other properties you want.

Remarks are presented under the following headings:

*Derivation*  
*Tolerance*

## Derivation

We wish to solve

$$AX = B \quad (1)$$

when  $A$  is triangular. Let us consider the lower-triangular case first. `solverlower()` is up to handling full matrices for  $B$  and  $X$ , but let us assume  $X$ :  $n \times 1$  and  $B$ :  $m \times 1$ :

$$\begin{bmatrix} a_{11} & 0 & 0 \dots & 0 \\ a_{21} & 0 & 0 \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ a_{m1} & a_{m2} & a_{m3} & a_{mn} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_n \end{bmatrix} = \begin{bmatrix} b_1 \\ b_2 \\ \vdots \\ b_m \end{bmatrix}$$

The first equation to be solved is

$$a_{11}x_1 = b_1$$

and the solution is simply

$$x_1 = \frac{b_1}{a_{11}} \quad (2)$$

The second equation to be solved is

$$a_{21}x_1 + a_{22}x_2 = b_2$$

and because we have already solved for  $x_1$ , the solution is simply

$$x_2 = \frac{b_2 - a_{21}x_1}{a_{22}} \quad (3)$$

We proceed similarly for the remaining rows of  $A$ . If there are additional columns in  $B$  and  $X$ , we can then proceed to handling each remaining column just as we handled the first column above.

In the upper-triangular case, the formulas are similar except that you start with the last row of  $A$ .

## Tolerance

In (2) and (3), we divide by the diagonal elements of  $A$ . If element  $a_{ii}$  is less than  $eta$  in absolute value, the corresponding  $x_i$  is set to zero.  $eta$  is given by

$$eta = 1e-13 * \text{trace}(\text{abs}(A)) / \text{rows}(A)$$

If you specify  $tol > 0$ , the value you specify is used to multiply  $eta$ . You may instead specify  $tol \leq 0$ , and then the negative of the value you specify is used in place of  $eta$ ; see [M-1] [tolerance](#).

## Conformability

`solverlower(A, B, rank, tol, d)`, `solveupper(A, B, rank, tol, d)`:

*input:*

$A:$       $n \times n$   
 $B:$       $n \times k$   
 $tol:$      $1 \times 1$  (optional)  
 $d:$       $1 \times 1$  (optional)

*output:*

$rank:$      $1 \times 1$  (optional)  
 $result:$     $n \times k$

`_solverlower(A, B, tol, d)`, `_solveupper(A, B, tol, d)`:

*input:*

$A:$       $n \times n$   
 $B:$       $n \times k$   
 $tol:$      $1 \times 1$  (optional)  
 $d:$       $1 \times 1$  (optional)

*output:*

$B:$       $n \times k$   
 $result:$     $1 \times 1$  (contains rank)

## Diagnostics

`solverlower(A, B, ...)`, `_solverlower(A, B, ...)`, `solveupper(A, B, ...)`, and `_solveupper(A, B, ...)` do not verify that the upper (lower) triangle of  $A$  contains zeros; they just use the lower (upper) triangle of  $A$ .

`_solverlower(A, B, ...)` and `_solveupper(A, B, ...)` do not abort with error if  $B$  is a view but can produce results subject to considerable roundoff error.

## Also see

[M-5] [cholsolve\(\)](#) — Solve  $AX=B$  for  $X$  using Cholesky decomposition

[M-5] [lusolve\(\)](#) — Solve  $AX=B$  for  $X$  using LU decomposition

[M-5] [qrsolve\(\)](#) — Solve  $AX=B$  for  $X$  using QR decomposition

[M-5] [svsolve\(\)](#) — Solve  $AX=B$  for  $X$  using singular value decomposition

[M-5] [solve\\_tol\(\)](#) — Tolerance used by solvers and inverters

[M-4] [matrix](#) — Matrix functions

## Syntax

<i>transmorphic matrix</i>	<code>sort(transmorphic matrix X, real rowvector idx)</code>
<i>void</i>	<code>_sort(transmorphic matrix X, real rowvector idx)</code>
<i>transmorphic matrix</i>	<code>jumble(transmorphic matrix X)</code>
<i>void</i>	<code>_jumble(transmorphic matrix X)</code>
<i>real colvector</i>	<code>order(transmorphic matrix X, real rowvector idx)</code>
<i>real colvector</i>	<code>unordered(real scalar n)</code>
<i>void</i>	<code>_collate(transmorphic matrix X, real colvector p)</code>

where

1.  $X$  may not be a pointer matrix.
2.  $p$  must be a permutation column vector, a  $1 \times c$  vector containing the integers  $1, 2, \dots, c$  in some order.

## Description

`sort(X, idx)` returns  $X$  with rows in ascending or descending order of the columns specified by  $idx$ . For instance, `sort(X, 1)` sorts  $X$  on its first column; `sort(X, (1,2))` sorts  $X$  on its first and second columns (meaning rows with equal values in their first column are ordered on their second column). In general, the  $i$ th sort key is column `abs(idx[i])`. Order is ascending if `idx[i] > 0` and descending otherwise. Ascending and descending are defined in terms of [\[M-5\] abs\(\)](#) (length of elements) for complex.

`_sort(X, idx)` does the same as `sort(X, idx)`, except that  $X$  is sorted in place.

`jumble(X)` returns  $X$  with rows in random order. For instance, to shuffle a deck of cards numbered 1 to 52, one could code `jumble(1::52)`. See `rseed()` in [\[M-5\] runiform\(\)](#) for information on setting the random-number seed.

`_jumble(X)` does the same as `jumble(X)`, except that  $X$  is jumbled in place.

`order(X, idx)` returns the permutation vector—see [\[M-1\] permutation](#)—that would put  $X$  in ascending (descending) order of the columns specified by  $idx$ . A row-permutation vector is a  $1 \times c$  column vector containing the integers  $1, 2, \dots, c$  in some order. Vectors  $(1\backslash 2\backslash 3)$ ,  $(1\backslash 3\backslash 2)$ ,  $(2\backslash 1\backslash 3)$ ,  $(2\backslash 3\backslash 1)$ ,  $(3\backslash 1\backslash 2)$ , and  $(3\backslash 2\backslash 1)$  are examples. Row-permutation vectors are used to specify the order in which the rows of a matrix  $X$  are to appear. If  $p$  is a row-permutation vector, `X[p, .]` returns  $X$  with its rows in the order of  $p$ ;  $p = (3\backslash 2\backslash 1)$  would reverse the rows of  $X$ . `order(X, idx)` returns the row-permutation vector that would sort  $X$  and, as a matter of fact, `sort(X, idx)` is implemented as `X[order(X, idx), .]`.

`unorder(n)` returns a  $1 \times n$  permutation vector for placing the rows in random order. Random numbers are calculated by `runiform()`; see `rseed()` in [M-5] `runiform()` for information on setting the random-number seed. `jumble()` is implemented in terms of `unorder()`: `jumble(X)` is equivalent to `X[unorder(rows(X)), .]`.

`_collate(X, p)` is equivalent to `X = X[p, .]`; it changes the order of the rows of *X*. `_collate()` is used by `_sort()` and `_jumble()` and has the advantage over subscripting in that no extra memory is required when the result is to be assigned back to itself. Consider

$$X = X[p, .]$$

There will be an instant after `X[p, .]` has been calculated but before the result has been assigned back to *X* when two copies of *X* exist. `_collate(X, p)` avoids that. `_collate()` is not a substitute for subscripting in all cases; `_collate()` requires *p* be a permutation vector.

## Remarks

If *X* is complex, the ordering is defined in terms of [M-5] `abs()` of its elements.

Also see `invorder()` and `revorder()` in [M-5] `invorder()`. Let *p* be the permutation vector returned by `order()`:

$$p = \text{order}(X, \dots)$$

Then `X[p, .]` are the sorted rows of *X*. `revorder()` can be used to reverse sort order: `X[revorder(p), .]` are the rows of *X* in the reverse of the order of `X[p, .]`. `invorder()` provides the inverse transform: If `Y = X[p, .]`, then `X = Y[invorder(p), .]`.

## Conformability

`sort(X, idx)`, `jumble(X)`:

<i>X</i> :	$r_1 \times c_1$
<i>idx</i> :	$1 \times c_2, c_2 \leq c_1$
result:	$r_1 \times c_1$

`_sort(X, idx)`, `_jumble(X)`:

<i>X</i> :	$r_1 \times c_1$
<i>idx</i> :	$1 \times c_2, c_2 \leq c_1$
result:	void; <i>X</i> row order modified

`order(X, idx)`:

<i>X</i> :	$r_1 \times c_1$
<i>idx</i> :	$1 \times c_2, c_2 \leq c_1$
result:	$r_1 \times 1$

`unorder(n)`:

<i>n</i> :	$1 \times 1$
result:	$n \times 1$

`_collate(X, p)`:

<i>X</i> :	$r \times c$
<i>p</i> :	$r \times 1$
result:	void; <i>X</i> row order modified

## Diagnostics

`sort(X, idx)` aborts with error if any element of `abs(idx)` is less than 1 or greater than `rows(X)`.

`_sort(X, idx)` aborts with error if any element of `abs(idx)` is less than 1 or greater than `rows(X)`, or if `X` is a view.

`_jumble(X)` aborts with error if `X` is a view.

`order(X, idx)` aborts with error if any element of `abs(idx)` is less than 1 or greater than `rows(X)`.

`unorder(n)` aborts with error if  $n < 1$ .

`_collate(X, p)` aborts with error if `p` is not a permutation vector or if `X` is a view.

## Also see

[M-5] [invector\(\)](#) — Permutation vector manipulation

[M-5] [uniqrows\(\)](#) — Obtain sorted, unique values

[M-4] [manipulation](#) — Matrix manipulation

# Title

[M-5] **soundex()** — Convert string to soundex code

## Syntax

*string matrix*            **soundex**(*string matrix s*)

*string matrix* **soundex\_nara**(*string matrix s*)

## Description

**soundex**(*s*) returns the soundex code for a string, *s*. The soundex code consists of a letter followed by three numbers: the letter is the first letter of the name and the numbers encode the remaining consonants. Similar sounding consonants are encoded by the same number.

**soundex\_nara**(*s*) returns the U.S. Census soundex code for a string, *s*. The soundex code consists of a letter followed by three numbers: the letter is the first letter of the name and the numbers encode the remaining consonants. Similar sounding consonants are encoded by the same number.

When *s* is not a scalar, these functions return element-by-element results.

## Remarks

**soundex**("Ashcraft") returns "A226".

**soundex\_nara**("Ashcraft") returns "A261".

## Conformability

**soundex**(*s*), **soundex\_nara**(*s*):

<i>s</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

None.

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] **spline3()** — Cubic spline interpolation

## Syntax

*real matrix*    **spline3**(*real vector x*, *real vector y*)

*real vector*    **spline3eval**(*real matrix spline\_info*, *real vector x*)

## Description

**spline3**(*x*, *y*) returns the coefficients of a cubic natural spline  $S(x)$ . The elements of *x* must be strictly monotone increasing.

**spline3eval**(*spline\_info*, *x*) uses the information returned by **spline3**() to evaluate and return the spline at the abscissas *x*. Elements of the returned result are set to missing if outside the range of the spline. *x* is assumed to be monotonically increasing.

## Remarks

**spline3**() and **spline3eval**() is a translation into Mata of [Herriot and Reinsch](#) (CUBNATSPLINE) (1973).

For  $xx$  in  $[x_i, x_{i+1})$ :

$$S(xx) = \{(d_i t + c_i)t + b_i\}t + y_i$$

with  $t = xx - x_i$ .

**spline3**() returns  $(b, c, d, x, y)$  or, if *x* and *y* are row vectors,  $(b, c, d, x', y')$ .

## Conformability

**spline3**(*x*, *y*):

<i>x</i> :	$n \times 1$	or	$1 \times n$
<i>y</i> :	$n \times 1$	or	$1 \times n$
<i>result</i> :	$n \times 5$		

**spline3eval**(*spline\_info*, *x*):

<i>spline_info</i> :	$n \times 5$		
<i>x</i> :	$m \times 1$	or	$1 \times m$
<i>result</i> :	$m \times 1$	or	$1 \times m$

## Diagnostics

**spline3**(*x*, *y*) requires that *x* be in ascending order.

**spline3eval**(*spline\_info*, *x*) requires that *x* be in ascending order.

## Reference

Herriot, J. G., and C. H. Reinsch. 1973. Algorithm 472: Procedures for natural spline interpolation [E1]. *Communications of the ACM* 16: 763–768.

## Also see

[\[M-4\] mathematical](#) — Important mathematical functions



# Title

[M-5] **sqrt()** — Square root

## Syntax

*numeric matrix*    **sqrt**(*numeric matrix Z*)

## Description

**sqrt**(*Z*) returns the elementwise square root of *Z*.

## Conformability

<b>sqrt</b> ( <i>Z</i> )	
<i>Z</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

**sqrt**(*Z*) returns missing when *Z* is real and  $Z < 0$ ; that is, **sqrt**(-4) = . but **sqrt**(-4+0i) = 2i.

## Also see

- [M-5] **cholesky()** — Cholesky square-root decomposition
- [M-4] **scalar** — Scalar mathematical functions

## Syntax

```

void          st_addobs(real scalar n)

void          st_addobs(real scalar n, real scalar nofill)

real scalar  _st_addobs(real scalar n)

real scalar  _st_addobs(real scalar n, real scalar nofill)
    
```

## Description

`st_addobs(n)` adds *n* observations to the current Stata dataset.

`st_addobs(n, nofill)` does the same thing but saves computer time by not filling in the additional observations with the appropriate missing-value code if *nofill*  $\neq$  0. `st_addobs(n, 0)` is equivalent to `st_addobs(n)`. Use of `st_addobs()` with *nofill*  $\neq$  0 is not recommended. If you specify *nofill*  $\neq$  0, it is your responsibility to ensure that the added observations ultimately are filled in or removed before control is returned to Stata.

`_st_addobs(n)` and `_st_addobs(n, nofill)` perform the same action as `st_addobs(n)` and `st_addobs(n, nofill)`, except that they return 0 if successful and the appropriate Stata return code otherwise (otherwise usually being caused by insufficient memory). Where `_st_addobs()` would return nonzero, `st_addobs()` aborts with error.

## Remarks

There need not be any variables defined to add observations. If you are attempting to create a dataset from nothing, you can add the observations first and then add the variables, or you can add the variables and then add the observations. Use `st_addvar()` (see [\[M-5\] st\\_addvar\(\)](#)) to add variables.

## Conformability

```

st_addobs(n, nofill):
    n:          1 × 1
    nofill:     1 × 1   (optional)
    result:     void

_st_addobs(n, nofill):
    n:          1 × 1
    nofill:     1 × 1   (optional)
    result:     1 × 1
    
```

## Diagnostics

`st_addobs(n [, nofill])` and `_st_addobs(n [, nofill])` abort with error if  $n < 0$ . They do nothing if  $n = 0$ .

`st_addobs()` aborts with error if there is insufficient memory to add the requested number of observations.

`_st_addobs()` aborts with error if  $n < 0$  but otherwise returns the appropriate Stata return code if the observations cannot be added. If they are added, 0 is returned.

`st_addobs()` and `_st_addobs()` do not set `st_updata()` (see [M-5] `st_updata()`); you must set it if you want it set.

## Also see

[M-4] `stata` — Stata interface functions

## Syntax

```

real rowvector    st_addvar(type, name)
real rowvector    st_addvar(type, name, nofill)
real rowvector    _st_addvar(type, name)
real rowvector    _st_addvar(type, name, nofill)
    
```

where

*type*:    *string scalar* or *rowvector* containing "byte", "int", "long", "float",  
          "double", or "str#"
          or
          *real scalar* or *rowvector* containing # (interpreted as str#)

*name*:    *string rowvector* containing new variable names

*nofill*:   *real scalar* containing 0 or non-0

## Description

st\_addvar(*type*, *name*) adds new variable *name*(s) of type *type* to the Stata dataset. Returned are the variable indices of the new variables. st\_addvar() aborts with error (and adds no variables) if any of the variables already exist or cannot be added for other reasons.

st\_addvar(*type*, *name*, *nofill*) does the same thing. *nofill* ≠ 0 specifies that the variables' values are not to be filled in with missing values. st\_addvar(*type*, *name*, 0) is the same as st\_addvar(*type*, *name*). Use of *nofill* ≠ 0 is not, in general, recommended. See [Using nofill](#) in *Remarks* below.

\_st\_addvar() does the same thing as st\_addvar() except that, rather than aborting with error if the new variable cannot be added, returned is a 1 × 1 scalar containing the negative of the appropriate Stata return code.

## Remarks

Remarks are presented under the following headings:

- [Creating a new variable](#)
- [Creating new variables](#)
- [Creating new string variables](#)
- [Creating a new temporary variable](#)
- [Creating temporary variables](#)
- [Handling errors](#)
- [Using nofill](#)

## Creating a new variable

To create new variable `myvar` as a double, code

```
idx = st_addvar("double", "myvar")
```

or

```
(void) st_addvar("double", "myvar")
```

You use the first form if you will subsequently need the variable's index number, or you use the second form otherwise.

## Creating new variables

You can add more than one variable. For instance,

```
idx = st_addvar("double", ("myvar1", "myvar2"))
```

adds two new variables, both of type double.

```
idx = st_addvar(("double", "float"), ("myvar1", "myvar2"))
```

also adds two new variables, but this time, `myvar1` is double and `myvar2` is float.

## Creating new string variables

Creating string variables is no different from any other type:

```
idx = st_addvar(("str10", "str5"), ("myvar1", "myvar2"))
```

creates `myvar1` as a `str10` and `myvar2` as a `str5`.

There is, however, another way to specify the types.

```
idx = st_addvar((10, 5), ("myvar1", "myvar2"))
```

also creates `myvar1` as a `str10` and `myvar2` as a `str5`.

```
idx = st_addvar(10, ("myvar1", "myvar2"))
```

creates both variables as `str10`s.

## Creating a new temporary variable

Function `st_tempname()` (see [\[M-5\] `st\_tempname\(\)`](#)) returns temporary variable names. To create a temporary variable as a double, code

```
idx = st_addvar("double", st_tempname())
```

or code

```
(void) st_addvar("double", name=st_tempname())
```

You use the first form if you will subsequently need the variable's index, or you use the second form if you will subsequently need the variable's name. You will certainly need one or the other. If you will need both, code

```
idx = st_addvar("double", name=st_tempname())
```

## Creating temporary variables

`st_tempname()` can return a vector of temporary variable names.

```
idx = st_addvar("double", st_tempname(5))
```

creates five temporary variables, each of type double.

## Handling errors

There are three common reasons why `st_addvar()` might fail: the variable name is invalid or a variable under that name already exists or there is insufficient memory to add another variable. If there is a problem adding a variable, `st_addvar()` will abort with error. If you wish to avoid the traceback log and just have Stata issue an error, use `_st_addvar()` and code

```
if ((idx = _st_addvar("double", "myvar")) < 0) exit(error(-idx))
```

If you are adding multiple variables, look at the first element of what `_st_addvar()` returns:

```
if ((idx = _st_addvar(types, names))[1] < 0) exit(error(-idx))
```

## Using *nofill*

The three-argument versions of `st_addvar()` and `_st_addvar()` allow you to avoid filling in the values of the newly created variable. Filling in those values with missing really is a waste of time if the next thing you are going to do is fill in the values with something else. On the other hand, it is important that all the observations be filled in on the new variable before control is returned to Stata, and this includes returning to Stata because of subsequent error or the user pressing *Break*. Thus use of *nofill*  $\neq 0$  is not, in general, recommended. Filling in values really does not take that long.

If you are determined to save the computer time, however, see [M-5] `setbreakintr()`. To do things right, you need to set the break key off, create your variable, fill it in, and turn break-key processing back on.

There is, however, a case in which use of *nofill*  $\neq 0$  is acceptable and such effort is not required: when you are creating a temporary variable. Temporary variables vanish in any case, and it does not matter whether they are filled in before they vanish.

Temporary variables in fact vanish not when Mata ends but when the ado-file calling Mata ends, if there is an ado-file. We will assume there is an ado-file because that is the only case in which you would be creating a temporary variable anyway. Because they do not disappear until later, there is the possibility of there being an issue if the variable is not filled in. If we assume, however, that your Mata program is correctly written and does fill in the variable ultimately, then the chances of a problem are minimal. If the user presses *Break* or there is some other problem in your program that causes Mata to abort, the ado-file will be aborted, too, and the variable will vanish.

Let us add that Stata will not crash if a variable is not filled in, even if it regains control. The danger is that the user will look at the variable or, worse, use it and be baffled by what he or she sees, which might concern not only odd values but also NaNs and worse.

## Conformability

`st_addvar(type, name, nofill):`

*type*:  $1 \times 1$  or  $1 \times k$   
*name*:  $1 \times k$   
*nofill*:  $1 \times 1$  (optional)  
*result*:  $1 \times k$

`_st_addvar(type, name, nofill):`

*type*:  $1 \times 1$  or  $1 \times k$   
*name*:  $1 \times k$   
*nofill*:  $1 \times 1$  (optional)  
*result*:  $1 \times k$  or, if error,  $1 \times 1$

## Diagnostics

`st_addvar(type, name, nofill)` aborts with error if

1. *type* is not equal to a valid Stata variable type and it is not a number that would form a valid `str#` variable type;
2. *name* is not a valid variable name;
3. a variable named *name* already exists;
4. there is insufficient memory to add another variable.

`_st_addvar(type, name, nofill)` aborts with error for reason 1 above, but otherwise, it returns the negative value of the appropriate Stata return code.

Both functions, when creating multiple variables, create either all the variables or none of them. Whether creating one variable or many, if variables are created, `st_updata()` (see [M-5] [st\\_updata\(\)](#)) is set unless all variables are temporary; see [M-5] [st\\_tempname\(\)](#).

## Reference

Gould, W. W. 2006. [Mata Matters: Creating new variables—sounds boring, isn't.](#) *Stata Journal* 6: 112–123.

## Also see

[M-5] [st\\_store\(\)](#) — Modify values stored in current Stata dataset

[M-5] [st\\_tempname\(\)](#) — Temporary Stata names

[M-4] [stata](#) — Stata interface functions

## Syntax

<i>real scalar</i>	<code>_st_data(real scalar i, real scalar j)</code>	
<i>real matrix</i>	<code>st_data(real matrix i, rowvector j)</code>	(1,2)
<i>real matrix</i>	<code>st_data(real matrix i, rowvector j, scalar selectvar)</code>	(1,2,3)
<i>string scalar</i>	<code>_st_sdata(real scalar i, real scalar j)</code>	
<i>string matrix</i>	<code>st_sdata(real matrix i, rowvector j)</code>	(1,2)
<i>string matrix</i>	<code>st_sdata(real matrix i, rowvector j, scalar selectvar)</code>	(1,2,3)

where

1. *i* may be specified as a  $1 \times 1$  scalar, as a  $1 \times 1$  scalar containing missing, as a column vector of observation numbers, as a row vector specifying an observation range, or as a  $k \times 2$  matrix specifying both.

- a. `st_data(1, 2)` returns the first observation on the second variable.
- b. `st_data(., 2)` returns all observations on the second variable.
- c. `st_data((1\2\5), 2)` returns observations 1, 2, and 5 on the second variable.
- d. `st_data((1,5), 2)` returns observations 1 through 5 on the second variable.
- e. `st_data((1,5\7,9), 2)` returns observations 1 through 5 and observations 7 through 9 on the second variable.

When a range is specified, any element of the range ( $i_1, i_2$ ) may be specified to contribute zero observations if  $i_2 = i_1 - 1$ .

2. *j* may be specified as a real row vector or as a string scalar or string row vector.
  - a. `st_data(., .)` returns the values of all variables, all observations of the Stata dataset.
  - b. `st_data(., 1)` returns the value of the first variable, all observations.
  - c. `st_data(., (3,1,9))` returns the values of the third, first, and ninth variables of all observations.
  - d. `st_data(., ("mpg", "weight"))` returns the values of variables `mpg` and `weight`, all observations.
  - e. `st_data(., ("mpg weight"))` does the same as d above.
  - f. `st_data(., ("gnp", "l.gnp"))` returns the values of `gnp` and the lag of `gnp`, all observations.



g. `st_data(., ("gnp l.gnp"))` does the same as `f` above.

h. `st_data(., ("mpg i.rep78"))` returns the value of `mpg` and the 5 pseudovariates associated with `i.rep78`. There are 5 pseudovariates because we are imagining that `auto.dta` is in memory; the actual number is a function of the values taken on by the variable in the sample specified. Factor variables can be specified only with string scalars; specifying `("mpg", "i.rep78")` will not work.

3. *selectvar* may be specified as real or as a string. Observations for which *selectvar*  $\neq$  0 will be selected. If *selectvar* is real, it is interpreted as a variable number. If string, *selectvar* should contain the name of a Stata variable.

Specifying *selectvar* as `"` or as missing `.` has the same result as not specifying *selectvar*; no observations are excluded.

Specifying *selectvar* as 0 means that observations with missing values of the variables specified by *j* are to be excluded.

## Description

`_st_data(i, j)` returns the numeric value of the *i*th observation of the *j*th Stata variable. Observations are numbered 1 through `st_nobs()`. Variables are numbered 1 through `st_nvar()`.

`st_data(i, j)` is similar to `_st_data(i, j)` except

1. *i* may be specified as a vector or matrix to obtain multiple observations simultaneously,
2. *j* may be specified using names or indices (indices are faster), and
3. *j* may be specified to obtain multiple variables simultaneously.

The net effect is that `st_data()` can return a scalar (the value of one variable in one observation), a row vector (the value of many variables in an observation), a column vector (the value of a variable in many observations), or a matrix (the value of many variables in many observations).

`st_data(i, j, selectvar)` works like `st_data(i, j)` except that only observations for which *selectvar*  $\neq$  0 are returned.

`_st_sdata()` and `st_sdata()` are the string variants of `_st_data()` and `st_data()`. `_st_data()` and `st_data()` are for use with numeric variables; they return missing `.` when used with string variables. `_st_sdata()` and `st_sdata()` are for use with string variables; they return empty string `"` when used with numeric variables.

## Remarks

Remarks are presented under the following headings:

*Description of `_st_data()` and `_st_sdata()`*

*Description of `st_data()` and `st_sdata()`*

*Details of observation subscripting using `st_data()` and `st_sdata()`*

Description of `_st_data()` and `_st_sdata()`

`_st_data()` returns one variable’s value in one observation. You refer to variables and observations by their numbers. The first variable in the Stata dataset is 1; the first observation is 1.

<code>_st_data(1, 1)</code>	value of 1st obs., 1st variable
<code>_st_data(1, 2)</code>	value of 1st obs., 2nd variable
<code>_st_data(2, 1)</code>	value of 2nd obs., 1st variable

`_st_sdata()` works the same way. `_st_data()` is for use with numeric variables, and `_st_sdata()` is for use with string variables.

`_st_data()` and `_st_sdata()` are the fastest way to obtain the value of a variable in one observation.

Description of `st_data()` and `st_sdata()`

`st_data()` can be used just like `_st_data()`, and used that way, it produces the same result.

Variables, however, can be referred to by their names or their numbers:

<code>st_data(1, 1)</code>	value of 1st obs., 1st variable
<code>st_data(1, 2)</code>	value of 1st obs., 2nd variable
<code>st_data(2, 1)</code>	value of 2nd obs., 1st variable
<code>st_data(1, "mpg")</code>	value of 1st obs, variable mpg
<code>st_data(2, "mpg")</code>	value of 2nd obs, variable mpg

Also, you may specify more than one variable:

<code>st_data(2, (1,2,3))</code>	value of 2nd obs., variables 1, 2, and 3
<code>st_data(2, ("mpg","weight","displ"))</code>	value of 2nd obs., variables mpg, weight, and displ
<code>st_data(2, "mpg weight displ")</code>	(same as previous)

Used this way, `st_data()` returns a row vector.

Similarly, you may obtain multiple observations:

<code>st_data((1\2\3), 10)</code>	values of obs. 1, 2, and 3, variable 10
<code>st_data((1,5), 10)</code>	values of obs. 1 through 5, variable 10
<code>st_data((1,5)\(7,9), 10)</code>	values of obs. 1 through 5 and 7 through 9, variable 10

`st_sdata()` works the same way as `st_data()`.

## Details of observation subscripting using `st_data()` and `st_sdata()`

1.  $i$  may be specified as a scalar: the specified, single observation is returned.  $i$  must be between 1 and `st_nobs()`; see [M-5] `st_nvar()`.
2.  $i$  may be specified as a scalar containing missing value: all observations are returned.
3.  $i$  may be specified as a column vector: the specified observations are returned. Each element of  $i$  must be between 1 and `st_nobs()` or may be missing. Missing is interpreted as `st_nobs()`.
4.  $i$  may be specified as a  $1 \times 2$  row vector: the specified range of observations is returned;  $(c_1, c_2)$  returns the  $c_2 - c_1 + 1$  observations  $c_1$  through  $c_2$ .  
  
 $c_2 - c_1 + 1$  must evaluate to a number greater than or equal to 0. In general,  $c_1$  and  $c_2$  must be between 1 and `st_nobs()`, but if  $c_2 - c_1 + 1 = 0$ , then  $c_1$  may be between 1 and `st_nobs()` + 1 and  $c_2$  may be between 0 and `st_nobs()`. Regardless,  $c_1 == .$  or  $c_2 == .$  is interpreted as `st_nobs()`.
5.  $i$  may be specified as a  $k \times 2$  matrix:  $((1,5) \backslash (7,7) \backslash (20,30))$  specifies observations 1 through 5, 7, and 20 through 30.

## Conformability

`_st_data(i, j)`, `_st_sdata(i, j)`:

$i$ :  $1 \times 1$   
 $j$ :  $1 \times 1$   
 result:  $1 \times 1$

`st_data(i, j)`, `st_sdata(i, j)`:

$i$ :  $n \times 1$  or  $n_2 \times 2$   
 $j$ :  $1 \times k$  or  $1 \times 1$  containing  $k$  elements when expanded  
 result:  $n \times k$

`st_data(i, j, selectvar)`, `st_sdata(i, j, selectvar)`:

$i$ :  $n \times 1$  or  $n_2 \times 2$   
 $j$ :  $1 \times k$  or  $1 \times 1$  containing  $k$  elements when expanded  
 $selectvar$ :  $1 \times 1$   
 result:  $(n - e) \times k$ , where  $e$  is number of observations excluded by  $selectvar$

## Diagnostics

`_st_data(i, j)` returns missing (.) if  $i$  or  $j$  is out of range; it does not abort with error.

`_st_sdata(i, j)` returns "" if  $i$  or  $j$  is out of range; it does not abort with error.

`st_data(i, j)` and `st_sdata(i, j)` abort with error if any element of  $i$  or  $j$  is out of range.  $j$  may be specified as variable names or variable indices. If names are specified, abbreviations are allowed. If you do not want this and no factor variables nor time-series-operated variables are specified, use `st_varindex()` (see [M-5] `st_varindex()`) to translate variable names into variable indices.

## Also see

[M-5] `st_view()` — Make matrix that is a view onto current Stata dataset

[M-5] `st_store()` — Modify values stored in current Stata dataset

[M-4] `stata` — Stata interface functions

[D] `putmata` — Put Stata variables into Mata and vice versa

Syntax

```
string colvector st_dir(cat, subcat, pattern)

string colvector st_dir(cat, subcat, pattern, adorn)
```

- where
- cat*: string scalar containing "local", "global", "r()", "e()", "s()", or "char"
  - subcat*: string scalar containing "macro", "numscalar", "strscalar", "matrix", or, if *cat*=="char", "\_dta" or a name.
  - pattern*: string scalar containing a pattern as defined in [M-5] [strmatch\(\)](#)
  - adorn*: string scalar containing 0 or non-0

The valid *cat*–*subcat* combinations and their meanings are

<i>cat</i>	<i>subcat</i>	Meaning
"local"	"macro"	Stata's local macros
"global"	"macro"	Stata's global macros
"global"	"numscalar"	Stata's numeric scalars
"global"	"strscalar"	Stata's string scalars
"global"	"matrix"	Stata's matrices
"r()"	"macro"	macros in r()
"r()"	"numscalar"	numeric scalars in r()
"r()"	"matrix"	matrices in r()
"e()"	"macro"	macros in e()
"e()"	"numscalar"	numeric scalars in e()
"e()"	"matrix"	matrices in e()
"s()"	"macro"	macros in s()
"char"	"_dta"	characteristics in _dta[]
"char"	"name"	characteristics in variable <i>name</i> []

st\_dir() returns an empty list if an invalid *cat*–*subcat* combination is specified.

## Description

`st_dir(cat, subcat, pattern)` and `st_dir(cat, subcat, pattern, adorn)` return a column vector containing the names matching *pattern* of the Stata objects described by *cat*–*subcat*.

Argument *adorn* is optional; not specifying it is equivalent to specifying *adorn* = 0. By default, simple names are returned. If *adorn* ≠ 0 is specified, the name is adorned in the standard Stata way used to describe the object. Say that one is listing the macros in `e()` and one of the elements is `e(cmd)`. By default, the returned vector will contain an element equal to "cmd". With *adorn* ≠ 0, the element will be "e(cmd)".

For many objects, the adorned and unadorned forms of the names are the same.

## Conformability

```
st_dir(cat, subcat, pattern, adorn):
      cat:      1 × 1
      subcat:   1 × 1
      pattern:  1 × 1
      adorn:    1 × 1   (optional)
      result:   k × 1
```

## Diagnostics

`st_dir(cat, subcat, pattern)` and `st_dir(cat, subcat, pattern, adorn)` abort with error if *cat* or *subcat* is invalid. If the combination is invalid, however, `J(0,1,"")` is returned. *subcat*==*name* is considered invalid unless *cat*=="char".

`st_dir()` aborts with error if any of its arguments are views.

## Also see

[M-4] [stata](#) — Stata interface functions

## Syntax

```
void st_dropvar(transmorphic rowvector vars)
```

```
void st_dropobsin(real matrix range)
```

```
void st_dropobsif(real colvector select)
```

```
void st_keepvar(transmorphic rowvector vars)
```

```
void st_keepobsin(real matrix range)
```

```
void st_keepobsif(real colvector select)
```

## Description

`st_dropvar(vars)` drops the variables specified. *vars* is a row vector that may contain either variable names or variable indices. `st_dropvar(.)` drops all variables and observations.

`st_dropobsin()` and `st_dropobsif()` have to do with dropping observations.

`st_dropobsin(range)` specifies the observations to be dropped:

`st_dropobsin(5)` drops observation 5.

`st_dropobsin((5,9))` drops observations 5 through 9.

`st_dropobsin((5\8\12))` drops observations 5 and 8 and 12.

`st_dropobsin((5,7\8,11\13,13))` drops observations 5 through 7, 8 through 11, and 13.

`st_dropobsin(.)` drops all observations (but not the variables).

`st_dropobsin(J(0,1,.))` drops no observations (or variables).

`st_dropobsif(select)` specifies a `st_nobs()`  $\times$  1 vector. Observations *i* for which *select*<sub>*i*</sub>  $\neq$  0 are dropped.

`st_keepvar()`, `st_keepobsin()`, and `st_keepobsif()` do the same thing, except that the variables and observations to be kept are specified.

## Remarks

To drop all variables and observations, code any of the following:

```
st_dropvar(.)
st_keepvar(J(1,0,.))
st_keepvar(J(1,0,""))
```

All do the same thing. Dropping all the variables clears the dataset.

Dropping all the observations, however, leaves the variables in place.

## Conformability

`st_dropvar(vars)`, `st_keepvar(vars)`:

*vars*:  $1 \times k$   
*result*: *void*

`st_dropobsin(range)`, `st_keepobsin(range)`:

*range*:  $k \times 1$  or  $k \times 2$   
*result*: *void*

`st_dropobsif(select)`, `st_keepobsif(select)`:

*select*: `st_nobs()`  $\times$  1  
*result*: *void*

## Diagnostics

`st_dropvar(vars)` and `st_keepvar(vars)` abort with error if any element of *vars* is missing unless *vars* is  $1 \times 1$ , in which case they drop or keep all the variables.

`st_dropvar(vars)` and `st_keepvar(vars)` abort with error if any element of *vars* is not a valid variable index or name, or if *vars* is a view. If *vars* is specified as names, abbreviations are not allowed.

`st_dropvar()` and `st_keepvar()` set `st_updata()` (see [M-5] `st_updata()`) unless all variables dropped are temporary; see [M-5] `st_tempname()`.

`st_dropobsin(range)` and `st_keepobsin(range)` abort with error if any element of *range* is missing unless *range* is  $1 \times 1$ , in which case they drop or keep all the observations.

`st_dropobsin(range)` and `st_keepobsin(range)` abort with error if any element of *range* is not a valid observation number (is not between 1 and `st_nobs()` [see [M-5] `st_nvar()`] inclusive) or if *range* is a view.

`st_dropobsif(select)` and `st_keepobsif(select)` abort with error if *select* is a view.

`st_dropobsin()`, `st_dropobsif()`, `st_keepobsin()`, and `st_keepobsif()` set `st_updata()` if any observations are removed from the data.

Be aware that, after dropping any variables or observations, any previously constructed views (see [M-5] `st_view()`) are probably invalid because views are internally stored in terms of variable and observation numbers. Subsequent use of an invalid view may lead to unexpected results or an abort with error.

## Also see

[M-4] `stata` — Stata interface functions



## Syntax

```

string scalar  st_global(string scalar name)

void           st_global(string scalar name, string scalar contents)

void           st_global(string scalar name, string scalar contents, string scalar hcat)

string scalar  st_global_hcat(string scalar name)
    
```

where

1. *name* is to contain
  - a. global macro such as "myname"
  - b. `r()` macro such as "r(names)"
  - c. `e()` macro such as "e(cmd)"
  - d. `s()` macro such as "s(vars)"
  - e. `c()` macro such as "c(current\_date)"
  - f. dataset characteristic such as "\_dta[date]"
  - g. variable characteristic such as "mpg[note]"
2. `st_global(name)` returns the contents of the specified Stata global. It returns "" when the global does not exist.
3. `st_global(name, contents)` sets or resets the contents of the specified Stata global.
4. `st_global(name, "")` deletes the specified Stata global. It does this even if *name* is not a macro. `st_global("r(N)", "")` would delete `r(N)` whether it were a macro, scalar, or matrix.
5. `st_global(name, contents, hcat)` sets or resets the contents of the specified Stata global, and it sets or resets the hidden or historical status when *name* is an `e()` or `r()` value. Allowed *hcat* values are "visible", "hidden", "historical", and a string scalar release number such as "10", "10.1", or any string release number matching "#[#][. [#]]". See [\[P\] return](#) for a description of hidden and historical `r()` and `e()` values.  
  
 When `st_global(name, contents)` is used to set an `e()` or `r()` value, its *hcat* is set to "visible".
6. `st_global_hcat(name)` returns the *hcat* associated with an `e()` or `r()` value.

## Description

`st_global(name)` returns the contents of the specified Stata global.

`st_global(name, contents)` sets or resets the contents of the specified Stata global. If the Stata global did not previously exist, a new global is created. If the global did exist, the new contents replace the old.

`st_global(name, contents, hcat)` and `st_global_hcat(name)` are used to set and query the *hcat* corresponding to an `e()` or `r()` value. They are also rarely used. See [\[R\] saved results](#) and [\[P\] return](#) for more information.

## Remarks

Mata provides a suite of functions for obtaining and setting the contents of global macros, local macros, saved results, etc. It can sometimes be confusing to know which you should use. The table on the following page will help.

Stata component/action	function call
Local macro	
obtain contents	<code>contents = st_local("name")</code>
create/set/replace	<code>st_local("name", contents)</code>
delete	<code>st_local("name", "")</code>
Global macro	
obtain contents	<code>contents = st_global("name")</code>
create/set/replace	<code>st_global("name", contents)</code>
delete	<code>st_global("name", "")</code>
Global numeric scalar	
obtain contents	<code>value = st_numscalar("name")</code>
create/set/replace	<code>st_numscalar("name", value)</code>
delete	<code>st_numscalar("name", J(0,0,.))</code>
Global string scalar	
obtain contents	<code>contents = st_strscalar("name")</code>
create/set/replace	<code>st_strscalar("name", contents)</code>
delete	<code>st_strscalar("name", J(0,0,""))</code>
Global matrix	
obtain contents	<code>matrix = st_matrix("name")</code> <code>rowlabel = st_matrixrowstripe("name")</code> <code>collabel = st_matrixcolstripe("name")</code>
create/set/replace	<code>st_matrix("name", matrix)</code> <code>st_matrixrowstripe("name", rowlabel)</code> <code>st_matrixcolstripe("name", collabel)</code>
replace	<code>st_replacematrix("name", matrix)</code>
delete	<code>st_matrix("name", J(0,0,.))</code>
Characteristic	
obtain contents	<code>contents = st_global("name[name]")</code>
create/set/replace	<code>st_global("name[name]", contents)</code>
delete	<code>st_global("name[name]", "")</code>

Stata component/action	function call
<code>r()</code> results	
macro	
obtain contents	<code>contents = st_global("r(name)")</code>
create/set/replace	<code>st_global("r(name)", contents)</code>
numeric scalar	
obtain contents	<code>value = st_numscalar("r(name)")</code>
create/set/replace	<code>st_numscalar("r(name)", value)</code>
matrix	
obtain contents	<code>matrix = st_matrix("r(name)")</code> <code>rowlabel = st_matrixrowstripe("r(name)")</code> <code>collabel = st_matrixcolstripe("r(name)")</code>
create/set/replace	<code>st_matrix("r(name)", matrix)</code> <code>st_matrixrowstripe("r(name)", rowlabel)</code> <code>st_matrixcolstripe("r(name)", collabel)</code>
replace	<code>st_replacematrix("r(name)", matrix)</code>
IN ALL CASES	
delete	<code>st_global("r(name)", "")</code>
to delete all of <code>r()</code>	<code>st_rclear()</code>
<code>e()</code> results	same as <code>r()</code> results, but code <code>e(name)</code> and <code>st_eclear()</code>
<code>s()</code> results	
macro	
obtain contents	<code>contents = st_global("s(name)")</code>
create/set/replace	<code>st_global("s(name)", contents)</code>
delete	<code>st_global("s(name)", "")</code>
to delete all of <code>s()</code>	<code>st_sclear()</code>
<code>c()</code> results	
macro	
obtain contents	<code>contents = st_global("c(name)")</code>
numeric scalar	
obtain contents	<code>value = st_numscalar("c(name)")</code>

See [M-5] [st\\_local\(\)](#), [M-5] [st\\_numscalar\(\)](#), [M-5] [st\\_matrix\(\)](#), and [M-5] [st\\_rclear\(\)](#).

## Conformability

```

st_global(name):
    name:      1 × 1
    result:    1 × 1

st_global(name, contents):
    name:      1 × 1
    contents:  1 × 1
    result:    void

st_global(name, contents, hcat):
    name:      1 × 1
    contents:  1 × 1
    hcat:      1 × 1
    result:    void

st_global_hcat(name):
    name:      1 × 1
    result:    1 × 1

```

## Diagnostics

`st_global(name)` returns "" if the name contained in *name* is not defined. `st_global(name)` aborts with error if the name is malformed, such as `st_global("invalid name")`.

`st_global(name, contents)` aborts with error if the name contained in *name* is malformed. The maximum length of strings in Mata is significantly longer than in Stata. `st_global()` truncates what is stored at the appropriate maximum length if that is necessary.

`st_global_hcat(name)` returns "visible" when *name* is not an `e()` or `r()` value and returns "" when *name* is an `e()` or `r()` value that does not exist.

## Reference

Gould, W. W. 2008. [Mata Matters: Macros](#). *Stata Journal* 8: 401–412.

## Also see

[M-5] [st\\_rclear\(\)](#) — Clear `r()`, `e()`, or `s()`

[M-4] [stata](#) — Stata interface functions

# Title

[M-5] **st\_isfmt()** — Whether valid %fmt

## Syntax

```
real scalar  st_isfmt(string scalar s)  
real scalar  st_isnumfmt(string scalar s)  
real scalar  st_isstrfmt(string scalar s)
```

## Description

`st_isfmt(s)` returns 1 if *s* contains a valid Stata *%fmt* and 0 otherwise.  
`st_isnumfmt(s)` returns 1 if *s* contains a valid Stata numeric *%fmt* and 0 otherwise.  
`st_isstrfmt(s)` returns 1 if *s* contains a valid Stata string *%fmt* and 0 otherwise.

## Conformability

```
st_isfmt(s), st_isnumfmt(s), st_isstrfmt(s):  
  s:      1 × 1  
  result: 1 × 1
```

## Diagnostics

`st_isfmt(s)`, `st_isnumfmt(s)`, and `st_isstrfmt(s)` abort with error if *s* is a view.

## Also see

[M-4] **stata** — Stata interface functions

# Title

[M-5] `st_isname()` — Whether valid Stata name

## Syntax

*real scalar*    `st_isname(string scalar s)`  
*real scalar*    `st_islmmname(string scalar s)`

## Description

`st_isname(s)` returns 1 if *s* contains a valid Stata name and 0 otherwise.  
`st_islmmname(s)` returns 1 if *s* contains a valid Stata local-macro name and 0 otherwise.

## Conformability

`st_isname(s), st_islmmname(s):`  
      *s:*         $1 \times 1$   
      *result:*     $1 \times 1$

## Diagnostics

`st_isname(s)` aborts with error if *s* is a view (but `st_islmmname()` does not).

## Also see

[\[M-4\] `stata`](#) — Stata interface functions

# Title

[M-5] **st\_local()** — Obtain strings from and put strings into Stata macros

## Syntax

```
string scalar  st_local(string scalar name)  
void          st_local(string scalar name, string scalar contents)
```

Note: `st_local(name, "")` deletes.

## Description

`st_local(name)` returns the contents of the specified local macro.

`st_local(name, contents)` sets or resets the contents of the specified local macro. If the macro did not previously exist, a new macro is created. If it did previously exist, the new contents replace the old.

## Remarks

See [\[M-5\] st\\_global\(\)](#) and [\[M-5\] st\\_rclear\(\)](#).

## Conformability

```
st_local(name):  
  name:      1 × 1  
  result:    1 × 1  
  
st_local(name, contents):  
  name:      1 × 1  
  contents:  1 × 1  
  result:    void
```

## Diagnostics

`st_local(name)` returns "" if the name contained in *name* is not defined. `st_local(name)` aborts with error if the name is malformed.

`st_local(name, contents)` aborts with error if the name contained in *name* is malformed.

## Reference

Gould, W. W. 2008. [Mata Matters: Macros](#). *Stata Journal* 8: 401–412.



## Also see

[M-4] [stata](#) — Stata interface functions

# Title

[M-5] `st_macroexpand()` — Expand Stata macros in string

## Syntax

```
string scalar    st_macroexpand(string scalar s)  
real scalar     _st_macroexpand(S, string scalar s)
```

Note: the type of *S* does not matter; it is replaced and becomes a string scalar.

## Description

`st_macroexpand(s)` returns *s* with any quoted or dollar sign–prefixed macros expanded.

`_st_macroexpand(S, s)` places in *S* the contents of *s* with any quoted or dollar sign–prefixed macros expanded and returns a Stata return code (it returns 0 if all went well).

## Remarks

Be careful coding string literals containing quoted or prefixed macros because macros are also expanded at compile time. For instance, consider

```
s = st_macroexpand("regress 'varlist'")
```

'varlist' will be substituted with its value at compile time. What you probably want is

```
s = st_macroexpand("regress " + "'" + "varlist" + "'")
```

## Conformability

`st_macroexpand(s)`:

<i>s</i> :	1 × 1
<i>result</i> :	1 × 1

`_st_macroexpand(S, s)`:

*input*:

<i>s</i> :	1 × 1
------------	-------

*output*:

<i>S</i> :	1 × 1
<i>result</i> :	1 × 1

## Diagnostics

`st_macroexpand(s)` aborts with error if *s* is too long (exceedingly unlikely) or if macro expansion fails (also unlikely).

`_st_macroexpand(S, s)` aborts with error if *s* is too long.

## Also see

[M-4] [stata](#) — Stata interface functions

## Syntax

```

real matrix    st_matrix(string scalar name)
string matrix st_matrixrowstripe(string scalar name)
string matrix st_matrixcolstripe(string scalar name)

void           st_matrix(string scalar name, real matrix X)
void           st_matrix(string scalar name, real matrix X, string scalar hcat)
void           st_matrixrowstripe(string scalar name, string matrix s)
void           st_matrixcolstripe(string scalar name, string matrix s)

void           st_replacematrix(string scalar name, real matrix X)

string scalar st_matrix_hcat(name)

```

where

1. All functions allow *name* to be
  - a. global matrix name such as "mymatrix",
  - b. `r()` matrix such as "r(Z)", or
  - c. `e()` matrix such as "e(V)".
2. `st_matrix(name)` returns the contents of the specified Stata matrix. It returns `J(0,0,.)` if the matrix does not exist.
3. `st_matrix(name, X)` sets or resets the contents of the specified Stata matrix. Row and column stripes are set to the default `r1, r2, ...`, and `c1, c2, ...`.
4. `st_replacematrix(name, X)` is an alternative way to replace existing Stata matrices. The number of rows and columns of *X* must match the Stata matrix being replaced, and in return, the row and column stripes are not replaced.
5. `st_matrix(name, X)` deletes the specified Stata matrix if *value*==`J(0,0,.)` (if *value* is  $0 \times 0$ ).
6. Neither `st_matrix()` nor `st_replacematrix()` can be used to set, replace, or delete special Stata `e()` matrices `e(b)`, `e(V)`, or `e(Cns)`. Only Stata commands `ereturn post` and `ereturn repost` can be used to set these special matrices; see [P] [ereturn](#). Also see [M-5] [stata\(\)](#) for executing Stata commands from Mata.
7. `st_matrix(name, X, hcat)` sets or resets the specified Stata matrix and sets the hidden or historical status when setting a Stata `e()` or `r()` matrix. Allowed *hcat* values are "visible", "hidden", "historical", and a string scalar release number such as "10", "10.1", or

any string release number matching `"#[#][.#[#]]"`. See [\[P\] return](#) for a description of hidden and historical saved results.

8. `st_matrix_hcat(name)` returns the *hcat* associated with a Stata `e()` or `r()` matrix.
9. `st_matrixrowstripe()` and `st_matrixcolstripe()` allow querying and resetting the row and column stripes of existing or previously created Stata matrices.

## Description

`st_matrix(name)` returns the contents of Stata's matrix *name*, or it returns `J(0,0,.)` if the matrix does not exist.

`st_matrixrowstripe(name)` returns the row stripe associated with the matrix *name*, or it returns `J(0,2,"")` if the matrix does not exist.

`st_matrixcolstripe(name)` returns the column stripe associated with the matrix *name*, or it returns `J(0,2,"")` if the matrix does not exist.

`st_matrix(name, X)` sets or resets the contents of the Stata matrix *name* to be *X*. If the matrix did not previously exist, a new matrix is created. If the matrix did exist, the new contents replace the old. Either way, the row and column stripes are also reset to contain `"r1"`, `"r2"`, ..., and `"c1"`, `"c2"`, ....

`st_matrix(name, X)` deletes the Stata matrix *name* when *X* is  $0 \times 0$ : `st_matrix(name, J(0,0,.))` deletes Stata matrix *name* or does nothing if *name* does not exist.

`st_matrixrowstripe(name, s)` and `st_matrixcolstripe(name, s)` change the contents to be *s* of the row and column stripe associated with the already existing Stata matrix *name*. In either case, *s* must be  $n \times 2$ , where *n* = the number of rows (columns) of the underlying matrix.

`st_matrixrowstripe(name, s)` and `st_matrixcolstripe(name, s)` reset the row and column stripe to be `"r1"`, `"r2"`, ..., and `"c1"`, `"c2"`, ..., when *s* is  $0 \times 2$  (that is, `J(0,2,"")`).

`st_replacematrix(name, X)` resets the contents of the Stata matrix *name* to be *X*. The existing Stata matrix must have the same number of rows and columns as *X*. The row stripes and column stripes remain unchanged.

`st_matrix(name, X, hcat)` and `st_matrix_hcat(name)` are used to set and query the *hcat* corresponding to a Stata `e()` or `r()` matrix. They are also rarely used. See [\[R\] saved results](#) and [\[P\] return](#) for more information.

## Remarks

Remarks are presented under the following headings:

*Processing Stata's row and column stripes*  
*Stata's matsize is irrelevant*

Also see [\[M-5\] st\\_global\(\)](#) and [\[M-5\] st\\_rclear\(\)](#).

## Processing Stata's row and column stripes

Both row stripes and column stripes are presented in the same way: each row of *s* represents the *eq:op.name* associated with a row or column of the underlying matrix. The first column records *eq*, and the second column records *op.name*. For instance, given the following Stata matrix

		eq2:	eq2:
	L.	L.	L.
	turn	turn	turn
mpg	1	2	3
L.mpg	5	6	7
eq2:mpg	9	10	11
eq2:L.mpg	13	14	15

`st_matrixrowstripe(name)` returns the  $4 \times 2$  string matrix

" "	"mpg"
" "	"L.mpg"
"eq2"	"mpg"
"eq2"	"L.mpg"

and `st_matrixcolstripe(name)` returns

" "	"turn"
" "	"L.turn"
"eq2"	"turn"
"eq2"	"L.turn"

## Stata's matsize is irrelevant

Matrices in Stata are limited to `matsize` (see [\[R\] matsize](#)), a number between 10 and 11,000. Mata matrices have no such limits.

When getting a matrix, the `matsize` limit plays no role.

When putting a matrix, the `matsize` limit is ignored; meaning that, to use the matrix in Stata, the user may have to reset `matsize` or, if the matrix is too large, the user may not be able to use the matrix at all.

## Conformability

```
st_matrix(name):
  name:      1 × 1
  result:    m × n  (0 × 0 if not found)
```

```
st_matrixrowstripe(name):
  name:      1 × 1
  result:    m × 2  (0 × 2 if not found)
```

```
st_matrixcolstripe(name):
  name:      1 × 1
  result:    n × 2  (0 × 2 if not found)
```

```

st_matrix(name, X):
    name:    1 × 1
    X:       r × c  (0 × 0 means delete)
    result:   void

st_matrix(name, X, hcat):
    name:    1 × 1
    X:       r × c
    hcat:    1 × 1
    result:   void

st_matrixrowstripe(name, s):
    name:    1 × 1
    s:       r × 2  (0 × 2 means default "r1", "r2", ...)
    result:   void

st_matrixcolstripe(name, s):
    name:    1 × 1
    s:       c × 2  (0 × 2 means default "c1", "c2", ...)
    result:   void

st_replacematrix(name, X):
    name:    1 × 1
    X:       m × n  (0 × 0 means delete)
    result:   void

st_matrix_hcat(name):
    name:    1 × 1
    result:   1 × 1

```

## Diagnostics

`st_matrix(name)`, `st_matrixrowstripe(name)`, and `st_matrixcolstripe(name)` abort with error if *name* is malformed. Also,

1. `st_matrix(name)` returns `J(0,0,.)` if Stata matrix *name* does not exist.
2. `st_matrixrowstripe(name)` and `st_matrixcolstripe(name)` return `J(0,2,"")` if Stata matrix *name* does not exist. There is no possibility that matrix *name* might exist and not have row and column stripes.

`st_matrix(name, X)`, `st_matrixrowstripe(name, s)`, and `st_matrixcolstripe(name, s)` abort with error if *name* is malformed. Also,

1. `st_matrixrowstripe(name, s)` aborts with error if `rows(s)` is not equal to the number of rows of Stata matrix *name* and `rows(s) != 0`, or if `cols(s) != 2`.
2. `st_matrixcolstripe(name, s)` aborts with error if `cols(s)` is not equal to the number of columns of Stata matrix *name* and `cols(s) != 0`, or if `cols(s) != 2`.

`st_replacematrix(name, X)` aborts with error if Stata matrix *name* does not have the same number of rows and columns as *X*. `st_replacematrix()` also aborts with error if Stata matrix *name* does not exist and `X != J(0,0,.)`; `st_replacematrix()` does nothing if the matrix does not exist and `X = J(0,0,.)`. `st_replacematrix()` aborts with error if *name* is malformed.

`st_matrix(name, X, hcat)` aborts with error if *hcat* is not an allowed value.

`st_matrix_hcat(name)` returns "visible" when *name* is not a Stata `e()` or `r()` matrix and returns "" when *name* is an `e()` or `r()` value that does not exist.

## Also see

[M-5] [st\\_rclear\(\)](#) — Clear `r()`, `e()`, or `s()`

[M-4] [stata](#) — Stata interface functions

**[M-5] st\_numscalar()** — Obtain values from and put values into Stata scalars

## Syntax

```

real    st_numscalar(string scalar name)

void    st_numscalar(string scalar name, real value)

void    st_numscalar(string scalar name, real value, string scalar hcat)

string  st_numscalar_hcat(string scalar name)

string  st_strscalar(string scalar name)

void    st_strscalar(string scalar name, string value)

```

where

1. Functions allow *name* to be
  - a. global scalar such as "myname",
  - b. `r()` scalar such as "r(mean)",
  - c. `e()` scalar such as "e(N)", or
  - d. `c()` scalar such as "c(namelen)".

Note that string scalars never appear in `r()` and `e()`; thus (b) and (c) do not apply to `st_strscalar()`.

2. `st_numscalar(name)` and `st_strscalar(name)` return the value of the specified Stata scalar. They return a  $1 \times 1$  result if the specified Stata scalar exists and return a  $0 \times 0$  result otherwise.
3. `st_numscalar(name, value)` and `st_strscalar(name, value)` set or reset the contents of the specified Stata scalar.
4. `st_numscalar(name, value)` and `st_strscalar(name, value)` delete the specified Stata scalar if `value==J(0,0,.)` (if `value` is  $0 \times 0$ ).
5. `st_numscalar(name, value, hcat)` sets or resets the specified Stata scalar and sets or resets the hidden or historical status when *name* is an `e()` or `r()` value. Allowed *hcat* values are "visible", "hidden", "historical", and a string scalar release number such as "10", "10.1", or any string release number matching "`#`[`#`][`.`[`#`[`#`]]". See [\[P\] return](#) for a description of hidden and historical saved results.

When `st_numscalar(name, value)` is used to set an `e()` or `r()` value, its *hcat* is set to "visible".

There is no three-argument form of `st_strscalar()` because there are no `r()` or `e()` string scalar values.



## Description

`st_numscalar(name)` returns the value of the specified Stata numeric scalar, or it returns `J(0,0,.)` if the scalar does not exist.

`st_numscalar(name, value)` sets or resets the value of the specified numeric scalar, assuming `value != J(0,0,.)`. `st_numscalar(name, value)` deletes the specified scalar if `value == J(0,0,.)`. `st_numscalar("x", J(0,0,.))` erases the scalar `x`, or it does nothing if scalar `x` did not exist.

`st_strscalar(name)` returns the value of the specified Stata string scalar, or it returns `J(0,0,"")` if the scalar does not exist.

`st_strscalar(name, value)` sets or resets the value of the specified scalar, assuming `value != J(0,0,"")`. `st_strscalar(name, value)` deletes the specified scalar if `value == J(0,0,"")`. `st_strscalar("x", J(0,0,""))` erases the scalar `x`, or it does nothing if scalar `x` did not exist.

Concerning deletion of a scalar, it does not matter whether you code `st_numscalar(name, J(0,0,.))` or `st_strscalar(name, J(0,0,""))`; both yield the same result.

`st_numscalar(name, value, hcat)` and `st_numscalar_hcat(name)` are used to set and query the `hcat` corresponding to an `e()` or `r()` value. They are also rarely used. See [\[R\] saved results](#) and [\[P\] return](#) for more information.

## Remarks

See [\[M-5\] st\\_global\(\)](#) and [\[M-5\] st\\_rclear\(\)](#).

## Conformability

`st_numscalar(name), st_strscalar(name):`

<i>name:</i>	$1 \times 1$
<i>result:</i>	$1 \times 1$ or $0 \times 0$

`st_numscalar(name, value), st_strscalar(name, value):`

<i>name:</i>	$1 \times 1$
<i>value:</i>	$1 \times 1$ or $0 \times 0$
<i>result:</i>	<i>void</i>

`st_numscalar(name, value, hcat):`

<i>name:</i>	$1 \times 1$
<i>value:</i>	$1 \times 1$
<i>hcat:</i>	$1 \times 1$
<i>result:</i>	<i>void</i>

`st_numscalar(name):`

<i>name:</i>	$1 \times 1$
<i>result:</i>	$1 \times 1$

## Diagnostics

All functions abort with error if *name* is malformed.

`st_numscalar(name)` and `st_strscalar(name)` return `J(0,0,.)` or `J(0,0,"")` if Stata scalar *name* does not exist. They abort with error, however, if the name is malformed.

`st_numscalar(name, value, hcat)` aborts with error if *hcat* is not an allowed value.

`st_numscalar_hcat(name)` returns "visible" when *name* is not an `e()` or `r()` value and returns "" when *name* is an `e()` or `r()` value that does not exist.

## Also see

[M-5] [st\\_rclear\(\)](#) — Clear `r()`, `e()`, or `s()`

[M-4] [stata](#) — Stata interface functions

# Title

[M-5] **st\_nvar()** — Numbers of variables and observations

## Syntax

*real scalar*    `st_nvar()`

*real scalar*    `st_nobs()`

## Description

`st_nvar()` returns the number of variables defined in the dataset currently loaded in Stata.

`st_nobs()` returns the number of observations defined in the dataset currently loaded in Stata.

## Conformability

`st_nvar()`, `st_nobs()`:  
*result:*         $1 \times 1$

## Diagnostics

None.

## Also see

[M-4] **stata** — Stata interface functions

# Title

**[M-5] st\_rclear()** — Clear `r()`, `e()`, or `s()`

## Syntax

```
void st_rclear()
```

```
void st_eclear()
```

```
void st_sclear()
```

## Description

`st_rclear()` clears Stata's `r()` saved results.

`st_eclear()` clears Stata's `e()` saved results.

`st_sclear()` clears Stata's `s()` saved results.

## Remarks

Returning results in `r()`, `e()`, or `s()` is one way of communicating results calculated in Mata back to Stata; see [\[M-1\] ado](#). See [\[R\] saved results](#) for a description of `e()`, `r()`, and `s()`.

Use `st_rclear()`, `st_eclear()`, or `st_sclear()` to clear results, and then use `st_global()` to define macros, `st_numscalar()` to define scalars, and `st_matrix()` to define Stata matrices in `r()`, `e()`, or `s()`. For example,

```
st_rclear()
st_global("r(name)", "tab")      see \[M-5\] st\_global\(\)
st_numscalar("r(N)", n1+n2)      see \[M-5\] st\_numscalar\(\)
st_matrix("r(table)", X+Y)       see \[M-5\] st\_matrix\(\)
```

It is not necessary to clear before saving, but it is considered good style unless it is your intention to add to previously saved results.

If a saved result already exists, `st_global()`, `st_numscalar()`, and `st_matrix()` may be used to redefine it and even to redefine it to a different type. For instance, continuing with our example, later in the same code might appear

```
if (...) {
    st_matrix("r(name)", X)
}
```

Saved result `r(name)` was previously defined as a macro containing `"tab"`, and, even so, can now be redefined to become a matrix.

If you want to eliminate a particular saved result, use `st_global()` to change its contents to `""`:

```
st_global("r(name)", "")
```

Do this regardless of the type of the saved result. Here we use `st_global()` to clear saved result `r(name)`, which might be a macro and might be a matrix.

## Conformability

`st_rclear()`, `st_eclear()`, and `st_sclear()` take no arguments and return void.

## Diagnostics

`st_rclear()`, `st_eclear()`, and `st_sclear()` cannot fail.

## Also see

[M-5] [st\\_global\(\)](#) — Obtain strings from and put strings into global macros

[M-5] [st\\_numscalar\(\)](#) — Obtain values from and put values into Stata scalars

[M-5] [st\\_matrix\(\)](#) — Obtain and put Stata matrices

[M-4] [stata](#) — Stata interface functions

**[M-5] st\_store()** — Modify values stored in current Stata dataset

## Syntax

```
void  _st_store(real scalar i, real scalar j, real scalar x)

void  st_store(real matrix i, rowvector j, real matrix X) (1,2)

void  st_store(real matrix i, rowvector j, scalar selectvar, real matrix X) (1,2,3)

void  _st_sstore(real scalar i, real scalar j, string scalar s)

void  st_sstore(real matrix i, rowvector j, string matrix X) (1,2)

void  st_sstore(real matrix i, rowvector j, scalar selectvar, string matrix X) (1,2,3)
```

where

1. *i* may be specified in the same way as with `st_data()`.
2. *j* may be specified in the same way as with `st_data()`, except that time-series operators may not be specified.
3. *selectvar* may be specified in the same way as with `st_data()`.

See [\[M-5\] st\\_data\(\)](#).

## Description

These functions mirror `_st_data()`, `st_data()`, and `st_sdata()`. Rather than returning the contents from the Stata dataset, these commands change those contents to be as given by the last argument.

## Remarks

See [\[M-5\] st\\_data\(\)](#).

## Conformability

```
_st_store(i, j, x), _st_sstore(i, j, x):
    i:      1 × 1
    j:      1 × 1
    x:      1 × 1
    result: void

st_store(i, j, X), st_sstore(i, j, X):
    i:      n × 1    or    n2 × 2
    j:      1 × k
    X:      n × k
    result: void
```

```
st_store(i, j, selectvar, X), st_sstore(i, j, selectvar, X):
```

```

      i:       $n \times 1$    or    $n_2 \times 2$ 
      j:       $1 \times k$ 
selectvar:   $1 \times 1$ 
      X:       $(n - e) \times k$ , where e is number of observations excluded by selectvar
      result: void
```

## Diagnostics

`_st_store(i, j, x)` and `_st_sstore(i, j, s)` do nothing if *i* or *j* is out of range; they do not abort with error.

`st_store(i, j, X)` and `st_sstore(i, j, s)` abort with error if any element of *i* or *j* is out of range. *j* may be specified as a vector of variable names or as a vector of variable indices. If names are specified, abbreviations are allowed. If you do not want this, use `st_varindex()` (see [M-5] [st\\_varindex\(\)](#)) to translate variable names into variable indices.

`st_store()` and `st_sstore()` abort with error if *X* is not [p-conformable](#) with the matrix that `st_data()` (`st_sdata()`) would return.

## Also see

[M-5] [st\\_data\(\)](#) — Load copy of current Stata dataset

[M-5] [st\\_addvar\(\)](#) — Add variable to current Stata dataset

[M-4] [stata](#) — Stata interface functions

[D] [putmata](#) — Put Stata variables into Mata and vice versa

## Syntax

```
void st_subview(X, transmorphic matrix V, real matrix i, real matrix j)
```

where

1. The type of  $X$  does not matter; it is replaced.
2.  $V$  is typically a view, but that is not required.  $V$ , however, must be real or string.

## Description

`st_subview( $X$ ,  $V$ ,  $i$ ,  $j$ )` creates new view matrix  $X$  from existing view matrix  $V$ .  $V$  is to have been created from a previous call to `st_view()` (see [\[M-5\] st\\_view\(\)](#)) or `st_subview()`.

Although `st_subview()` is intended for use with view matrices, it may also be used when  $V$  is a regular matrix. Thus code may be written in such a way that it will work without regard to whether a matrix is or is not a view.

$i$  may be specified as a  $1 \times 1$  scalar, a  $1 \times 1$  scalar containing missing, as a column vector of row numbers, as a row vector specifying a row-number range, or as a  $k \times 2$  matrix specifying both:

- a. `st_subview( $X$ ,  $V$ , 1,2)` makes  $X$  equal to the first row of the second column of  $V$ .
- b. `st_subview( $X$ ,  $V$ , .,2)` makes  $X$  equal to all rows of the second column of  $V$ .
- c. `st_subview( $X$ ,  $V$ , (1\2\5),2)` makes  $X$  equal to rows 1, 2, and 5 of the second column of  $V$ .
- d. `st_subview( $X$ ,  $V$ , (1,5),2)` makes  $X$  equal to rows 1 through 5 of the second column of  $V$ .
- e. `st_subview( $X$ ,  $V$ , (1,5\7,9),2)` makes  $X$  equal to rows 1 through 5 and 7 through 9 of the second column of  $V$ .
- f. When a range is specified, any element of the range  $(i_1, i_2)$  may be set to contribute zero observations if  $i_2 = i_1 - 1$ . For example,  $(1,0)$  is not an error and neither is  $(1,0\5,7)$ .

$j$  may be specified in the same way as  $i$ , except transposed, to specify the selected columns:

- a. `st_subview( $X$ ,  $V$ , 2,.)` makes  $X$  equal to all columns of the second row of  $V$ .
- b. `st_subview( $X$ ,  $V$ , 2,(1,2,5))` makes  $X$  equal to columns 1, 2, and 5 of the second row of  $V$ .
- c. `st_subview( $X$ ,  $V$ , 2,(1\5))` makes  $X$  equal to columns 1 through 5 of the second row of  $V$ .
- d. `st_subview( $X$ ,  $V$ , 2,((1\5),(7\9)))` makes  $X$  equal to columns 1 through 5 and 7 through 9 of the second row of  $V$ .



- e. When a range is specified, any element of the range  $(j_1 \setminus j_2)$  may be set to contribute zero columns if  $j_2 = j_1 - 1$ . For example,  $(1 \setminus 0)$  is not an error and neither is  $((1 \setminus 0), (5 \setminus 7))$ .

Obviously, notations for  $i$  and  $j$  can be specified simultaneously:

- `st_subview(X, V, ., .)` makes  $X$  a duplicate of  $V$ .
- `st_subview(X, V, ., (1\5))` makes  $X$  equal to columns 1 through 5 of all rows of  $X$ .
- `st_subview(X, V, (10,25), (1\5))` makes  $X$  equal to columns 1 through 5 of rows 10 through 25 of  $X$ .

Also, `st_subview()` may be used to create views with duplicate variables or observations from  $V$ .

## Remarks

Say that you need to make a calculation on matrices  $X$  and  $Y$ , which might be views. Perhaps the calculation is `invsym(X'X)*X'Y`. Regardless, you start as follows:

```
st_view(X, ., "v2 v3 v4", 0)
st_view(Y, ., "v1 v7"    , 0)
```

You are already in trouble. You smartly coded fourth argument as 0, meaning exclude the missing values, but you do not know that the same observations were excluded in the manufacturing of  $X$  as in the manufacturing of  $Y$ .

If you had previously created a `touse` variable in your dataset marking the observations to be used in the calculation, one solution would be

```
st_view(X, ., "v2 v3 v4", "touse")
st_view(Y, ., "v1 v7"    , "touse")
```

That solution is recommended, but let's assume you did not do that. The other solution is

```
st_view(M, ., "v2 v3 v4 v1 v7", 0)
st_subview(X, M, ., (1,2,3))
st_subview(Y, M, ., (4,5))
```

The first call to `st_view()` will eliminate observations with missing values on any of the variables, and the second two `st_subview()` calls will create the matrices you wanted, obtaining them from the correctly formed  $M$ . Basically, the two `st_subview()` calls amount to the same thing as

```
X = M[, (1,2,3)]
Y = M[, (4,5)]
```

but you do not want to code that because then matrices  $X$  and  $Y$  would contain copies of the data, and you are worried that the dataset might be large.

For a second example, let's pretend that you are processing a panel dataset and making calculations from matrix *X* within panel. Your code looks something like

```
st_view(id, ., "panelid", 0)
for (i=1; i<=rows(id); i=j+1) {
    j = endobs(id, i)
    st_view(X, (i,j), "v1 v2 ...", 0)
    ...
}
```

where you have previously written function `endobs()` to be

```
scalar endobs(vector id, scalar i)
{
    scalar    j
    for (j=i+1; j<=rows(id); j++) {
        if (id[j]!=id[i]) return(j-1)
    }
    return(rows(id))
}
```

In any case, there could be a problem. Missing values of variable `panelid` might not align with missing values of variables `v1`, `v2`, etc. The result could be that observation and row numbers are not in accordance or that there appears to be a group that, in fact, has all missing data. The right way to handle the problem is

```
st_view(M, ., "panelid v1 v2 ...", 0)
st_subview(id, M, ., 1)
for (i=1; i<=rows(id); i=j+1) {
    j = endobs(id, i)
    st_subview(X, M, (i,j), (2\cols(M)))
    ...
}
```

## Conformability

`st_subview(X, V, i, j):`

*input:*

<i>V</i> :	$r \times c$	
<i>i</i> :	$1 \times 1, n \times 1,$	or $n_2 \times 2$
<i>j</i> :	$1 \times 1, 1 \times k,$	or $2 \times k_2$

*output:*

<i>X</i> :	$n \times k$
------------	--------------

## Diagnostics

`st_subview(X, V, i, j)` aborts with error if *i* or *j* are out of range. *i* and *j* refer to row and column numbers of *V*, not observation and variable numbers of the underlying Stata dataset.

## Also see

[M-5] `st_view()` — Make matrix that is a view onto current Stata dataset

[M-5] `select()` — Select rows or columns

[M-4] `stata` — Stata interface functions

# Title

[M-5] **st\_tempname()** — Temporary Stata names

## Syntax

```
string scalar      st_tempname()
string rowvector   st_tempname(real scalar n)

string scalar      st_tempfilename()
string rowvector   st_tempfilename(real scalar n)
```

## Description

`st_tempname()` returns a Stata temporary name, the same as would be returned by Stata's `tempvar` and `tempname` commands; see [P] [macro](#).

`st_tempname(n)` returns *n* temporary Stata names,  $n \geq 0$ .

`st_tempfilename()` returns a Stata temporary filename, the same as would be returned by Stata's `tempfile` command; see [P] [macro](#).

`st_tempfilename(n)` returns *n* temporary filenames,  $n \geq 0$ .

## Remarks

Remarks are presented under the following headings:

*Creating temporary objects*  
*When temporary objects will be eliminated*

### Creating temporary objects

`st_tempname()`s can be used to name Stata's variables, matrices, and scalars. Although in Stata a distinction is drawn between `tempvars` and `tempnames`, there is no real distinction, and so `st_tempname()` handles both in Mata. For instance, one can create a temporary variable by coding

```
idx = st_addvar("double", st_tempname())
```

See [M-5] [st\\_addvar\(\)](#).

One creates a temporary file by coding

```
fh = fopen(st_tempfilename(), "w")
```

See [M-5] [fopen\(\)](#).

## When temporary objects will be eliminated

Temporary objects do not vanish when the Mata function ends, nor when Mata itself ends. They are removed when the ado-file (or do-file) calling Mata terminates.

Forget Mata for a minute. Stata eliminates temporary variables and files when the program that created them ends. That same rule applies to Mata: Stata eliminates them, not Mata, and that means that the ado-file or do-file that called Mata will eliminate them when that ado-file or do-file ends. Temporary variables and files are not eliminated by Mata when the Mata function ends. Thus Mata functions can create temporary objects for use by their ado-file callers, should that prove useful.

## Conformability

`st_tempname()`, `st_tempfilename()`:

*result*:  $1 \times 1$

`st_tempname(n)`, `st_tempfilename(n)`:

*n*:  $1 \times 1$

*result*:  $1 \times n$

## Diagnostics

`st_tempname(n)` and `st_tempfilename(n)` abort with error if  $n < 0$  and return `J(1,0,"")` if  $n = 0$ .

## Also see

[M-5] [st\\_addvar\(\)](#) — Add variable to current Stata dataset

[M-4] [stata](#) — Stata interface functions

# Title

**[M-5] st\_tsrevar()** — Create time-series *op.varname* variables

## Syntax

```
real rowvector st_tsrevar(string rowvector s)
```

```
real rowvector _st_tsrevar(string rowvector s)
```

## Description

`st_tsrevar(s)` is the equivalent of Stata's [\[TS\] tsrevar](#) programming command: it generates temporary variables containing the evaluation of any *op.varname* combinations appearing in *s*.

`_st_tsrevar(s)` does the same thing as `st_tsrevar()`. The two functions differ in how they respond to invalid elements of *s*. `st_tsrevar()` aborts with error, and `_st_tsrevar()` places missing in the appropriate element of the returned result.

## Remarks

Both of these functions help achieve efficiency when using views and time-series variables. Assume that in *vars*, you have a list of Stata variable names, some of which might contain time-series *op.varname* combinations such as `l.gnp`. For example, *vars* might contain

```
vars = "gnp r l.gnp"
```

If you wanted to create in *V* a view on the data, you would usually code

```
st_view(V, ., vars)
```

We are not going to do that, however, because we plan to do many calculations with *V* and, to speed execution, we want any *op.varname* combinations evaluated just once, as *V* is created. Of course, if efficiency were our only concern, we would code

```
V = st_data(., vars)
```

Assume, however, that we have lots of data, so memory is an issue, and yet we still want as much efficiency as possible given the constraint of not copying the data. The solution is to code

```
st_view(V, ., st_tsrevar(tokens(vars)))
```

`st_tsrevar()` will create temporary variables for each *op.varname* combination (`l.gnp` in our example), and then return the Stata variable indices of each of the variables, whether newly created or already existing. If `gnp` was the second variable in the dataset, `r` was the 23rd, and in total there were 54 variables, then returned by `st_tsrevar()` would be `(2, 23, 55)`. Variable 55 is new, created by `st_tsrevar()`, and it contains the values of `l.gnp`. The new variable is temporary and will be dropped automatically at the appropriate time.

## Conformability

```
st_tsrevar(s), _st_tsrevar(s):  
    s:           $1 \times c$   
    result:      $1 \times c$ 
```

## Diagnostics

`st_tsrevar()` aborts with error if any variable name is not found or any *op.varname* combination is invalid.

`_st_tsrevar()` puts missing in the appropriate element of the returned result for any variable name that is not found or any *op.varname* combination that is invalid.

## Also see

[M-5] `st_varname()` — Obtain variable names from variable indices

[M-4] `stata` — Stata interface functions

[M-5] `st_varindex()` — Obtain variable indices from variable names

# Title

[M-5] **st\_updata()** — Determine or set data-have-changed flag

## Syntax

```
real scalar  st_updata()  
void         st_updata(real scalar value)
```

## Description

`st_updata()` returns 0 if the data in memory have not changed since they were last saved and returns 1 otherwise.

`st_updata(value)` sets the data-have-changed flag to 0 if *value* = 0 and 1 otherwise.

## Remarks

Stata's `describe` command (see [\[D\] describe](#)) reports whether the data have changed since they were last saved. Stata's `use` command (see [\[D\] use](#)) refuses to load a new dataset if the data currently in memory have not been saved since they were last changed. Other components of Stata also react to the data-have-changed flag.

`st_updata()` allows you to respect that same flag.

Also, as a Mata programmer, you must set the flag if your function changes the data in memory. Mata attempts to set the flag for you (for instance, when you add a new variable using `st_addvar()` [see [\[M-5\] st\\_addvar\(\)](#)]), but there are other places where the flag ought to be set, and you must do so. For instance, Mata does not set the flag every time you change a value in the dataset. Setting the flag what may be many thousands of times would reduce performance too much.

Moreover, even when Mata does set the flag, it might do so inappropriately, because the logic of your program fooled Mata. For instance, perhaps you added a variable and later dropped it. In such cases, the appropriate code is

```
priorupdatavalue = st_updata()  
...  
st_updata(priorupdatavalue)
```

## Conformability

```
st_updata():  
  result:      1 × 1  
  
st_updata(value):  
  value:      1 × 1  
  result:      void
```



## Diagnostics

None.

## Also see

[M-4] [stata](#) — Stata interface functions

## Syntax

```

string scalar  st_varformat(scalar var)

void           st_varformat(scalar var, string scalar fmt)

string scalar  st_varlabel(scalar var)

void           st_varlabel(scalar var, string scalar label)

string scalar  st_varvalue label(scalar var)

void           st_varvalue label(scalar var, string scalar labelname)
    
```

where *var* contains a Stata variable name or a Stata variable index.

## Description

`st_varformat(var)` returns the display format associated with *var*, such as "%9.0gc".  
`st_varformat(var, fmt)` changes *var*'s display format.

`st_varlabel(var)` returns the variable label associated with *var*, such as "Sex of Patient", or it returns "" if *var* has no variable label. `st_varformat(var, label)` changes *var*'s variable label.

`st_varvalue label(var)` returns the value-label name associated with *var*, such as "origin", or it returns "" if *var* has no value label. `st_varvalue label(var, labelname)` changes the value-label name associated with *var*.

## Conformability

```

st_varformat(var), st_varlabel(var), st_varvalue label(var):
    var:      1 × 1
    result:   1 × 1

st_varformat(var, fmt), st_varlabel(var, label), st_varvalue label(var, labelname):
    var:      1 × 1
    value:    1 × 1
    result:   void
    
```

## Diagnostics

In all functions, if *var* is specified as a name, abbreviations are not allowed.

All functions abort with error if *var* is not a valid Stata variable.

`st_varformat(var, fmt)` aborts with error if *fmt* does not contain a valid display format for *var*.

`st_varlabel(var, label)` will truncate *label* if it is too long.

`st_varvalue_label(var, labelname)` aborts with error if *var* is a Stata string variable or if *labelname* does not contain a valid name (assuming *labelname* is not ""). It is not required, however, that the label name exist.

## Also see

[M-4] [stata](#) — Stata interface functions

**[M-5] `st_varindex()`** — Obtain variable indices from variable names

## Syntax

```

real rowvector    st_varindex(string rowvector s)

real rowvector    st_varindex(string rowvector s, real scalar abbrev)

real rowvector    _st_varindex(string rowvector s)

real rowvector    _st_varindex(string rowvector s, real scalar abbrev)

```

## Description

`st_varindex(s)` returns the variable index associated with each variable name recorded in *s*. `st_varindex(s)` does not allow variable-name abbreviations such as "pr" for "price".

`st_varindex(s, abbrev)` does the same thing but allows you to specify whether variable-name abbreviations are to be allowed. Abbreviations are allowed if *abbrev*  $\neq$  0. `st_varindex(s)` is equivalent to `st_varindex(s, 0)`.

`_st_varindex()` does the same thing as `st_varindex()`. The two functions differ in how they respond when a name is not found. `st_varindex()` aborts with error, and `_st_varindex()` places missing in the appropriate element of the returned result.

## Remarks

These functions require that each element of *s* contain a variable name, such as

```
s = ("price", "mpg", "weight")
```

If you have one string containing multiple names

```
s = ("price mpg weight")
```

then use `tokens()` to split it into the desired form, as in

```
k = st_varindex(tokens(s))
```

See [M-5] [tokens\(\)](#).

## Conformability

```

st_varindex(s, abbrev), _st_varindex(s, abbrev):
    s:          1 × k
    abbrev:     1 × 1   (optional)
    result:     1 × k

```

## Diagnostics

`st_varindex()` aborts with error if any name is not found.

`_st_varindex()` puts missing in the appropriate element of the returned result for any name that is not found.

## Also see

[M-5] `st_varname()` — Obtain variable names from variable indices

[M-5] `tokens()` — Obtain tokens from string

[M-4] `stata` — Stata interface functions

# Title

[M-5] **st\_varname()** — Obtain variable names from variable indices

## Syntax

*string rowvector*    **st\_varname**(*real rowvector k*)

*string rowvector*    **st\_varname**(*real rowvector k*, *real scalar tsmat*)

## Description

**st\_varname**(*k*) returns the Stata variable names associated with the variable indices stored in *k*. For instance, with the automobile data in memory

```
names = st_varname((1..3))
```

results in *names* being ("make", "price", "mpg").

**st\_varname**(*k*, *tsmap*) does the same thing but allows you to specify whether you want the actual or logical variable names of any time-series-operated variables created by the Mata function **st\_tsrevar**() (see [M-5] **st\_tsrevar**()) or by the Stata command **tsrevar** (see [TS] **tsrevar**).

**st\_varname**(*k*) is equivalent to **st\_varname**(*k*, 0); actual variable names are returned.

**st\_varname**(*k*, 1) returns logical variable names.

## Remarks

To understand the actions of **st\_varname**(*k*, 1), pretend that variable 58 was created by **st\_tsrevar**():

```
k = st_tsrevar(("gnp", "r", "l.gnp"))
```

Pretend that *k* now contains (12, 5, 58). Variable 58 is a new, temporary variable, containing *l.gnp* values. Were you to ask for the actual names of the variables

```
actualnames = st_varname(k)
```

*actualnames* would contain ("gnp", "r", "\_\_00004a"), although the name of the last variable will vary because it is a temporary variable. Were you to ask for the logical names,

```
logicalnames = st_varname(k, 1)
```

you would get back ("gnp", "r", "L.gnp").

## Conformability

**st\_varname**(*k*, *tsmap*)

<i>k</i> :	$1 \times c$	
<i>tsmap</i> :	$1 \times 1$	(optional)
<i>result</i> :	$1 \times c$	

## Diagnostics

`st_varname(k)` and `st_varname(k, tmap)` abort with error if any element of *k* is less than 1 or greater than `st_nvar()`; see [M-5] `st_nvar()`.

## Also see

[M-5] `st_varindex()` — Obtain variable indices from variable names

[M-5] `st_tsrevar()` — Create time-series op.varname variables

[M-4] `stata` — Stata interface functions

# Title

[M-5] **st\_varrename()** — Rename Stata variable

## Syntax

`void st_varrename(scalar var, string scalar newname)`

where *var* contains a Stata variable name or a Stata variable index.

## Description

`st_varrename(var, newname)` changes the name of *var* to *newname*.

If *var* is specified as a name, abbreviations are not allowed.

## Conformability

<code>st_varrename(<i>var</i>, <i>newname</i>):</code>	
<i>var</i> :	$1 \times 1$
<i>newname</i> :	$1 \times 1$
<i>result</i> :	<i>void</i>

## Diagnostics

`st_varrename(var, newname)` aborts with error if *var* is not a valid Stata variable or if *newname* is not a valid name or if a variable named *newname* already exists.

## Also see

[\[M-4\] stata](#) — Stata interface functions



# Title

[M-5] **st\_vartype()** — Storage type of Stata variable

## Syntax

```
string scalar  st_vartype(scalar var)
real scalar    st_isnumvar(scalar var)
real scalar    st_isstrvar(scalar var)
```

where *var* contains a Stata variable name or a Stata variable index.

## Description

In all the functions, if *var* is specified as a name, abbreviations are not allowed.

`st_vartype(var)` returns the [storage type](#) of the *var*, such as `float`, `double`, or `str18`.

`st_isnumvar(var)` returns 1 if *var* is a numeric variable and 0 otherwise.

`st_isstrvar(var)` returns 1 if *var* is a string variable and 0 otherwise.

## Remarks

`st_isstrvar(var)` and `st_isnumvar(var)` are antonyms. Both functions are provided merely for convenience; they tell you nothing that you cannot discover from `st_vartype(var)`.

## Conformability

```
st_vartype(var):
  var:      1 × 1
  result:    1 × 1

st_isnumvar(var), st_isstrvar(var):
  var:      1 × 1
  result:    1 × 1
```

## Diagnostics

All functions abort with error if *var* is not a valid Stata variable.

## Also see

[\[M-4\] stata](#) — Stata interface functions

# Title

**[M-5] st\_view()** — Make matrix that is a view onto current Stata dataset

## Syntax

```
void st_view(V, real matrix i, rowvector j)
```

```
void st_view(V, real matrix i, rowvector j, scalar selectvar)
```

```
void st_sview(V, real matrix i, rowvector j)
```

```
void st_sview(V, real matrix i, rowvector j, scalar selectvar)
```

where

1. The type of *V* does not matter; it is replaced.
2. *i* may be specified in the same way as with `st_data()`.
3. *j* may be specified in the same way as with `st_data()`. Factor variables and time-series-operated variables may be specified.
4. *selectvar* may be specified in the same way as with `st_data()`.

See [\[M-5\] st\\_data\(\)](#).

## Description

`st_view()` and `st_sview()` create a matrix that is a view onto the current Stata dataset.

## Remarks

Remarks are presented under the following headings:

[Overview](#)

[Advantages and disadvantages of views](#)

[When not to use views](#)

[Cautions when using views 1: Conserving memory](#)

[Cautions when using views 2: Assignment](#)

[Efficiency](#)

## Overview

`st_view()` serves the same purpose as `st_data()`—and `st_sview()` serves the same purpose as `st_sdata()`—except that, rather than returning a matrix that is a copy of the underlying values, `st_view()` and `st_sview()` create a matrix that is a view onto the Stata dataset itself.

To understand the distinction, consider

```
X = st_data(., "mpg displ weight")
```

and

```
st_view(X, ., "mpg displ weight")
```

Both commands fill in matrix `X` with the same data. However, were you to code

```
X[2,1] = 123
```

after the `st_data()` setup, you would change the value in the matrix `X`, but the Stata dataset would remain unchanged. After the `st_view()` setup, changing the value in the matrix would cause the value of `mpg` in the second observation to change to 123.

## Advantages and disadvantages of views

Views make it easy to change the dataset, and that can be an advantage or a disadvantage, depending on your goals.

Putting that aside, views are in general better than copies because 1) they take less time to set up and 2) they consume less memory. The memory savings can be considerable. Consider a 100,000-observation dataset on 30 variables. Coding

```
X = st_data(., .)
```

creates a new matrix that is 24 MB in size. Meanwhile, the total storage requirement for

```
st_view(X, ., .)
```

is roughly 128 bytes!

There is a cost; when you use the matrix `X`, it takes longer to access the individual elements. You would have to do a lot of calculation with `X`, however, before that sum of the longer access times would equal the initial savings in setup time, and even then, the longer access time is probably worth the savings in memory.

## When not to use views

Do not use views as a substitute for scalars. If you are going to loop through the data an observation at a time, and if every usage you will make of `X` is in scalar calculations, use `_st_data()`. There is nothing faster for that problem.

Putting aside that extreme, views become more efficient relative to copies the larger they are; that is, it is more efficient to use `st_data()` for small amounts of data, especially if you are going to make computationally intensive calculations with it.

## Cautions when using views 1: Conserving memory

If you are using views, it is probably because you are concerned about memory, and if you are, you want to be careful to avoid making copies of views. Copies of views are not views; they are copies. For instance,

```
st_view(V, ., .)
Y = V
```

That innocuous looking `Y = V` just made a copy of the entire dataset, meaning that if the dataset had 100,000 observations on 30 variables, `Y` now consumes 24 MB. Coding `Y = V` may be desirable in certain circumstances, but in general, it is better to set up another view.

Similarly, watch out for subscripts. Consider the following code fragment

```
st_view(V, ., .)
for (i=1; i<=cols(V); i++) {
    sum = colsum(V[,i])
    ...
}
```

The problem in the above code is the `V[,i]`. That creates a new column vector containing the values from the *i*th column of *V*. Given 100,000 observations, that new column vector needs 800k of memory. Better to code would be

```
for (i=1; i<=cols(V); i++) {
    st_view(v, ., i)
    sum = colsum(v)
    ...
}
```

If you need *V* and *v*, that is okay. You can have many views of the data setup simultaneously.

Similarly, be careful using views with operators.  $X'X$  makes a copy of *X* in the process of creating the transpose. Use functions such as `cross()` (see [M-5] [cross\(\)](#)) that are designed to minimize the use of memory.

Do not be overly concerned about this issue. Making a copy of a column of a view amounts to the same thing as introducing a temporary variable in a Stata program—something that is done all the time.

## Cautions when using views 2: Assignment

The ability to assign to a view and so change the underlying data can be either convenient or dangerous, depending on your goals. When making such assignments, there are two things you need be aware of.

The first is more of a Stata issue than it is a Mata issue. Assignment does not cause promotion. Coding

```
V[1,2] = 4059.125
```

might store 4059.125 in the first observation of the second variable of the view. Or, if that second variable is an `int`, what will be stored is 4059, or if it is a `byte`, what will be stored is missing.

The second caution is a Mata issue. To reassign all the values of the view, code

```
V[,.] = matrix_expression
```

Do not code

```
V = matrix_expression
```

The second expression does not assign to the underlying dataset, it redefines *V* to be a regular matrix.

Mata does not allow the use of views as the destination of assignment when the view contains factor variables or time-series-operated variables such as `i.rep78` or `l.gnp`.

## Efficiency

Whenever possible, specify argument  $i$  of `st_view(V, i, j)` and `st_sview(V, i, j)` as `.` (missing value) or as a row vector range (for example,  $(i_1, i_2)$ ) rather than as a column vector list.

Specify argument  $j$  as a real row vector rather than as a string whenever `st_view()` and `st_sview()` are used inside loops with the same variables (and the view does not contain factor variables nor time-series-operated variables). This prevents Mata from having to look up the same names over and over again.

## Conformability

`st_view(V, i, j), st_sview(V, i, j):`

*input:*

$i:$        $n \times 1$     or     $n_2 \times 2$   
 $j:$        $1 \times k$     or     $1 \times 2$  containing  $k$  elements when expanded

*output:*

$V:$        $n \times k$

`st_view(V, i, j, selectvar), st_sview(V, i, j, selectvar):`

*input:*

$i:$        $n \times 1$     or     $n_2 \times 2$   
 $j:$        $1 \times k$     or     $1 \times 2$  containing  $k$  elements when expanded

*selectvar:*     $1 \times 1$

*output:*

$V:$        $(n - e) \times k$ , where  $e$  is number of observations excluded by *selectvar*

## Diagnostics

`st_view(i, j[, selectvar])` and `st_sview(i, j[, selectvar])` abort with error if any element of  $i$  is outside the range of observations or if a variable name or index recorded in  $j$  is not found. Variable-name abbreviations are allowed. If you do not want this and no factor variables nor time-series-operated variables are specified, use `st_varindex()` (see [M-5] [st\\_varindex\(\)](#)) to translate variable names into variable indices.

`st_view()` and `st_sview()` abort with error if any element of  $i$  is out of range as described under the heading [Details of observation subscripting using st\\_data\(\) and st\\_sdata\(\)](#) in [M-5] [st\\_data\(\)](#).

Some functions do not allow views as arguments. If `example(X)` does not allow views, you can still use it by coding

```
... example(X=V) ...
```

because that will make a copy of view  $V$  in  $X$ . Most functions that do not allow views mention that in their *Diagnostics* section, but some do not because it was unexpected that anyone would want to use a view in that case. If a function does not allow a view, you will see in the traceback log:

```
: myfunction(...)
      example(): 3103 view found where array required
      mysub():   - function returned error
      myfunction(): - function returned error
      <istmt>:    - function returned error
r(3103);
```

The above means that function `example()` does not allow views.

## Reference

Gould, W. W. 2005. [Mata Matters: Using views onto the data](#). *Stata Journal* 5: 567–573.

## Also see

[M-5] [st\\_subview\(\)](#) — Make view from view

[M-5] [select\(\)](#) — Select rows or columns

[M-5] [st\\_viewvars\(\)](#) — Variables and observations of view

[M-5] [st\\_data\(\)](#) — Load copy of current Stata dataset

[M-4] [stata](#) — Stata interface functions

[D] [putmata](#) — Put Stata variables into Mata and vice versa

# Title

[M-5] **st\_viewvars()** — Variables and observations of view

## Syntax

*real rowvector*    **st\_viewvars**(*matrix V*)

*real vector*        **st\_viewobs**(*matrix V*)

where  $V$  is required to be a view.

## Description

**st\_viewvars**( $V$ ) returns the indices of the Stata variables corresponding to the columns of  $V$ .

**st\_viewobs**( $V$ ) returns the Stata observation numbers corresponding to the rows of  $V$ . Returned is either a  $1 \times 2$  row vector recording the observation range or an  $N \times 1$  column vector recording the individual observation numbers.

## Remarks

The results returned by these two functions are suitable for inclusion as arguments in subsequent calls to **st\_view()** and **st\_sview()**; see [M-5] **st\_view()**.

## Conformability

**st\_viewvars**( $V$ ):

$V$ :         $N \times k$   
*result*:     $1 \times k$

**st\_viewobs**( $V$ ):

$V$ :         $N \times k$   
*result*:     $1 \times 2$    or    $N \times 1$

## Diagnostics

**st\_viewvars**( $V$ ) and **st\_viewobs**( $V$ ) abort with error if  $V$  is not a view.

## Also see

[M-5] **st\_view()** — Make matrix that is a view onto current Stata dataset

[M-4] **stata** — Stata interface functions

## Syntax

```
real scalar    st_vlexists(name)  
void           st_vldrop(name)  
string matrix st_vlmap(name, real matrix values)  
real matrix   st_vlsearch(name, string matrix text)  
void          st_vlload(name, values, text)  
void          st_vlmodify(name, real colvector values, real colvector text)
```

where *name* is *string scalar* and where the types of *values* and *text* in `st_vlload()` are irrelevant because they are replaced.

## Description

`st_vlexists(name)` returns 1 if value label *name* exists and returns 0 otherwise.

`st_vldrop(name)` drops value label *name* if it exists.

`st_vlmap(name, values)` maps *values* through value label *name* and returns the result.

`st_vlsearch(name, text)` does the reverse; it returns the value corresponding to the text.

`st_vlload(name, values, text)` places value label *name* into *values* and *text*.

`st_vlmodify(name, values, text)` creates a new value label or modifies an existing one.

## Remarks

Value labels are named and record a mapping from numeric values to text. For instance, a value label named `sex1b1` might record that 1 corresponds to male and 2 to female. Values labels are attached to Stata numeric variables. If a Stata numeric variable had the value label `sex1b1` attached to it, then the 1s and 2s in the variable would display as male and female. How other values would appear—if there were other values—would not be affected.

Remarks are presented under the following headings:

[Value-label mapping](#)

[Value-label creation and editing](#)

[Loading value labels](#)



## Value-label mapping

Let us consider value label `sexlbl` mapping 1 to male and 2 to female.

`st_vlmap("sexlbl", values)` would map the  $r \times c$  matrix values through `sexlbl` and return an  $r \times c$  string matrix containing the result. Any values for which there was no mapping would result in `"`. Thus

```
: res = st_vlmap("sexlbl", 1)
: res
male
: res = st_vlmap("sexlbl", (2,3,1))
: res
```

	1	2	3
1	female		male

`st_vlsearch(name, text)` performs the reverse mapping:

```
: txt = st_vlsearch("sexlbl", ("female","", "male"))
: txt
```

	1	2	3
1	2	.	1

## Value-label creation and editing

`st_vlmodify(name, values, text)` creates new value labels and modifies existing ones.

If value label `sexlbl` did not exist, coding

```
: st_vlmodify("sexlbl", (1\2), ("male\"female"))
```

would create it. If the value label did previously exist, the above would modify it so that 1 now corresponds to male and 2 to female, regardless of what 1 or 2 previously corresponded to, if they corresponded to anything. Other mappings that might have been included in the value label remain unchanged. Thus

```
: st_vlmodify("sexlbl", 3, "unknown")
```

would add another mapping to the label. Values are deleted by specifying the text as `"`, so

```
: st_vlmodify("sexlbl", 3, "")
```

would remove the mapping for 3 (if there was a mapping). If you remove all the mappings, the value label itself is automatically dropped:

```
: st_vlmodify("sexlbl", (1\2), ("\""))
```

results in value label `sexlbl` being dropped if 1 and 2 were the final mappings in it.

## Loading value labels

`st_vlload(name, values, text)` returns the value label in *values* and *text*, where you can do with it as you please. Thus you could code

```
st_vlload("sexlbl", values, text)
...
st_vldrop("sexlbl")
st_vlmodify("sexlbl", values, text)
```

## Conformability

```
st_vlexists(name):
    name:      1 × 1
    result:    1 × 1

st_vldrop(name):
    name:      1 × 1
    result:    void

st_vlmap(name, values):
    name:      1 × 1
    values:    r × c
    result:    r × c

st_vlsearch(name, text):
    name:      1 × 1
    text:      r × c
    result:    r × c

st_vlload(name, values, text):

    input:
        name:      1 × 1

    output:
        values:    k × 1
        text:      k × 1

st_vlmodify(name, values, text):
    name:      1 × 1
    values:    m × 1
    text:      m × 1
    result:    void
```

## Diagnostics

The only conditions under which the above functions abort with error is when *name* is malformed or Mata is out of memory. Functions tolerate all other problems.

`st_vldrop(name)` does nothing if value label *name* does not exist.

`st_vlmap(name, values)` returns `J(rows(values), cols(values), "")` if value label *name* does not exist. When the value label does exist, individual values for which there is no recorded mapping are returned as "".

`st_vlsearch(name, text)`: returns `J(rows(values), cols(values), .)` if value label *name* does not exist. When the value label does exist, individual text values for which there is no corresponding value are returned as `.` (missing).

`st_vlload(name, values, text)`: sets *values* and *text* to be  $0 \times 1$  when value label *name* does not exist.

`st_vlmodify(name, values, text)`: creates the value label if it does not already exist. Value labels may map only integers and `.a`, `.b`, ..., `.z`. Attempts to insert a mapping for `.` are ignored. Noninteger values are truncated to integer values. If an element of *text* is `"`, then the corresponding mapping is removed.

## Also see

[M-4] [stata](#) — Stata interface functions

## Syntax

```

void          stata(cmd)
void          stata(cmd, nooutput)
void          stata(cmd, nooutput, nomacroexpand)

real scalar   _stata(cmd)
real scalar   _stata(cmd, nooutput)
real scalar   _stata(cmd, nooutput, nomacroexpand)
    
```

where

<i>cmd</i> :	<i>string scalar</i>
<i>nooutput</i> :	<i>real scalar</i>
<i>nomacroexpand</i> :	<i>real scalar</i>

## Description

`stata(cmd)` executes the Stata command contained in the string scalar *cmd*. Output from the command appears at the terminal, and any macros contained in *cmd* are expanded.

`stata(cmd, nooutput)` does the same thing, but if *nooutput*  $\neq$  0, output produced by the execution is not displayed. `stata(cmd, 0)` is equivalent to `stata(cmd)`.

`stata(cmd, nooutput, nomacroexpand)` does the same thing but, before execution, suppresses expansion of any macros contained in *cmd* if *nomacroexpand*  $\neq$  0. `stata(cmd, 0, 0)` is equivalent to `stata(cmd)`.

`_stata()` repeats the syntaxes of `stata()`. The difference is that, whereas `stata()` aborts with error if the execution results in a nonzero return code, `_stata()` returns the resulting return code.

## Remarks

The command you execute may invoke a process that causes another instance of Mata to be invoked. For instance, Stata program *A* calls Mata function *m1()*, which executes `stata()` to invoke Stata program *B*, which in turn calls Mata function *m2()*, which . . .

`stata(cmd)` and `_stata(cmd)` execute *cmd* at the current run level. This means that any local macros refer to local macros in the caller's space. Consider the following:

```

program example
    ...
    local x = "value from A"
    mata: myfunc()
    display "'x'"
    ...
end

mata void myfunc()
{
    stata("local x = "new value"")
}

```

After example executes `mata: myfunc()`, `'x'` will be "new value".

That `stata()` and `_stata()` work that way was intentional: Mata functions can modify the caller's environment so that they may create temporary variables for the caller's use, etc., and you only have to exercise a little caution. Executing `stata()` functions to run other ado-files and programs will cause no problems because other ado-files and programs create their own new environment in which temporary variables, local macros, etc., are private.

Also, do not use `stata()` or `_stata()` to execute a multiline command or to execute the first line of what could be considered a multiline command. Once the first line is executed, Stata will fetch the remaining lines from the caller's environment. For instance, consider

---

```

mata void myfunc()
{
    stata("if (1==1) {")
}

mata: myfunc()
display "hello"
}

```

---

In the example above, `myfunc()` will consume the `display "hello"` and `}` lines.

## Conformability

```

stata(cmd, nooutput, nomacroexpand):
    cmd:      1 × 1
    nooutput: 1 × 1 (optional)
    nomacroexpand: 1 × 1 (optional)
    result:   void

```

```

_stata(cmd, nooutput, nomacroexpand):
    cmd:      1 × 1
    nooutput: 1 × 1 (optional)
    nomacroexpand: 1 × 1 (optional)
    result:   1 × 1

```

## Diagnostics

`stata()` aborts with error if *cmd* is too long (exceedingly unlikely), if macro expansion fails, or if execution results in a nonzero return code.

`_stata()` aborts with error if *cmd* is too long.

## Also see

[M-3] [mata stata](#) — Execute Stata command

[M-4] [stata](#) — Stata interface functions

# Title

[M-5] **stataversion()** — Version of Stata being used

## Syntax

*real scalar*   stataversion()

*real scalar*   statatasetversion()

*void*   statatasetversion(*real scalar version*)

Note: the version number is multiplied by 100: Stata 2.0 is 200, Stata 5.1 is 510, and Stata 12.0 is 1200.

## Description

stataversion() returns the version of Stata/Mata that is running, multiplied by 100. For instance, if you have Stata 12 installed on your computer, stataversion() returns 1200.

statatasetversion() returns the version of Stata that has been set by the user—the version of Stata that Stata is currently emulating—multiplied by 100. Usually stataversion() == statatasetversion(). If the user has set a previous version—say, version 8 by typing version 8 in Stata—statatasetversion() will return a number less than stataversion().

statatasetversion(*version*) allows you to reset the version being emulated. Results are the same as using Stata's version command; see [P] [version](#). *version*, however, is specified as an integer equal to 100 times the version you want.

## Remarks

It is usually not necessary to reset statatasetversion(). If you do reset statatasetversion(), good form is to set it back when you are finished:

```
current_version = statatasetversion()
statatasetversion(desired_version)
...
statatasetversion(current_version)
```

## Conformability

stataversion():

*result:*     1 × 1

statatasetversion():

*result:*     1 × 1

statatasetversion(*version*):

*version:*     1 × 1

*result:*     void

## Diagnostics

`statastversion(version)` aborts with error if *version* is less than 100 or greater than `stataversion()`.

## Also see

[M-5] [bufio\(\)](#) — Buffered (binary) I/O

[M-5] [byteorder\(\)](#) — Byte order used by computer

[M-4] [programming](#) — Programming functions



# Title

[M-5] **strdup()** — String duplication

## Syntax

$n * s$

$s * n$

$n : * s$

$s : * n$

where  $n$  is real and  $s$  is string.

## Description

There is no `strdup()` function. Instead, the multiplication operator is used:

```
3*"example" = "exampleexampleexample"
0*"this" = ""
```

## Conformability

$n*s, s*n:$	
$n:$	$1 \times 1$
$s:$	$r \times c$
$result:$	$r \times c$

$n:*s, s:*n:$	
$n:$	$r_1 \times c_1$
$s:$	$r_2 \times c_2, \quad n \text{ and } s \text{ c-conformable}$
$result:$	$\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

- If  $n < 0$ , the result is as if  $n = 0$ : "" is returned.
- If  $n$  is not an integer, the result is as if `trunc( $n$ )` were specified.

## Also see

[M-4] [string](#) — String manipulation functions

# Title

[M-5] **strlen()** — Length of string

## Syntax

*real matrix*    **strlen**(*string matrix s*)

## Description

**strlen**(*s*) returns the length of—the number of characters contained in—the string *s*.

When *s* is not a scalar, **strlen**() returns element-by-element results.

## Remarks

Stata understands **strlen**() as a synonym for its own **length**() function, so you can use the function named **strlen**() in both your Stata and Mata code. Do not, however, use **length**() in Mata when you mean **strlen**(). Mata's **length**() function returns the length (number of elements) of a vector.

## Conformability

**strlen**(*s*):

<i>s</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

**strlen**(*s*), when *s* is a binary string (a string containing binary 0), returns the overall length of the string, not the location of the binary 0. Use **strpos**(*s*, **char**(0)) if you want the location of the binary 0; see [M-5] **strpos**).

## Also see

- [M-5] **strpos**() — Find substring in string
- [M-5] **fmtwidth**() — Width of %fmt
- [M-4] **string** — String manipulation functions

# Title

[M-5] **strmatch()** — Determine whether string matches pattern

## Syntax

*real matrix*    **strmatch**(*string matrix s*, *string matrix pattern*)

## Description

**strmatch**(*s*, *pattern*) returns 1 if *s* matches *pattern* and 0 otherwise.

When arguments are not scalar, **strmatch()** returns element-by-element results.

## Remarks

In *pattern*, \* means that 0 or more characters go here and ? means that exactly one character goes here. Thus *pattern*="\*" matches anything and *pattern*="?p\*x" matches all strings whose second character is *p* and whose last character is *x*.

Stata understands **strmatch()** as a synonym for its own [strmatch\(\)](#) function, so you can use the **strmatch()** function in both your Stata and Mata code.

## Conformability

**strmatch**(*s*, *pattern*):

<i>s</i> :	$r_1 \times c_1$
<i>pattern</i> :	$r_2 \times c_2$ , <i>s</i> and <i>pattern</i> r-conformable
<i>result</i> :	$\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

In **strmatch**(*s*, *pattern*), if *s* or *pattern* contain a binary 0 (they usually would not), the strings are considered to end at that point.

## Also see

[M-4] [string](#) — String manipulation functions

# Title

[M-5] **stroofreal()** — Convert real to string

## Syntax

```
string matrix stroofreal(real matrix R)  
  
string matrix stroofreal(real matrix R, string matrix format)
```

## Description

`stroofreal(R)` returns *R* as a string using Stata's %9.0g format. `stroofreal(R)` is equivalent to `stroofreal(R, "%9.0g")`.

`stroofreal(R, format)` returns *R* as a string formatted using *format*.

Leading blanks are trimmed from the result.

When arguments are not scalar, `stroofreal()` returns element-by-element results.

## Conformability

```
stroofreal(R, format):  
  R:       $r_1 \times c_1$   
  format:  $r_2 \times c_2$ , R and format r-conformable (optional)  
  result:  $\max(r_1, r_2) \times \max(c_1, c_2)$ 
```

## Diagnostics

`stroofreal(R, format)` returns "." if *format* is invalid.

## Also see

- [M-5] [strtoreal\(\)](#) — Convert string to real
- [M-4] [string](#) — String manipulation functions

# Title

[M-5] **strpos()** — Find substring in string

## Syntax

*real matrix*    **strpos**(*string matrix haystack*, *string matrix needle*)

## Description

**strpos**(*haystack*, *needle*) returns the location of the first occurrence of *needle* in *haystack* or 0 if *needle* does not occur.

When arguments are not scalar, **strpos**() returns element-by-element results.

## Remarks

When working with binary strings, one can find the location of the binary 0 using **strpos**(*s*, **char**(0)).

## Conformability

**strpos**(*haystack*, *needle*):

*haystack:*         $r_1 \times c_1$

*needle:*          $r_2 \times c_2$ ,    *haystack* and *needle* r-conformable

*result:*          $\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

**strpos**(*haystack*, *needle*) returns 0 if *needle* is not found in *haystack*.

## Also see

[\[M-4\] string](#) — String manipulation functions

# Title

[M-5] **strreverse()** — Reverse string

## Syntax

*string matrix*   **strreverse**(*string matrix s*)

## Description

**strreverse**(*s*) returns *s* with its characters in reverse order.

When *s* is not a scalar, **strreverse**() returns element-by-element results.

## Remarks

Stata understands **strreverse**() as a synonym for its own **reverse()** function, so you can use the **strreverse**() name in both your Stata and Mata code.

## Conformability

<b>strreverse</b> ( <i>s</i> )	
<i>s</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

None.

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] **strtoname()** — Convert a string to a Stata name

## Syntax

*string matrix*    **strtoname**(*string matrix s*, *real scalar p*)

*string matrix*    **strtoname**(*string matrix s*)

## Description

**strtoname**(*s*, *p*) returns *s* translated into a Stata name. Each character in *s* that is not allowed in a Stata name is converted to an underscore character, `_`. If the first character in *s* is a numeric character and *p* is not 0, then the result is prefixed with an underscore. The result is truncated to 32 characters.

**strtoname**(*s*) is equivalent to **strtoname**(*s*, 1).

When arguments are not scalar, **strtoname**() returns element-by-element results.

## Remarks

**strtoname**("StataName") returns "StataName".

**strtoname**("not a Stata name") returns "not\_a\_Stata\_name".

**strtoname**("0 is off") returns "\_0\_is\_off".

**strtoname**("0 is off", 0) returns "0\_is\_off".

## Conformability

```
strtoname(s, p):  
    s:       $r \times c$   
    p:       $1 \times 1$   
    result:  $r \times c$ 
```

```
strtoname(s):  
    s:       $r \times c$   
    result:  $r \times c$ 
```

## Diagnostics

None.

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] **strtoreal()** — Convert string to real

## Syntax

*real matrix*    **strtoreal**(*string matrix S*)  
*real scalar*    **\_strtoreal**(*string matrix S, R*)

## Description

**strtoreal**(*S*) returns *S* converted to real. Elements of *S* that cannot be converted are returned as . (missing value).

**\_strtoreal**(*S, R*) does the same as above—it returns the converted values in *R*—and it returns the number of elements that could not be converted. In such cases, the corresponding value of *R* contains . (missing).

## Remarks

**strtoreal**("1.5") returns (numeric) 1.5.

**strtoreal**("−2.5e+1") returns (numeric) −25.

**strtoreal**("not a number") returns (numeric) . (missing).

Typically, **strtoreal**(*S*) and **\_strtoreal**(*S, R*) are used with scalars, but if applied to a vector or matrix *S*, element-by-element results are returned.

In performing the conversion, leading and trailing blanks are ignored: "1.5" and " 1.5 " both convert to (numeric) 1.5, but "1.5 kilometers" converts to . (missing). Use **strtoreal**(**tokens**(*S*)[1]) to convert just the first space-delimited part.

All Stata numeric formats are understood, such as 0, 1, −2, 1.5, 1.5e+2, and −1.0x+8, as well as the missing-value codes ., .a, .b, ..., .z.

Thus using **strtoreal**(*S*), if an element of *S* converts to . (missing), you cannot tell whether the element was valid and equal to "." or the element was invalid and so defaulted to . (missing), such as if *S* contained "cat" or "dog" or "1.5 kilometers".

When it is important to distinguish between these cases, use **\_strtoreal**(*S, R*). The conversion is returned in *R* and the function returns the number of elements that were invalid. If **\_strtoreal**() returns 0, then all values were valid.



## Conformability

```

strtoreal(S):
  input:
    S:       $r \times c$ 
  output:
    result:   $r \times c$ 

_strtoreal(S, R):
  input:
    S:       $r \times c$ 
  output:
    R:       $r \times c$ 
    result:   $1 \times 1$ 

```

## Diagnostics

`strtoreal(S)` returns a missing value wherever an element of *S* cannot be converted to a number.

`_strtoreal(S, R)` does the same, but the result is returned in *R*.

## Also see

[M-5] [strofreal\(\)](#) — Convert real to string

[M-4] [string](#) — String manipulation functions

## Syntax

*string matrix*    **stritrim**(*string matrix s*)

*string matrix*    **strltrim**(*string matrix s*)

*string matrix*    **strrtrim**(*string matrix s*)

*string matrix*    **strtrim**(*string matrix s*)

## Description

**stritrim**(*s*) returns *s* with all consecutive, internal blanks collapsed to one blank.

**strltrim**(*s*) returns *s* with leading blanks removed.

**strrtrim**(*s*) returns *s* with trailing blanks removed.

**strtrim**(*s*) returns *s* with leading and trailing blanks removed.

When *s* is not a scalar, these functions return element-by-element results.

## Remarks

Stata understands **stritrim()**, **strltrim()**, **strrtrim()**, and **strtrim()**, as synonyms for its own **itrim()**, **ltrim()**, **rtrim()**, and **trim()** functions, so you can use the **str\*()** names in both your Stata and Mata code.

## Conformability

**stritrim**(*s*), **strltrim**(*s*), **strrtrim**(*s*), **strtrim**(*s*):

<i>s</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

None.

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] **strupper()** — Convert string to uppercase (lowercase)

## Syntax

*string matrix*    **strupper**(*string matrix s*)

*string matrix*    **strlower**(*string matrix s*)

*string matrix*    **strproper**(*string matrix s*)

## Description

**strupper**(*s*) returns *s*, converted to uppercase.

**strlower**(*s*) returns *s*, converted to lowercase.

**strproper**(*s*) returns a string with the first letter capitalized and any other letters capitalized that immediately follow characters that are not letters; all other letters are converted to lowercase.

When *s* is not a scalar, these functions return element-by-element results.

## Remarks

**strproper**("mR. joHn a. sMitH") returns Mr. John A. Smith.

**strproper**("jack o'reilly") returns Jack O'Reilly.

**strproper**("2-cent's worth") returns 2-Cent'S Worth.

Also, Stata understands **strupper()**, **strlower()**, and **strproper()** as synonyms for its own **upper()**, **lower()**, and **proper()** functions, so you can use the **str\*()** names in both your Stata and Mata code.

## Conformability

**strupper**(*s*), **strlower**(*s*), **strproper**(*s*):

<i>s</i> :	$r \times c$
<i>result</i> :	$r \times c$

## Diagnostics

None.

## Also see

[M-4] **string** — String manipulation functions

# Title

[M-5] **substr()** — Substitute text

## Syntax

```
string matrix  substr(string matrix s, string matrix old, string matrix new)

string matrix  substr(string matrix s, string matrix old, string matrix new,
                      real matrix cnt)

string matrix  subinword(string matrix s, string matrix old, string matrix new)

string matrix  subinword(string matrix s, string matrix old, string matrix new,
                        real matrix cnt)
```

## Description

`substr(s, old, new)` returns *s* with all occurrences of *old* changed to *new*.

`substr(s, old, new, cnt)` returns *s* with the first *cnt* occurrences of *old* changed to *new*. All occurrences are changed if *cnt* contains missing.

`subinword(s, old, new)` returns *s* with all occurrences of *old* on word boundaries changed to *new*.

`subinword(s, old, new, cnt)` returns *s* with the first *cnt* occurrences of *old* on word boundaries changed to *new*. All occurrences are changed if *cnt* contains missing.

When arguments are not scalar, these functions return element-by-element results.

## Remarks

`substr("th thin man", "th", "the")` returns “the thein man”.

`subinword("th thin man", "th", "the")` returns “the thin man”.

## Conformability

```
substr(s, old, new, cnt), subinword(s, old, new, cnt):
  s:      r1 × c1
  old:    r2 × c2
  new:    r3 × c3
  cnt:    r4 × c4 (optional);  s, old, new, cnt r-conformable
  result: max(r1,r2,r3,r4) × max(c1,c2,c3,c4)
```

## Diagnostics

`substr(s, old, new, cnt)` and `subinword(s, old, new, cnt)` treat *cnt* < 0 as if *cnt* = 0 was specified; the original string *s* is returned.

---

## Also see

[M-4] [string](#) — String manipulation functions

# Title

[M-5] **sublowertriangle()** — Return a matrix with zeros above a diagonal

## Syntax

```
numeric matrix sublowertriangle(numeric matrix A [, numeric scalar p])
void           _sublowertriangle(numeric matrix A [, numeric scalar p])
```

where argument *p* is optional.

## Description

`sublowertriangle(A, p)` returns *A* with the elements above a diagonal set to zero. In the returned matrix,  $A[i, j] = 0$  for all  $i - j < p$ . If it is not specified, *p* is set to zero.

`_sublowertriangle()` mirrors `sublowertriangle()` but modifies *A*. `_sublowertriangle(A, p)` sets  $A[i, j] = 0$  for all  $i - j < p$ . If it is not specified, *p* is set to zero.

## Remarks

Remarks are presented under the following headings:

*Get lower triangle of a matrix*  
*Nonsquare matrices*

### Get lower triangle of a matrix

If *A* is a square matrix, then `sublowertriangle(A, 0) = lowertriangle(A)`. `sublowertriangle()` is a generalization of `lowertriangle()`.

We begin by defining *A*

```
: A = (1, 2, 3 \ 4, 5, 6 \ 7, 8, 9)
```

`sublowertriangle(A, 0)` returns *A* with zeros above the main diagonal as does `lowertriangle()`:

```
: sublowertriangle(A, 0)
      1  2  3
1      1  0  0
2      4  5  0
3      7  8  9
```

sublowertriangle(A, 1) returns A with zeros in the main diagonal and above.

```
: sublowertriangle(A, 1)
1  2  3
```

1	0	0	0
2	4	0	0
3	7	8	0

sublowertriangle(A, p) can take negative  $p$ . For example, setting  $p = -1$  yields

```
: sublowertriangle(A, -1)
```

	1	2	3
1	1	2	0
2	4	5	6
3	7	8	9

## Nonsquare matrices

sublowertriangle() and \_sublowertriangle() may be used with nonsquare matrices.

For instance, we define a nonsquare matrix A

```
: A = (1, 2, 3, 4 \ 5, 6, 7, 8 \ 9, 10, 11, 12)
```

We use sublowertriangle() to obtain the lower triangle of A:

```
: sublowertriangle(A, 0)
```

	1	2	3	4
1	1	0	0	0
2	5	6	0	0
3	9	10	11	0

## Conformability

sublowertriangle(A, p):

*input:*

A:  $r \times c$

p:  $1 \times 1$  (optional)

*output:*

result:  $r \times c$

\_sublowertriangle(A, p):

*input:*

A:  $r \times c$

p:  $1 \times 1$  (optional)

*output:*

A:  $r \times c$

## Diagnostics

None.

## Also see

[M-4] [manipulation](#) — Matrix manipulation



# Title

[M-5] `_substr()` — Substitute into string

## Syntax

*void* `_substr(string scalar s, string scalar tosub, real scalar pos)`

## Description

`_substr(s, tosub, pos)` substitutes *tosub* into *s* at position *pos*. The first position of *s* is *pos* = 1. `_substr()` may be used with text or binary strings.

Do not confuse `_substr()` with `substr()`, which extracts substrings; see [M-5] [substr\(\)](#).

## Remarks

If *s* contains “abcdef”, then `_substr(s, "XY", 2)` changes *s* to contain “aXYdef”.

## Conformability

`_substr(s, tosub, pos):`

<i>input:</i>	
<i>s:</i>	$1 \times 1$
<i>tosub:</i>	$1 \times 1$
<i>pos:</i>	$1 \times 1$
<i>output:</i>	
<i>s:</i>	$1 \times 1$

## Diagnostics

`_substr(s, tosub, pos)` does nothing if *tosub*==“”.

`_substr(s, tosub, pos)` may not be used to extend *s*: `_substr()` aborts with error if substituting *tosub* into *s* would result in a string longer than the original *s*. `_substr()` also aborts with error if *pos* ≤ 0 or *pos* ≥ . unless *tosub* is “”.

`_substr(s, tosub, pos)` aborts with error if *s* or *tosub* are views.

## Also see

[M-4] [string](#) — String manipulation functions

## Syntax

```
string matrix  substr(string matrix s, real matrix b, real matrix l)

string matrix  substr(string matrix s, real matrix b)
```

## Description

substr(*s*, *b*, *l*) returns the substring of *s* starting at position *b* and continuing for a length of *l*, where

1. *b* specifies the starting position; the first character of the string is *b* = 1.
2. *b* > 0 is interpreted as distance from the start of the string; *b* = 2 means starting at the second character.
3. *b* < 0 is interpreted as distance from the end of string; *b* = -1 means starting at the last character; *b* = -2 means starting at the second from the last character.
4. *l* specifies the length; *l* = 2 means for two characters.
5. *l* < 0 is treated the same as *l* = 0: no characters are copied.
6. *l* ≥ . is interpreted to mean to the end of the string.

substr(*s*, *b*) is equivalent to substr(*s*, *b*, .) for strings that do not contain binary 0. If there is a binary 0 to the right of *b*, the substring from *b* up to but not including the binary 0 is returned.

When arguments are not scalar, substr() returns element-by-element results.

## Conformability

```
substr(s, b, l):
    s:       $r_1 \times c_1$ 
    b:       $r_2 \times c_2$ 
    l:       $r_3 \times c_3$ ; s, b, and l r-conformable
    result:  $\max(r_1, r_2, r_3) \times \max(c_1, c_2, c_3)$ 

substr(s, b):
    s:       $r_1 \times c_1$ 
    b:       $r_2 \times c_2$ ; s and b r-conformable
    result:  $\max(r_1, r_2) \times \max(c_1, c_2)$ 
```

## Diagnostics

In substr(*s*, *b*, *l*) and substr(*s*, *b*), if *b* describes a position before the beginning of the string or after the end, "" is returned. If *b* + *l* describes a position to the right of the end of the string, results are as if a smaller value for *l* were specified.

---

## Also see

[M-4] [string](#) — String manipulation functions

## Syntax

<i>numeric colvector</i>	<code>rowsum(<i>numeric matrix Z</i> [ , <i>missing</i> ])</code>
<i>numeric rowvector</i>	<code>colsum(<i>numeric matrix Z</i> [ , <i>missing</i> ])</code>
<i>numeric scalar</i>	<code>sum(<i>numeric matrix Z</i> [ , <i>missing</i> ])</code>
<i>numeric colvector</i>	<code>quadrowsum(<i>numeric matrix Z</i> [ , <i>missing</i> ])</code>
<i>numeric rowvector</i>	<code>quadcolsum(<i>numeric matrix Z</i> [ , <i>missing</i> ])</code>
<i>numeric scalar</i>	<code>quadsum(<i>numeric matrix Z</i> [ , <i>missing</i> ])</code>

where optional argument *missing* is a real scalar that determines how missing values in *Z* are treated:

1. Specifying *missing* as 0 is equivalent to not specifying the argument; missing values in *Z* are treated as contributing 0 to the sum.
2. Specifying *missing* as 1 (or nonzero) specifies that missing values in *Z* are to be treated as missing values and to turn the sum to missing.

## Description

`rowsum(Z)` and `rowsum(Z, missing)` return a column vector containing the sum over the rows of *Z*.

`colsum(Z)` and `colsum(Z, missing)` return a row vector containing the sum over the columns of *Z*.

`sum(Z)` and `sum(Z, missing)` return a scalar containing the sum over the rows and columns of *Z*.

`quadrowsum()`, `quadcolsum()`, and `quadsum()` are quad-precision variants of the above functions. The sum is accumulated in quad precision and then rounded to double precision and returned.

Argument *missing* determines how missing values are treated. If *missing* is not specified, results are the same as if *missing* = 0 were specified: missing values are treated as zero. If *missing* = 1 is specified, missing values are treated as missing values.

These functions may be used with real or complex matrix *Z*.

## Remarks

All functions return the same type as the argument, real if argument is real, complex if complex.

## Conformability

`rowsum(Z, missing)`, `quadrowsum(Z, missing)`:

<i>Z</i> :	$r \times c$	
<i>missing</i> :	$1 \times 1$	(optional)
<i>result</i> :	$r \times 1$	

`colsum(Z, missing)`, `quadcolsum(Z, missing)`:

<i>Z</i> :	$r \times c$	
<i>missing</i> :	$1 \times 1$	(optional)
<i>result</i> :	$1 \times c$	

`sum(Z, missing)`, `quadsum(Z, missing)`:

<i>Z</i> :	$r \times c$	
<i>missing</i> :	$1 \times 1$	(optional)
<i>result</i> :	$1 \times 1$	

## Diagnostics

If *missing* = 0, missing values are treated as contributing zero to the sum; they do not turn the sum to missing. Otherwise, missing values turn the sum to missing.

## Also see

[M-5] [mean\(\)](#) — Means, variances, and correlations

[M-5] [runningsum\(\)](#) — Running sum of vector

[M-5] [cross\(\)](#) — Cross products

[M-4] [mathematical](#) — Important mathematical functions

[M-4] [utility](#) — Matrix utility functions

# Title

[M-5] **svd()** — Singular value decomposition

## Syntax

*void*                      **svd**(*numeric matrix A*, *U*, *s*, *Vt*)

*real colvector*      **svdsv**(*numeric matrix A*)

*void*                      **\_svd**(*numeric matrix A*, *s*, *Vt*)

*real colvector*      **\_svdsv**(*numeric matrix A*)

*real scalar*          **\_svd\_la**(*numeric matrix A*, *s*, *Vt*)

## Description

**svd**(*A*, *U*, *s*, *Vt*) calculates the singular value decomposition of *A*:  $m \times n$ ,  $m \geq n$ , returning the result in *U*, *s*, and *Vt*. Singular values returned in *s* are sorted from largest to smallest.

**svdsv**(*A*) returns the singular values of *A*:  $m \times n$ ,  $m \geq n$  or  $m < n$  (that is, no restriction), in a column vector of length  $\min(m,n)$ . *U* and *Vt* are not calculated.

**\_svd**(*A*, *s*, *Vt*) does the same as **svd**(), except that it returns *U* in *A*. Use of **\_svd**() conserves memory.

**\_svdsv**(*A*) does the same as **svdsv**(), except that, in the process, it destroys *A*. Use of **\_svdsv**() conserves memory.

**\_svd\_la**() is the interface into the [M-1] **LAPACK** SVD routines and is used in the implementation of the previous functions. There is no reason you should want to use it.

## Remarks

Remarks are presented under the following headings:

*[Introduction](#)*

*[Possibility of convergence problems](#)*

Documented here is the thin SVD, appropriate for use with *A*:  $m \times n$ ,  $m \geq n$ . See [M-5] **fullsvd**() for the full SVD, appropriate for use in all cases. The relationship between the two is discussed in *[Relationship between the full and thin SVDs](#)* in [M-5] **fullsvd**().

Use of the thin SVD—the functions documented here—is preferred when  $m \geq n$ .

## Introduction

The SVD is used to compute accurate solutions to linear systems and least-squares problems, to compute the 2-norm, and to determine the numerical rank of a matrix.

The singular value decomposition (SVD) of  $A: m \times n, m \geq n$ , is given by

$$A = U * \text{diag}(s) * V'$$

where

$$\begin{array}{ll} U: & m \times n \text{ and } U'U = I(n) \\ s: & n \times 1 \\ V: & n \times n \text{ and orthogonal (unitary)} \end{array}$$

When  $A$  is complex, the transpose operator  $'$  is understood to mean the [conjugate transpose](#) operator.

Vector  $s$  contains the singular values, and those values are real even when  $A$  is complex.  $s$  is ordered so that the largest singular value appears first, then the next largest, and so on.

Function `svd(A, U, s, Vt)` returns  $U$ ,  $s$ , and  $Vt = V'$ .

Function `svdsv(A)` returns  $s$ , omitting the calculation of  $U$  and  $Vt$ . Also, whereas `svd()` is suitable for use only in the case  $m \geq n$ , `svdsv()` may be used in all cases.

## Possibility of convergence problems

It is possible, although exceedingly unlikely, that the SVD routines could fail to converge. `svd()`, `svdsv()`, `_svd()`, and `_svdsv()` then return singular values in  $s$  equal to missing.

In coding, it is perfectly acceptable to ignore this possibility because (1) it is so unlikely and (2) even if the unlikely event occurs, the missing values will properly reflect the situation. If you do wish to check, in addition to checking `missing(s)>0` (see [\[M-5\] missing\(\)](#)), you must also check `missing(A)==0` because that is the other reason  $s$  could contain missing values. Convergence was not achieved if `missing(s) > 0 & missing(A)==0`. If you are calling one of the destructive-of- $A$  versions of SVD, remember to check `missing(A)==0` before extracting singular values.

## Conformability

`svd(A, U, s, Vt):`

*input:*

$$A: \quad m \times n, \quad m \geq n$$

*output:*

$$U: \quad m \times n$$

$$s: \quad n \times 1$$

$$Vt: \quad n \times n$$

```
svdsv(A):
    A:       $m \times n$ ,  $m \geq n$  or  $m < n$ 
    result:  $\min(m, n) \times 1$ 

_svd(A, s, Vt):
    input:
        A:       $m \times n$ ,  $m \geq n$ 
    output:
        A:       $m \times n$ , contains  $U$ 
        s:       $n \times 1$ 
        Vt:      $n \times n$ 

_svds(A):
    input:
        A:       $m \times n$ ,  $m \geq n$  or  $m < n$ 
    output:
        A:       $0 \times 0$ 
        result:  $\min(m, n) \times 1$ 

_svd_la(A, s, Vt):
    input:
        A:       $m \times n$ ,  $m \geq n$ 
    output:
        A:       $m \times n$ , contains  $U$ 
        s:       $n \times 1$ 
        Vt:      $n \times n$ 
    result:     $1 \times 1$ 
```

## Diagnostics

`svd(A, U, s, Vt)` and `_svd(A, s, Vt)` return missing results if *A* contains missing. In all other cases, the routines should work, but there is the unlikely possibility of convergence problems, in which case missing results will also be returned; see [Possibility of convergence problems](#) above.

`svdsv(A)` and `_svdsv(A)` return missing results if *A* contains missing values or if there are convergence problems.

`_svd()` and `_svdsv()` abort with error if *A* is a view.

Direct use of `_svd_la()` is not recommended.

Singular value decompositions have multiple roots, as is engagingly explained by [Stewart \(1993\)](#). Eugenio Beltrami (Italy, 1835–1900), Camille Jordan (France, 1838–1922), and James Joseph Sylvester (Britain, 1814–1897) came to them through what we now call linear algebra, while Erhard Schmidt (Germany, 1876–1959) and Hermann Klaus Hugo Weyl (Germany, 1885–1955) approached them from integral equations. Although none of them used the matrix terminology or notation that is familiar to modern workers, seeing the structure within sets of equations was a more familiar task to them than it was to us. The terminology “singular values” appears to come from the literature on integral equations. The use of SVDs as workhorses in modern numerical analysis owes most to Gene Howard Golub (United States, 1932–2007).



## References

- Stewart, G. W. 1993. On the early history of the singular value decomposition. *SIAM Review* 35: 551–566.
- Trefethen, L. N. 2007. Obituary: Gene H. Golub (1932–2007). *Nature* 450: 962.

## Also see

- [M-5] `fullsvd()` — Full singular value decomposition
- [M-5] `svsolve()` — Solve  $AX=B$  for  $X$  using singular value decomposition
- [M-5] `pinv()` — Moore–Penrose pseudoinverse
- [M-5] `norm()` — Matrix and vector norms
- [M-5] `rank()` — Rank of matrix
- [M-4] `matrix` — Matrix functions

## Syntax

```

numeric matrix    svsolve(A, B)
numeric matrix    svsolve(A, B, rank)
numeric matrix    svsolve(A, B, rank, tol)

real scalar       _svsolve(A, B)
real scalar       _svsolve(A, B, tol)
    
```

where

```

A:    numeric matrix
B:    numeric matrix
rank: irrelevant; real scalar returned
tol:  real scalar
    
```

## Description

`svsolve(A, B, ...)`, uses singular value decomposition to solve  $AX = B$  and return  $X$ . When  $A$  is singular, `svsolve()` computes the minimum-norm least-squares generalized solution. When *rank* is specified, in it is placed the rank of  $A$ .

`_svsolve(A, B, ...)` does the same thing, except that it destroys the contents of  $A$  and it overwrites  $B$  with the solution. Returned is the rank of  $A$ .

In both cases, *tol* specifies the tolerance for determining whether  $A$  is of full rank. *tol* is interpreted in the standard way—as a multiplier for the default if  $tol > 0$  is specified and as an absolute quantity to use in place of the default if  $tol \leq 0$  is specified.

## Remarks

`svsolve(A, B, ...)` is suitable for use with square or nonsquare, full-rank or rank-deficient matrix  $A$ . When  $A$  is of full rank, `qrsolve()` returns the same solution as `lusolve()` (see [\[M-5\] lusolve\(\)](#)), ignoring roundoff error. When  $A$  is singular, `svsolve()` returns the minimum-norm least-squares generalized solution. `qrsolve()` (see [\[M-5\] qrsolve\(\)](#)), an alternative, returns a generalized least-squares solution that amounts to dropping rows of  $A$ .

Remarks are presented under the following headings:

[Derivation](#)  
[Relationship to inversion](#)  
[Tolerance](#)

## Derivation

We wish to solve for  $X$

$$AX = B \tag{1}$$

Perform singular value decomposition on  $A$  so that we have  $A = USV'$ . Then (1) can be rewritten

$$USV'X = B$$

Premultiplying by  $U'$  and remembering that  $U'U = I$ , we have

$$SV'X = U'B$$

Matrix  $S$  is diagonal and thus its inverse is easily calculated, and we have

$$V'X = S^{-1}U'B$$

When we premultiply by  $V$ , remembering that  $VV' = I$ , the solution is

$$X = VS^{-1}U'B \tag{2}$$

See [M-5] `svd()` for more information on the SVD.

## Relationship to inversion

For a general discussion, see *Relationship to inversion* in [M-5] `lusolve()`.

For an inverse based on the SVD, see [M-5] `pinv()`. `pinv(A)` amounts to `svsolve(A, I(rows(A)))`, although `pinv()` has separate code that uses less memory.

## Tolerance

In (2) above, we are required to calculate the inverse of diagonal matrix  $S$ . The generalized solution is obtained by substituting zero for the  $i$ th diagonal element of  $S^{-1}$ , where the  $i$ th diagonal element of  $S$  is less than or equal to  $eta$  in absolute value. The default value of  $eta$  is

$$eta = \text{epsilon}(1) * \text{rows}(A) * \max(S)$$

If you specify  $tol > 0$ , the value you specify is used to multiply  $eta$ . You may instead specify  $tol \leq 0$  and then the negative of the value you specify is used in place of  $eta$ ; see [M-1] *tolerance*.

## Conformability

`svsolve(A, B, rank, tol):`

*input:*

*A:*  $m \times n$

*B:*  $m \times k$

*tol:*  $1 \times 1$  (optional)

*output:*

*rank:*  $1 \times 1$  (optional)

*result:*  $n \times k$

`_svsolve(A, B, tol):`

*input:*

*A:*  $m \times n$

*B:*  $m \times k$

*tol:*  $1 \times 1$  (optional)

*output:*

*A:*  $0 \times 0$

*B:*  $m \times k$

*result:*  $1 \times 1$

## Diagnostics

`svsolve(A, B, ...)` and `_svsolve(A, B, ...)` return missing results if  $A$  or  $B$  contain missing.

`_svsolve(A, B, ...)` aborts with error if  $A$  (but not  $B$ ) is a view.

## Also see

[M-5] [`solvelower\(\)`](#) — Solve  $AX=B$  for  $X$ ,  $A$  triangular

[M-5] [`cholsolve\(\)`](#) — Solve  $AX=B$  for  $X$  using Cholesky decomposition

[M-5] [`lusolve\(\)`](#) — Solve  $AX=B$  for  $X$  using LU decomposition

[M-5] [`qrsolve\(\)`](#) — Solve  $AX=B$  for  $X$  using QR decomposition

[M-4] [`matrix`](#) — Matrix functions

[M-4] [`solvers`](#) — Functions to solve  $AX=B$  and to obtain  $A$  inverse

# Title

[M-5] **swap()** — Interchange contents of variables

## Syntax

*void*   **swap**(*transmorphic matrix A*, *transmorphic matrix B*)

## Description

**swap**(*A*, *B*) interchanges the contents of *A* and *B*. *A* and *B* are not required to be of the same type or dimension.

## Remarks

There is no faster way than **swap**(*A*, *B*) to assign *A=B* when you do not care about the contents of *B* after the assignment. For instance, you have the code

*A* = *B*  
*B* = ...(*matrix expression*)...

Faster is

**swap**(*A*, *B*)  
*B* = ...(*matrix expression*)...

The execution time of **swap**() is independent of the size of *A* and *B*, and **swap**() conserves memory to boot. Pretend that *B* is a 900 × 900 matrix. After *A=B* is executed, but before *B* is reassigned, two copies of the 900 × 900 matrix exist. That does not happen with **swap**().

## Conformability

**swap**(*A*, *B*):

*input:*

*A:*     *r*<sub>1</sub> × *c*<sub>1</sub>  
*B:*     *r*<sub>2</sub> × *c*<sub>2</sub>

*output:*

*A:*     *r*<sub>2</sub> × *c*<sub>2</sub>  
*B:*     *r*<sub>1</sub> × *c*<sub>1</sub>

## Diagnostics

**swap**(*A*, *B*) works only with variables. Do not code, for instance, **swap**(*A*[*i*,*j*], *A*[*j*,*i*]). It is not an error, but it will have no effect.

## Also see

[M-4] [programming](#) — Programming functions

Title

[M-5] Toeplitz() — Toeplitz matrices

Syntax

*numeric matrix*    `Toeplitz(numeric colvector cI, numeric rowvector rI)`

Description

`Toeplitz(cI, rI)` returns the Toeplitz matrix defined by *cI* being its first column and *rI* being its first row. A Toeplitz matrix *T* is characterized by  $T[i, j] = T[i - 1, j - 1]$ ,  $i, j > 1$ . In a Toeplitz matrix, each diagonal is constant.

Vectors *cI* and *rI* specify the first column and first row of *T*.

Remarks

`cI[1]` is used to fill  $T[1, 1]$ , and `rI[1]` is not used.

To obtain the symmetric (Hermitian) Toeplitz matrix, code `Toeplitz(v, v')` (if *v* is a column vector), or `Toeplitz(v', v)` if *v* is a row vector.

Conformability

`Toeplitz(cI, rI)`:

<i>cI</i> :	$r \times 1$
<i>rI</i> :	$1 \times c$
<i>result</i> :	$r \times c$

Diagnostics

None.

Otto Toeplitz (1881–1940) was born in Breslau, Germany (now Wrocław, Poland), and educated there in mathematics. He researched and taught at universities in Göttingen, Kiel, and Bonn, making many contributions to algebra and analysis, but he was dismissed in 1935 for being a Jew. Toeplitz emigrated to Palestine in 1939 but died a few months later in Jerusalem. He was fascinated by the history of mathematics and wrote a popular work with Hans Rademacher, *The Enjoyment of Mathematics*.

Reference

Robinson, A. 1976. Toeplitz, Otto. In Vol. 13 of *Dictionary of Scientific Biography*, ed. C. C. Gillispie. New York: Scribner's.

## Also see

[M-4] [standard](#) — Functions to create standard matrices



## Syntax

```

t = tokeninit([wchars [, pchars [, qchars [, allownum [, allowhex]]]])
t = tokeninitstata()

void          tokenset(t, string scalar s)

string rowvector tokengetall(t)
string scalar  tokenget(t)
string scalar  tokenpeek(t)
string scalar  tokenrest(t)

real scalar    tokenoffset(t)
void           tokenoffset(t, real scalar offset)

string scalar  tokenwchars(t)
void           tokenwchars(t, string scalar wchars)

string rowvector tokenpchars(t)
void           tokenpchars(t, string rowvector pchars)

string rowvector tokenqchars(t)
void           tokenqchars(t, string rowvector qchars)

real scalar    tokenallownum(t)
void           tokenallownum(t, real scalar allownum)

real scalar    tokenallowhex(t)
void           tokenallowhex(t, real scalar allowhex)
    
```

where

*t* is *transmorphic* and contains the parsing environment information. You obtain a *t* from `tokeninit()` or `tokeninitstata()` and then pass *t* to the other functions.

*wchars* is a *string scalar* containing the characters to be treated as white space, such as " ", (" "+char(9)), or "".

*pchars* is a *string rowvector* containing the strings to be treated as parsing characters, such as "" and (">", "<", ">=", "<="). "" and `J(1,0,"")` are given the same interpretation: there are no parsing characters.

*qchars* is a *string rowvector* containing the character pairs to be treated as quote characters. "" (that is, empty string) is given the same interpretation as `J(1,0,"")`; there are no quote characters. *qchars* = (""""") (that is, the two-character string quote indicates that " is to be treated as open quote and " is to be treated as close quote. *qchars* = (""""", ""'""'") indicates that, in addition, ' is to be treated as open quote and ' as close quote. In a syntax that did not use < and > as parsing characters, *qchars* = ("<>") would indicate that < is to be treated as open quote and > as close quote.

*allownum* is a *string scalar* containing 0 or 1. *allownum* = 1 indicates that numbers such as 12.23 and 1.52e+02 are to be returned as single tokens even in violation of other parsing rules.

*allowhex* is a *string scalar* containing 0 or 1. *allowhex* = 1 indicates that numbers such as 1.921fb54442d18X+001 and 1.0x+a are to be returned as single tokens even in violation of other parsing rules.

## Description

These functions provide advanced parsing. If you simply wish to convert strings into row vectors by separating on blanks, converting "mpg weight displ" into ("mpg", "weight", "displ"), see [M-5] [tokens\(\)](#).

## Remarks

Remarks are presented under the following headings:

### Concepts

[White-space characters](#)

[Parsing characters](#)

[Quote characters](#)

[Overrides](#)

[Setting the environment to parse on blanks with quote binding](#)

[Setting the environment to parse full Stata syntax](#)

[Setting the environment to parse tab-delimited files](#)

### Function overview

[tokeninit\(\)](#) and [tokeninitstata\(\)](#)

[tokenset\(\)](#)

[tokengetall\(\)](#)

[tokenget\(\)](#), [tokenpeek\(\)](#), and [tokenrest\(\)](#)

[tokenoffset\(\)](#)

[tokenwchars\(\)](#), [tokenpchars\(\)](#), and [tokenqchars\(\)](#)

[tokenallownum](#) and [tokenallowhex\(\)](#)

## Concepts

Parsing refers to splitting a string into pieces, which we will call tokens. Parsing as implemented by the `token*()` functions is defined by (1) the white-space characters *wchars*, (2) the parsing characters *pchars*, and (3) the quote characters *qchars*.

### White-space characters

Consider the string "this that what". If there are no white-space characters, no parsing characters, and no quote characters, that is, if *wchars* = *pchars* = *qchars* = "", then the result of parsing "this that what" would be one token that would be the string just as it is: "this that what".

If *wchars* were instead " ", then parsing "this that what" results in ("this", "that", "what"). Parsing "this that    what" (note the multiple blanks) would result in the same thing. White-space characters separate one token from the next but are not otherwise significant.

## Parsing characters

If we instead left *wchars* = "" and set *pchars* = " ", "this that what" parses into ("this", " ", "that", " ", "what") and parsing "this that    what" results in ("this", " ", "that", " ", " ", " ", "what").

*pchars* are like *wchars* except that they are themselves significant.

*pchars* do not usually contain space. A more reasonable definition of *pchars* is ("+", "-"). Then parsing "x+y" results in ("x", "+", "y"). Also, the parsing characters can be character combinations. If *pchars* = ("+", "-", "++", "--"), then parsing "x+y++" results in ("x", "+", "y", "++") and parsing "x+++y" results in ("x", "++", "+", "y"). Longer *pchars* are matched before shorter ones regardless of the order in which they appear in the *pchars* vector.

## Quote characters

*qchars* specifies the quote characters. Pieces of the string being parsed that are surrounded by quotes are returned as one token, ignoring the separation that would usually occur because of the *wchars* and *pchars* definitions. Consider the string

```
mystr= "x = y"
```

Let *wchars* = " " and *pchars* include "=". That by itself would result in the above string parsing into the five tokens

mystr	=	"x	=	y"
-------	---	----	---	----

Now let *qchars* = ('"', "'"); that is, *qchars* is the two-character string "". Parsing then results in the three tokens

mystr	=	"x = y"
-------	---	---------

Each element of *qchars* contains a character pair: the open character followed by the close character. We defined those two characters as " and ' above, that is, as being the same. The two characters can differ. We might define the first as ‘ and the second as ’. When the characters are different, quotations can nest. The quotation "he said "hello"" makes no sense because that parses into ("he said ", hello, ""). The quotation ‘he said ‘hello’’, however, makes perfect sense and results in the single token ‘he said ‘hello’’.

The quote characters can themselves be multiple characters. You can define open quote as ‘" and close as "': *qchars* = ('"', "'"). Or you can define multiple sets of quotation characters, such as *qchars* = ('"', "'"), ('"', "'").

The quote characters do not even have to be quotes at all. In some context you might find it convenient to specify them as ("()"). With that definition, "(2 × (3 + 2))" would parse into one token. Specifying them like this can be useful, but in general we recommend against it. It is usually better to write your code so that quote characters really are quote characters and to push the work of handling other kinds of nested expressions back onto the caller.

## Overrides

The `token*()` functions provide two overrides: *allownum* and *allowhex*. These have to do with parsing numbers. First, consider life without overrides. You have set `wchars = " "` and `pchars = ("=", "+", "-", "*", "/")`. You attempt to parse

```
y = x + 1e+13
```

The result is

y	=	x	+	1e	+	13
---	---	---	---	----	---	----

when what you wanted was

y	=	x	+	1e+13
---	---	---	---	-------

Setting *allownum* = 1 will achieve the desired result. *allownum* specifies that, when a token could be interpreted as a number, the number interpretation is to be taken even in violation of the other parsing rules.

Setting *allownum* = 1 will not find numbers buried in the middle of strings, such as the `1e+3` in `"xis1e+3"`, but if the number occurs at the beginning of the token according to the parsing rules set by *wchars* and *pchars*, *allownum* = 1 will continue the token in violation of those rules if that results in a valid number.

The override *allowhex* is similar and Stata specific. Stata (and Mata) provide a unique and useful way of writing hexadecimal floating-point numbers in a printable, short, and precise way:  $\pi$  can be written `1.921fb54442d18X+001`. Setting *allowhex* = 1 allows such numbers.

## Setting the environment to parse on blanks with quote binding

Stata's default rule for parsing do-file arguments is "parse on blanks and bind on quotes". The settings for duplicating that behavior are

```
wchars = " "
pchars = ( " " )
qchars = ( ' ' ' ' ' ' ' ' , ' " ' ' ' ' ' ' ' ' )
allownum = 0
allowhex = 0
```

This behavior can be obtained by coding

```
t = tokeninit(" ", " ", (' ' ' ' ' ' ' ' , ' " ' ' ' ' ' ' ' ' ), 0, 0)
```

or by coding

```
t = tokeninit()
```

because in `tokeninit()` the arguments are optional and "parse on blank with quote binding" is the default.

With those settings, parsing `'"first second "third fourth" fifth"'` results in `("first", "second", '"third fourth"', "fifth")`.

This result is a little different from that of Stata because the third token includes the quote binding characters. Assume that the parsed string was obtained by coding

```
res = tokengetall(r)
```

The following code will remove the open and close quotes, should that be desirable.

```
for (i=1; i<=cols(res); i++) {
    if (res[i]=='"') {
        res[i] = substr(res[i], 2, strlen(res[i])-2)
    }
    else if (substr(res[i], 1, 2)=="' + '"') {
        res[i] = substr(res[i], 3, strlen(res[i])-4)
    }
}
```

## Setting the environment to parse full Stata syntax

To parse full Stata syntax, the settings are

```
wchars = " "
pchars = ( "\", "~", "!", "=", ":", ";", ",",
            "?", "!", "@", "#", "==", "!=>=",
            "<=", "<", ">", "&", "|", "&&", "||",
            "+", "-", "++", "--", "*", "/", "^",
            "(", ")", "[", "]", "{", "}" )
qchars = ( '""', '""', char(96)+char(39))
allownum = 1
allowhex = 1
```

The above is a slight oversimplification. Stata is an interpretive language and Stata does not require users to type filenames in quotes, although Stata does allow it. Thus `"\"` is sometimes a parsing character and sometimes not, and the same is true of `"/`. As Stata parses a line from left to right, it will change *pchars* between two `tokenget()` calls when the next token could be or is known to be a filename. Sometimes Stata peeks ahead to decide which way to parse. You can do the same by using the `tokenpchars()` and `tokenpeek()` functions.

To obtain the above environment, code

```
t = tokeninitstata()
```

## Setting the environment to parse tab-delimited files

The `token*()` functions can be used to parse lines from tab-delimited files. A tab-delimited file contains lines of the form

```
<field1><tab><field2><tab><field3>
```

The parsing environment variables are

```
wchars = ""  
pchars = ( char(9) ) (i.e., tab)  
qchars = ( "" )  
allownum = 0  
allowhex = 0
```

To set this environment, code

```
t = tokeninit("", char(9), "", 0, 0)
```

Say that you then parse the line

```
Farber, William<tab> 2201.00<tab>12
```

The results will be

```
("Farber, William", char(9), " 2201.00", char(9), "12")
```

If the line were

```
Farber, William<tab><tab>12
```

the result would be

```
("Farber, William", char(9), char(9), "12")
```

The tab-delimited format is not well defined when the missing fields occur at the end of the line. A line with the last field missing might be recorded

```
Farber, William<tab> 2201.00<tab>
```

or

```
Farber, William<tab> 2201.00
```

A line with the last two fields missing might be recorded

```
Farber, William<tab><tab>
```

or

```
Farber, William<tab>
```

or

```
Farber, William
```

The following program would correctly parse lines with missing fields regardless of how they are recorded:

```

real rowvector readtabbed(transmorphic t, real scalar n)
{
    real scalar      i
    string rowvector res
    string scalar    token
    res = J(1, n, "")
    i = 1
    while ((token = tokenget(t))!="") {
        if (token==char(9)) i++
        else res[i] = token
    }
    return(res)
}

```

## Function overview

The basic way to proceed is to initialize the parsing environment and store it in a variable,

```
t = tokeninit(...)
```

and then set the string *s* to be parsed,

```
tokenset(t, s)
```

and finally use `tokenget()` to obtain the tokens one at a time (`tokenget()` returns "" when the end of the line is reached), or obtain all the tokens at once using `tokengetall(t)`. That is, either

```

while((token = tokenget(t)) != "") {
    ... process token ...
}

```

or

```

tokens = tokengetall(t)
for (i=1; i<=cols(tokens); i++) {
    ... process tokens[i] ...
}

```

After that, set the next string to be parsed,

```
tokenset(t, nextstring)
```

and repeat.

### tokeninit() and tokeninitstata()

`tokeninit()` and `tokeninitstata()` are alternatives. `tokeninitstata()` is generally unnecessary unless you are writing a fairly complicated function.

Whichever function you use, code

```
t = tokeninit(...)
```

or

```
t = tokeninitstata()
```

If you declare *t*, declare it `transmorphic`. *t* is in fact a structure containing all the details of your parsing environment, but that is purposely hidden from you so that you cannot accidentally modify the environment.

`tokeninit()` allows up to five arguments:

```
t = tokeninit(wchars, pchars, qchars, allownum, allowhex)
```

You may omit arguments from the end. If omitted, the default values of the arguments are

```
allowhex = 0
```

```
allownum = 0
```

```
qchars = ( "''''''", "''''''''" )
```

```
pchars = ( "" )
```

```
wchars = " "
```

## Notes

### 1. Concerning *wchars*:

- wchars* is a *string scalar*. The white-space characters appear one after the other in the string. The order in which the characters appear is irrelevant.
- Specify *wchars* as " " to treat blank as white space.
- Specify *wchars* as " "+char(9) to treat blank and *tab* as white space. Including *tab* is necessary only when strings to be parsed are obtained from a file; strings obtained from Stata already have the *tab* characters removed.
- Any character can be treated as a white-space character, including letters.
- Specify *wchars* as "" to specify that there are no white-space characters.

### 2. Concerning *pchars*:

- pchars* is a *string rowvector*. Each element of the vector is a separate parse character. The order in which the parse characters are specified is irrelevant.
- Specify *pchars* as ("+", "-") to make + and - parse characters.
- Parse characters may be character combinations such as ++ or >=. Character combinations may be up to four characters long.
- Specify *pchars* as "" or J(1,0,"") to specify that there are no parse characters. It makes no difference which you specify, but you will realize that J(1,0,"") is more logically consistent if you think about it.



### 3. Concerning *qchars*:

- a. *qchars* is a **string** rowvector. Each element of the vector contains the open followed by the close characters. The order in which sets of quote characters are specified is irrelevant.
- b. Specify *qchars* as ('""') to make " an open and close character.
- c. Specify *qchars* as ('""', '""') to make " and ' quote characters.
- d. Individual quote characters can be up to two characters long.
- e. Specify *qchars* as "" or J(1,0,"") to specify that there are no quote characters.

## tokenset()

After `tokeninit()` or `tokeninitstata()`, you are not yet through with initialization. You must `tokenset(s)` to specify the string scalar you wish to parse. You `tokenset()` one line, parse it, and if you have more lines, you `tokenset()` again and repeat the process. Often you will need to parse only one line. Perhaps you wish to write a program to parse the argument of a complicated option in a Stata ado-file. The structure is

```

program ...
    ...
    syntax ... [, ... MYoption(string) ...]
    mata: parseoption('"'myoption'"')
    ...
end

mata:
void parseoption(string scalar option)
{
    transmorphic      t
    t = tokeninit(...)
    tokenset(t, option)
    ...
}
end

```

### Notes

1. When you `tokenset(s)`, the contents of *s* are not stored. Instead, a pointer to *s* is stored. This approach saves memory and time, but it means that if you change *s* after setting it, you will change the subsequent behavior of the `token*()` functions.
2. Simply changing *s* is not sufficient to restart parsing. If you change *s*, you must `tokenset(s)` again.

## tokengetall()

You have two alternatives in how to process the tokens. You can parse the entire line into a row vector containing all the individual tokens by using `tokengetall()`,

```
tokens = tokengetall(t)
```

or you can use `tokenget()` to process the tokens one at a time, which is discussed in the next section.

Using `tokengetall()`, `tokens[1]` will be the first token, `tokens[2]` the second, and so on. There are, in total, `cols(tokens)` tokens. If the line was empty or contained only white-space characters, `cols(tokens)` will be 0.

### **tokenget(), tokenpeek(), and tokenrest()**

`tokenget()` returns the tokens one at a time and returns "" when the end of the line is reached. The basic loop for processing all the tokens in a line is

```
while ( (token = tokenget(t)) != "") {
    ...
}
```

`tokenpeek()` allows you to peek ahead at the next token without actually getting it, so whatever is returned will be returned again by the next call to `tokenget()`. `tokenpeek()` is suitable only for obtaining the next token after `tokenget()`. Calling `tokenpeek()` twice in a row will not return the next two tokens; it will return the next token twice. To obtain the next two tokens, code

```
...
current = tokenget(t)           // get the current token
...
t2 = t                         // copy parse environment
next_1 = tokenget(t2)           // peek at next token
next_1 = tokenget(t2)           // peek at token after that
...
current = tokenget(t)           // get next token
```

If you declare `t2`, declare it `transmorphic`.

`tokenrest()` returns the unparsed portion of the `tokenset()` string. Assume that you have just gotten the first token by using `tokenget()`. `tokenrest()` would return the rest of the original string, following the first token, unparsed. `tokenrest(t)` returns `substr(original_string, tokenoffset(t), .)`.

### **tokenoffset()**

`tokenoffset()` is useful only when you are using the `tokenget()` rather than `tokengetall()` style of programming. Let the original string you `tokenset()` be "this is an example". Right after you have `tokenset()` this string, `tokenoffset()` is 1:

```
    this is an example
    |
tokenoffset() = 1
```

After getting the first token (say it is "this"), `tokenoffset()` is 5:

```
    this is an example
    |
tokenoffset() = 5
```

`tokenoffset()` is always located on the first character following the last character parsed.

The syntax of `tokenoffset()` is

```
tokenoffset(t)
```

and

```
tokenoffset(t, newoffset)
```

The first returns the current offset value. The second resets the parser's location within the string.

### **tokenwchars(), tokenpchars(), and tokenqchars()**

`tokenwchars()`, `tokenpchars()`, and `tokenqchars()` allow resetting the current *wchars*, *pchars*, and *qchars*. As with `tokenoffset()`, they come in two syntaxes.

With one argument, *t*, they return the current value of the setting. With two arguments, *t* and *newvalue*, they reset the value.

Resetting in the midst of parsing is an advanced issue. The most useful of these functions is `tokenpchars()`, since for interactive grammars, it is sometimes necessary to switch on and off a certain parsing character such as `/`, which in one context means division and in another is a file separator.

### **tokenallownum and tokenallowhex()**

These two functions allow obtaining the current values of *allownum* and *allowhex* and resetting them.

## **Conformability**

`tokeninit(wchars, pchars, qchars, allownum, allowhex):`

<i>wchars</i> :	$1 \times 1$	(optional)
<i>pchars</i> :	$1 \times c_p$	(optional)
<i>qchars</i> :	$1 \times c_q$	(optional)
<i>allownum</i> :	$1 \times 1$	(optional)
<i>allowhex</i> :	$1 \times 1$	(optional)
<i>result</i> :	<i>transmorphic</i>	

`tokeninitstata():`

<i>result</i> :	<i>transmorphic</i>
-----------------	---------------------

`tokenset(t, s):`

<i>t</i> :	<i>transmorphic</i>
<i>s</i> :	$1 \times 1$
<i>result</i> :	<i>void</i>

`tokengetall(t):`

<i>t</i> :	<i>transmorphic</i>
<i>result</i> :	$1 \times k$

`tokenget(t), tokenpeek(t), tokenrest(t):`

<i>t</i> :	<i>transmorphic</i>
<i>result</i> :	$1 \times 1$

`tokenoffset(t), tokenwchars(t), tokenallownum(t), tokenallowhex(t):`

*t:* *transmorphic*

*result:*  $1 \times 1$

`tokenoffset(t, newvalue), tokenwchars(t, newvalue),  
tokenallownum(t, newvalue), tokenallowhex(t, newvalue):`

*t:* *transmorphic*

*newvalue:*  $1 \times 1$

*result:* *void*

`tokenpchars(t), tokenqchars(t):`

*t:* *transmorphic*

*result:*  $1 \times c$

`tokenpchars(t, newvalue), tokenqchars(t, newvalue):`

*t:* *transmorphic*

*newvalue:*  $1 \times c$

*result:* *void*

## Diagnostics

None.

## Also see

[M-5] [tokens\(\)](#) — Obtain tokens from string

[M-4] [programming](#) — Programming functions

[M-4] [string](#) — String manipulation functions

# Title

[M-5] **tokens()** — Obtain tokens from string

## Syntax

*string rowvector*    **tokens**(*string scalar s*)

*string rowvector*    **tokens**(*string scalar s*, *string scalar parsechars*)

## Description

**tokens**(*s*) returns the contents of *s*, split into words.

**tokens**(*s*, *parsechars*) returns the contents of *s* split into tokens based on *parsechars*.

**tokens**(*s*) is equivalent to **tokens**(*s*, " ").

If you need more advanced parsing, see [M-5] **tokenget**( ).

## Remarks

**tokens**( ) is commonly used to split a string containing a sequence of variable names into a row vector, each element of which contains one variable name:

```
tokens("mpg weight displacement") = ("mpg", "weight", "displacement")
```

Some Stata interface functions require that variable names be specified in this form. This is required, for instance, by **st\_varindex**( ); see [M-5] **st\_varindex**( ). If you had a string scalar *vars* containing one or more variable names, you could obtain their variable indices by coding

```
indices = st_varindex(tokens(vars))
```

## Conformability

**tokens**(*s*, *parsechars*)

<i>s</i> :	$1 \times 1$	
<i>parsechars</i> :	$1 \times 1$	(optional)
<i>result</i> :	$1 \times w$ ,	$w = \text{number of words (tokens) in } s$

## Diagnostics

If *s* contains "", **tokens**( ) returns J(1,0,"").

If *s* contains double-quoted or compound-double-quoted material, the quotes are stripped and that material is returned as one token. For example,

```
tokens('"this "is an" example"') = ("this", "is an", "example")
```

If *s* contains quoted material and the quotes do not match, results are as if the appropriate number of close quotes were added to the end of *s*. For example,

```
tokens('"this "is an example"') = ("this", "is an example")
```

## Also see

[M-5] [`invtokens\(\)`](#) — Concatenate string rowvector into string scalar

[M-5] [`tokenget\(\)`](#) — Advanced parsing

[M-4] [string](#) — String manipulation functions

# Title

[M-5] `trace()` — Trace of square matrix

## Syntax

*numeric scalar*    `trace(numeric matrix A)`

*numeric scalar*    `trace(numeric matrix A, numeric matrix B)`

*numeric scalar*    `trace(numeric matrix A, numeric matrix B, real scalar t)`

## Description

`trace(A)` returns the sum of the diagonal elements of  $A$ . Returned result is real if  $A$  is real, complex if  $A$  is complex.

`trace(A, B)` returns  $\text{trace}(AB)$ , the calculation being made without calculating or storing the off-diagonal elements of  $AB$ . Returned result is real if  $A$  and  $B$  are real and is complex otherwise.

`trace(A, B, t)` returns  $\text{trace}(AB)$  if  $t = 0$  and returns  $\text{trace}(A'B)$  otherwise, where, if either  $A$  or  $B$  is complex, transpose is understood to mean [conjugate transpose](#). Returned result is real if  $A$  and  $B$  are real and is complex otherwise.

## Remarks

`trace(A, B)` returns the same result as `trace(A*B)` but is more efficient if you do not otherwise need to calculate  $A*B$ .

`trace(A, B, 1)` returns the same result as `trace(A'B)` but is more efficient.

For real matrices  $A$  and  $B$ ,

$$\text{trace}(A') = \text{trace}(A)$$

$$\text{trace}(AB) = \text{trace}(BA)$$

and for complex matrices,

$$\text{trace}(A') = \text{conj}(\text{trace}(A))$$

$$\text{trace}(AB) = \text{trace}(BA)$$

where, for complex matrices, transpose is understood to mean conjugate transpose.

Thus for real matrices,

To calculate	Code
$\text{trace}(AB)$	<code>trace(A, B)</code>
$\text{trace}(A'B)$	<code>trace(A, B, 1)</code>
$\text{trace}(AB')$	<code>trace(A, B, 1)</code>
$\text{trace}(A'B')$	<code>trace(A, B)</code>

and for complex matrices,

To calculate	Code
$\text{trace}(AB)$	<code>trace(A, B)</code>
$\text{trace}(A'B)$	<code>trace(A, B, 1)</code>
$\text{trace}(AB')$	<code>conj(trace(A, B, 1))</code>
$\text{trace}(A'B')$	<code>conj(trace(A, B))</code>

Transpose in the first column means conjugate transpose.

## Conformability

<code>trace(A):</code>	
<i>A:</i>	$n \times n$
<i>result:</i>	$1 \times 1$
<code>trace(A, B):</code>	
<i>A:</i>	$n \times m$
<i>B:</i>	$m \times n$
<i>result:</i>	$1 \times 1$
<code>trace(A, B, t)</code>	
<i>A:</i>	$n \times m$ if $t = 0$ , $m \times n$ otherwise
<i>B:</i>	$m \times n$
<i>t:</i>	$1 \times 1$
<i>result:</i>	$1 \times 1$

## Diagnostics

- `trace(A)` aborts with error if  $A$  is not square.
- `trace(A, B)` and `trace(A, B, t)` abort with error if the matrices are not conformable or their product is not square.
- The trace of a  $0 \times 0$  matrix is 0.



## Also see

[M-4] [matrix](#) — Matrix functions

# Title

[M-5] `_transpose()` — Transposition in place

## Syntax

```
void _transpose(numeric matrix A)
```

## Description

`_transpose(A)` replaces  $A$  with  $A'$ . Coding `_transpose(A)` is equivalent to coding  $A = A'$ , except that execution can take a little longer and less memory is used. When  $A$  is complex,  $A$  is replaced with its conjugate transpose; see [M-5] [transposeonly\(\)](#) if transposition without conjugation is desired.

## Remarks

In some calculation, you need  $A'$

$X = \dots$  *calculation using  $A'$*   $\dots$

If  $A$  is large, you can save considerable memory by coding

```
_transpose(A)
X = ... calculation using A ...
_transpose(A)
```

## Conformability

`_transpose(A)`:

*input:*

$A:$        $r \times c$

*output:*

$A:$        $c \times r$

## Diagnostics

`_transpose(A)` aborts with error if  $A$  is a view.

## Also see

[M-2] [op\\_transpose](#) — Conjugate transpose operator

[M-5] [transposeonly\(\)](#) — Transposition without conjugation

[M-5] [conj\(\)](#) — Complex conjugate

[M-4] [manipulation](#) — Matrix manipulation

# Title

[M-5] `transposeonly()` — Transposition without conjugation

## Syntax

```
numeric matrix    transposeonly(numeric matrix A)  
void               _transposeonly(numeric matrix A)
```

## Description

`transposeonly(A)` returns  $A$  with its rows and columns interchanged. When  $A$  is real, the actions of `transposeonly(A)` are indistinguishable from coding  $A'$ ; see [M-2] [op\\_transpose](#). The returned result is the same, and the execution time is the same, too. When  $A$  is complex, however, `transposeonly(A)` is equivalent to coding `conj(A')`, but `transposeonly()` obtains the result more quickly.

`_transposeonly(A)` interchanges the rows and columns of  $A$  in place—without use of additional memory—and returns the transposed (but not conjugated) result in  $A$ .

## Remarks

`transposeonly()` is useful when you are coding in the programming, rather than the mathematical, sense. Say that you have two row vectors,  $a$  and  $b$ , and you want to place the two vectors together in a matrix  $R$ , and you want to turn them into column vectors. If  $a$  and  $b$  were certain to be real, you could just code

```
R = (a', b')
```

The above line, however, would result in not just the organization but also the values recorded in  $R$  changing if  $a$  or  $b$  were complex. The solution is to code

```
R = (transposeonly(a), transposeonly(b))
```

The above line will work for real or complex  $a$  and  $b$ . If you were concerned about memory consumption, you could instead code

```
R = (a \ b)  
_transposeonly(R)
```

## Conformability

`transposeonly(A)`:

```
  A:       $r \times c$   
result:    $c \times r$ 
```

`_transposeonly(A)`:

*input:*

$A:$        $r \times c$

*output:*

$A:$        $c \times r$

## Diagnostics

`_transposeonly(A)` aborts with error if  $A$  is a view.

## Also see

[M-2] [op\\_transpose](#) — Conjugate transpose operator

[M-5] [\\_transpose\(\)](#) — Transposition in place

[M-4] [manipulation](#) — Matrix manipulation

## Syntax

*real matrix*    **trunc**(*real matrix R*)

*real matrix*    **floor**(*real matrix R*)

*real matrix*    **ceil**(*real matrix R*)

*real matrix*    **round**(*real matrix R*)

*real matrix*    **round**(*real matrix R*, *real matrix U*)

## Description

These functions convert noninteger values to integers by moving toward 0, moving down, moving up, or rounding. These functions are typically used with scalar arguments, and they return a scalar in that case. When used with vectors or matrices, the operation is performed element by element.

**trunc**( $R$ ) returns the integer part of  $R$ .

**floor**( $R$ ) returns the largest integer  $i$  such that  $i \leq R$ .

**ceil**( $R$ ) returns the smallest integer  $i$  such that  $i \geq R$ .

**round**( $R$ ) returns the integer closest to  $R$ .

**round**( $R$ ,  $U$ ) returns the values of  $R$  rounded in units of  $U$  and is equivalent to **round**(( $R$ :/ $U$ ))\* $U$ . For instance, **round**( $R$ , 2) returns  $R$  rounded to the closest even number. **round**( $R$ , .5) returns  $R$  rounded to the closest multiple of one half. **round**( $R$ , 1) returns  $R$  rounded to the closest integer and so is equivalent to **round**( $R$ ).

## Remarks

Remarks are presented under the following headings:

[Relationship to Stata's functions](#)

[Examples of rounding](#)

## Relationship to Stata's functions

**trunc**() is equivalent to Stata's **int**() function.

**floor**(), **ceil**(), and **round**() are equivalent to Stata's functions of the same name.

## Examples of rounding

<i>x</i>	trunc( <i>x</i> )	floor( <i>x</i> )	ceil( <i>x</i> )	round( <i>x</i> )
1	1	1	1	1
1.3	1	1	2	1
1.6	1	1	2	2
−1	−1	−1	−1	−1
−1.3	−1	−2	−1	−1
−1.6	−1	−2	−1	−2

## Conformability

trunc(*R*), floor(*R*), ceil(*R*):

*R*:  $r \times c$   
*result*:  $r \times c$

round(*R*):

*R*:  $r \times c$   
*result*:  $r \times c$

round(*R*, *U*):

*R*:  $r_1 \times c_1$   
*U*:  $r_2 \times c_2$ , *R* and *U* r-conformable  
*result*:  $\max(r_1, r_2) \times \max(c_1, c_2)$

## Diagnostics

Most Stata and Mata functions return missing when arguments contain missing, and in particular, return . whether the argument is ., .a, .b, ..., .z. The logic is that performing the operation on a missing value always results in the same missing-value result. For example, sqrt(.a)==.

These functions, however, when passed a missing value, return the particular missing value. Thus trunc(.a)==.a, floor(.b)==.b, ceil(.c)==.c, and round(.d)==.d.

For round() with two arguments, this applies to the first argument and only when the second argument is not missing. If the second argument is missing (whether ., .a, ..., or .z), then . is returned.

## Also see

[M-4] [scalar](#) — Scalar mathematical functions

# Title

[M-5] **uniqrows()** — Obtain sorted, unique values

## Syntax

*transmorphic matrix*    **uniqrows**(*transmorphic matrix P*)

## Description

**uniqrows**(*P*) returns a sorted matrix containing the unique rows of *P*.

## Remarks

```
: x
      1  2  3
1      4  5  7
2      4  5  6
3      1  2  3
4      4  5  6

: uniqrows(x)
      1  2  3
1      1  2  3
2      4  5  6
3      4  5  7
```

## Conformability

**uniqrows**(*P*)

*P*:             $r_1 \times c_1$

*result*:        $r_2 \times c_1, \quad r_2 \leq r_1$

## Diagnostics

In **uniqrows**(*P*), if **rows**(*P*)==0, **J**(0, **cols**(*P*), **missingof**(*P*)) is returned.

If **rows**(*P*)>0 and **cols**(*P*)==0, **J**(1, 0, **missingof**(*P*)) is returned.

## Also see

- [M-5] **sort()** — Reorder rows of matrix
- [M-4] **manipulation** — Matrix manipulation

# Title

[M-5] **unitcircle()** — Complex vector containing unit circle

## Syntax

*complex colvector*    **unitcircle**(*real scalar n*)

## Description

**unitcircle**(*n*) returns a column vector containing  $C(\cos(\theta), \sin(\theta))$  for  $0 \leq \theta \leq 2\pi$  in *n* points.

## Conformability

**unitcircle**(*n*):

<i>n</i> :	$1 \times 1$
<i>result</i> :	$n \times 1$

## Diagnostics

None.

## Also see

[M-4] **standard** — Functions to create standard matrices



# Title

[M-5] **unlink()** — Erase file

## Syntax

```
void          unlink(real scalar filename)
real scalar _unlink(real scalar filename)
```

## Description

`unlink(filename)` erases *filename* if it exists, does nothing if *filename* does not exist, and aborts with error if *filename* exists but cannot be erased.

`_unlink(filename)` does the same, except that, if *filename* cannot be erased, rather than aborting with error, `_unlink()` returns a negative error code. `_unlink()` returns 0 if *filename* was erased or *filename* did not exist.

## Remarks

To remove directories, see `rmdir()` in [M-5] [chdir\(\)](#).

## Conformability

```
unlink(filename)
  filename:    1 × 1
  result:     void
```

```
_unlink(filename)
  filename:    1 × 1
  result:     1 × 1
```

## Diagnostics

`unlink(filename)` aborts with error when `_unlink()` would give a negative result.

`_unlink(filename)` returns a negative result if the file cannot be erased and returns 0 otherwise. If the file did not exist, 0 is returned. When there is an error, most commonly returned are -3602 (filename invalid) or -3621 (file is read-only).

## Also see

[M-4] [io](#) — I/O functions

# Title

**[M-5] valofexternal()** — Obtain value of external global

## Syntax

*transmorphic matrix*   `valofexternal(string scalar name)`

## Description

`valofexternal(name)` returns the contents of the external global matrix, vector, or scalar whose name is specified by *name*; it returns `J(0,0,.)` if the external global is not found.

Also see [Linking to external globals](#) in [M-2] **declarations**.

## Remarks

Also see [M-5] [findexternal\(\)](#). Rather than returning a pointer to the external global, as does `findexternal()`, `valofexternal()` returns the contents of the external global. This is useful when the external global contains a scalar:

```
tol = valofexternal("tolerance")
if (tol==J(0,0,.)) tol = 1e-6
```

Using `findexternal()`, one alternative would be

```
if ((p = findexternal("tolerance"))==NULL) tol = 1e-6
else tol = *p
```

For efficiency reasons, use of `valofexternal()` should be avoided with nonscalar objects; see [M-5] [findexternal\(\)](#).

## Conformability

```
valofexternal(name):
  name:      1 × 1
  result:    r × c    or    0 × 0 if not found
```

## Diagnostics

`valofexternal()` aborts with error if *name* contains an invalid name.

`valofexternal(name)` returns `J(0,0,.)` if *name* does not exist.

## Also see

[M-5] [findexternal\(\)](#) — Find, create, and remove external globals

[M-4] [programming](#) — Programming functions

Title

[M-5] **Vandermonde()** — Vandermonde matrices

Syntax

*numeric matrix*   **Vandermonde**(*numeric colvector x*)

Description

**Vandermonde**(*x*) returns the Vandermonde matrix containing the geometric progression of *x* in each row

$$\begin{bmatrix} 1 & x_1 & x_1^2 & x_1^3 & \dots & x_1^{n-1} \\ 1 & x_2 & x_2^2 & x_2^3 & \dots & x_2^{n-1} \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & x_n & x_n^2 & x_n^3 & \dots & x_n^{n-1} \end{bmatrix}$$

where  $n = \text{rows}(x)$ . Some authors use the transpose of the above matrix.

Remarks

Vandermonde matrices are useful in polynomial interpolation.

Conformability

**Vandermonde**(*x*):  
          *x*:         $n \times 1$   
      *result*:     $n \times n$

Diagnostics

None.

Alexandre-Théophile Vandermonde (1735–1796) was born in Paris. His first passion was music (particularly the violin) and he turned to mathematics only at the age of 35. Four papers dated 1771 and 1772 are his entire mathematical output, although all contain good work. He also worked in experimental science and the theory of music, arguing that musicians should ignore all theory and trust their trained ears, and was busy with various committees and other administration. Vandermonde was a strong supporter of the French Revolution. He is now best known for the Vandermonde determinant, even though it does not appear in any of his papers, and for the associated matrix. Lebesgue later conjectured that the attribution arises from a misreading of Vandermonde’s notation.

## Reference

Jones, P. S. 1976. Vandermonde, Alexandre-Théophile. In Vol. 13 of *Dictionary of Scientific Biography*, ed. C. C. Gillispie, 571–572. New York: Scribner’s.

## Also see

[\[M-4\]](#) **standard** — Functions to create standard matrices

Syntax

```
transmorphic colvector  vec(transmorphic matrix T)  
transmorphic colvector  vech(transmorphic matrix T)  
transmorphic matrix     invvech(transmorphic colvector v)
```

Description

`vec(T)` returns  $T$  transformed into a column vector with one column stacked onto the next.

`vech(T)` returns square and typically symmetric matrix  $T$  transformed into a column vector; only the lower half of the matrix is recorded.

`invvech(v)` returns `vech()`-style column vector  $v$  transformed into a symmetric (Hermitian) matrix.

Remarks

Remarks are presented under the following headings:

```
Example of vec()  
Example of vech() and invvech()
```

Example of `vec()`

```
: x  
      1  2  3  
1  [ 1  2  3  
2  [ 4  5  6  
  
: vec(x)  
      1  
1  [ 1  
2  [ 4  
3  [ 2  
4  [ 5  
5  [ 3  
6  [ 6
```

### Example of vech() and invvech()

```
: x
[symmetric]
      1  2  3
1      1
2      2  4
3      3  6  9

: v = vech(x)
: v
      1
1      1
2      2
3      3
4      4
5      6
6      9

: invvech(v)
[symmetric]
      1  2  3
1      1
2      2  4
3      3  6  9
```

### Conformability

```
vec(T):
      T:      r × c
result:      r * c × 1

vech(T):
      T:      n × n
result:      (n(n + 1)/2 × 1)

invvech(v):
      v:      (n(n + 1)/2 × 1)
result:      n × n
```

### Diagnostics

vec(T) cannot fail.

vech(T) aborts with error if T is not square. vech() records only the lower triangle of T; it does not require T be symmetric.

invvech(v) aborts with error if v does not have 0, 1, 3, 6, 10, ... rows.

---

## Also see

[M-4] [manipulation](#) — Matrix manipulation





**[M-6] Mata glossary of common terms**



# Title

[M-6] Glossary

## Description

Commonly used terms are defined here.

## Mata glossary

### arguments

The values a function receives are called the function’s arguments. For instance, in `lud(A, L, U)`,  $A$ ,  $L$ , and  $U$  are the arguments.

### array

An array is any indexed object that holds other objects as elements. Vectors are examples of 1-dimensional arrays. Vector  $\mathbf{v}$  is an array, and  $\mathbf{v}[1]$  is its first element. Matrices are 2-dimensional arrays. Matrix  $\mathbf{X}$  is an array, and  $\mathbf{X}[1, 1]$  is its first element. In theory, one can have 3-dimensional, 4-dimensional, and higher arrays, although Mata does not directly provide them. See [M-2] [subscripts](#) for more information on arrays in Mata.

Arrays are usually indexed by sequential integers, but in associative arrays, the indices are strings that have no natural ordering. Associative arrays can be 1-dimensional, 2-dimensional, or higher. If  $A$  were an associative array, then  $A[\text{“first”}]$  might be one of its elements. See [M-5] [asarray\(\)](#) for associative arrays in Mata.

### binary operator

A binary operator is an operator applied to two arguments. In 2–3, the minus sign is a binary operator, as opposed to the minus sign in `-9`, which is a [unary operator](#).

### broad type

Two matrices are said to be of the same broad type if the elements in each are numeric, are string, or are pointers. Mata provides two numeric types, real and complex. The term *broad type* is used to mask the distinction within numeric and is often used when discussing operators or functions. One might say, “The comma operator can be used to join the rows of two matrices of the same broad type,” and the implication of that is that one could join a real to a complex. The result would be complex. Also see [type](#), [eltype](#), and [orgtype](#).

c-conformability

Matrix, vector, or scalar  $A$  is said to be c-conformable with matrix, vector, or scalar  $B$  if they have the same number of rows and columns (they are *p-conformable*), or if they have the same number of rows and one is a vector, or if they have the same number of columns and one is a vector, or if one or the other is a scalar. c stands for colon; c-conformable matrices are suitable for being used with Mata's `:op` operators.  $A$  and  $B$  are c-conformable if and only if

$A$	$B$
$r \times c$	$r \times c$
$r \times 1$	$r \times c$
$1 \times c$	$r \times c$
$1 \times 1$	$r \times c$
$r \times c$	$r \times 1$
$r \times c$	$1 \times c$
$r \times c$	$1 \times 1$

The idea behind c-conformability is generalized elementwise operation. Consider  $C=A:*B$ . If  $A$  and  $B$  have the same number of rows and have the same number of columns, then  $||C_{ij}|| = ||A_{ij}*B_{ij}||$ . Now say that  $A$  is a column vector and  $B$  is a matrix. Then  $||C_{ij}|| = ||A_i*B_{ij}||$ : each element of  $A$  is applied to the entire row of  $B$ . If  $A$  is a row vector, each column of  $A$  is applied to the entire column of  $B$ . If  $A$  is a scalar,  $A$  is applied to every element of  $B$ . And then all the rules repeat, with the roles of  $A$  and  $B$  interchanged. See [M-2] [op\\_colon](#) for a complete definition.

class programming

See [object-oriented programming](#).

colon operators

Colon operators are operators preceded by a colon, and the colon indicates that the operator is to be performed elementwise.  $A:*B$  indicates element-by-element multiplication, whereas  $A*B$  indicates matrix multiplication. Colons may be placed in front of any operator. Usually one thinks of elementwise as meaning  $c_{ij} = a_{ij} <op> b_{ij}$ , but in Mata, elementwise is also generalized to include c-conformability. See [M-2] [op\\_colon](#).

column-major order

Matrices are stored as vectors. Column-major order specifies that the vector form of a matrix is created by stacking the columns. For instance,

: A

	1	2
1	1	4
2	2	5
3	3	6

is stored as

	1	2	3	4	5	6
1	1	2	3	4	5	6

in column-major order. The LAPACK functions use column-major order. Mata uses row-major order. See [row-major order](#).

## colvector

See [vector](#), [colvector](#), and [rowvector](#).

## complex

A matrix is said to be complex if its elements are complex numbers. Complex is one of two numeric types in Stata, the other being real. Complex is generally used to describe how a matrix is stored and not the kind of numbers that happen to be in it: complex matrix  $Z$  might happen to contain real numbers. Also see [type](#), [eltype](#), and [orgtype](#).

## condition number

The condition number associated with a numerical problem is a measure of that quantity's amenability to digital computation. A problem with a low condition number is said to be well conditioned, whereas a problem with a high condition number is said to be ill conditioned.

Sometimes reciprocals of condition numbers are reported and yet authors will still refer to them sloppily as condition numbers. Reciprocal condition numbers are often scaled between 0 and 1, with values near `epsilon(1)` indicating problems.

## conformability

Conformability refers to row-and-column matching between two or more matrices. For instance, to multiply  $A*B$ ,  $A$  must have the same number of columns as  $B$  has rows. If that is not true, then the matrices are said to be nonconformable (for multiplication).

Three kinds of conformability are often mentioned in the Mata documentation: [p-conformability](#), [c-conformability](#), and [r-conformability](#).

## conjugate

If  $z = a + bi$ , the conjugate of  $z$  is  $\text{conj}(z) = a - bi$ . The conjugate is obtained by reversing the sign of the imaginary part. The conjugate of a real number is the number itself.

## conjugate transpose

See [transpose](#).

## data matrix

A dataset containing  $n$  observations on  $k$  variables is often stored in an  $n \times k$  matrix. An observation refers to a row of that matrix; a variable refers to a column. When the rows are observations and the columns are variables, the matrix is called a data matrix.

## declarations

Declarations state the *eltype* and *orgtype* of functions, arguments, and variables. In

```
real matrix myfunc(real vector A, complex scalar B)
{
    real scalar i
    ...
}
```

the `real matrix` is a function declaration, the `real vector` and `complex scalar` are argument declarations, and `real scalar i` is a variable declaration. The `real matrix` states the function returns a real matrix. The `real vector` and `complex scalar` state the kind of arguments `myfunc()` expects and requires. The `real scalar i` helps Mata to produce more efficient compiled code.

Declarations are optional, so the above could just as well have read

```
function myfunc(A, B)
{
    ...
}
```

When you omit the function declaration, you must substitute the word `function`.

When you omit the other declarations, `transmorphic matrix` is assumed, which is fancy jargon for a matrix that can hold anything. The advantages of explicit declarations are that they reduce the chances you make a mistake either in coding or in using the function, and they assist Mata in producing more efficient code. Working interactively, most people omit the declarations.

See [\[M-2\] declarations](#) for more information.

## defective matrix

An  $n \times n$  matrix is defective if it does not have  $n$  linearly independent eigenvectors.

## dereference

Dereferencing is an action performed on pointers. Pointers contain memory addresses, such as 0x2a1228. One assumes something of interest is stored at 0x2a1228, say, a real scalar equal to 2. When one accesses that 2 via the pointer by coding `*p`, one is said to be dereferencing the pointer. Unary `*` is the dereferencing operator.

## diagonal matrix

A matrix is diagonal if its off-diagonal elements are zero;  $A$  is diagonal if  $A[i, j] == 0$  for  $i \neq j$ . Usually, diagonal matrices are also *square*. Some definitions require that a diagonal matrix also be a square matrix.

## diagonal of a matrix

The diagonal of a matrix is the set of elements  $A[i,j]$ .

## dyadic operator

Synonym for [binary operator](#).

## eigenvalues and eigenvectors

A scalar,  $\lambda$ , is said to be an eigenvalue of square matrix  $\mathbf{A}$ :  $n \times n$  if there is a nonzero column vector  $\mathbf{x}$ :  $n \times 1$  (called an eigenvector) such that

$$\mathbf{A}\mathbf{x} = \lambda\mathbf{x} \quad (1)$$

Equation (1) can also be written

$$(\mathbf{A} - \lambda\mathbf{I})\mathbf{x} = 0$$

where  $\mathbf{I}$  is the  $n \times n$  identity matrix. A nontrivial solution to this system of  $n$  linear homogeneous equations exists if and only if

$$\det(\mathbf{A} - \lambda\mathbf{I}) = 0 \quad (2)$$

This  $n$ th-degree polynomial in  $\lambda$  is called the characteristic polynomial or characteristic equation of  $A$ , and the eigenvalues  $\lambda$  are its roots, also known as the characteristic roots.

The eigenvector defined by (1) is also known as the right eigenvector, because matrix  $\mathbf{A}$  is postmultiplied by eigenvector  $\mathbf{x}$ . See [M-5] [eigensystem\(\)](#) and [left eigenvectors](#).

## eltype

See [type](#), [eltype](#), and [orgtype](#).

## epsilon(1), etc.

`epsilon(1)` refers to the unit roundoff error associated with a computer, also informally called machine precision. It is the smallest amount by which a number may differ from 1. For IEEE double-precision variables, `epsilon(1)` is approximately  $2.22045\text{e-}16$ .

`epsilon(x)` is the smallest amount by which a real number can differ from  $x$ , or an approximation thereof; see [M-5] [epsilon\(\)](#).

## exp

*exp* is used in syntax diagrams to mean “any valid expression may appear here”; see [M-2] [exp](#).

## external variable

See [global variable](#).

## function

The words *program* and *function* are used interchangeably. The programs that you write in Mata are in fact functions. Functions receive arguments and optionally return results.

Examples of functions that are included with Mata are `sqrt()`, `ttail()`, and `substr()`. Such functions are often referred to as the built-in functions or the library functions. Built-in functions refer to functions implemented in the C code that implements Mata, and library functions refer to functions written in the Mata programming language, but many users use the words interchangeably because how functions are implemented is of little importance. If you have a choice between using a built-in function and a library function, however, the built-in function will usually execute more quickly and the library function will be easier to use. Mostly, however, features are implemented one way or the other and you have no choice.

Also see [underscore functions](#).

For a list of the functions that Mata provides, see [\[M-4\] intro](#).

## generalized eigenvalues

A scalar,  $\lambda$ , is said to be a generalized eigenvalue of a pair of  $n \times n$  square numeric matrices  $\mathbf{A}$ ,  $\mathbf{B}$  if there is a nonzero column vector  $\mathbf{x}$ :  $n \times 1$  (called a generalized eigenvector) such that

$$\mathbf{A}\mathbf{x} = \lambda\mathbf{B}\mathbf{x} \quad (1)$$

Equation (1) can also be written

$$(\mathbf{A} - \lambda\mathbf{B})\mathbf{x} = 0$$

A nontrivial solution to this system of  $n$  linear homogeneous equations exists if and only if

$$\det(\mathbf{A} - \lambda\mathbf{B}) = 0 \quad (2)$$

In practice, the generalized eigenvalue problem for the matrix pair  $(\mathbf{A}, \mathbf{B})$  is usually formulated as finding a pair of scalars  $(w, b)$  and a nonzero column vector  $\mathbf{x}$  such that

$$w\mathbf{A}\mathbf{x} = b\mathbf{B}\mathbf{x}$$

The scalar  $w/b$  is a generalized eigenvalue if  $b$  is not zero.

Infinity is a generalized eigenvalue if  $b$  is zero or numerically close to zero. This situation may arise if  $\mathbf{B}$  is singular.

The Mata functions that compute generalized eigenvalues return them in two complex vectors,  $\mathbf{w}$  and  $\mathbf{b}$  of length  $n$ . If  $\mathbf{b}[i] = 0$ , the  $i$ th generalized eigenvalue is infinite, otherwise the  $i$ th generalized eigenvalue is  $\mathbf{w}[i]/\mathbf{b}[i]$ .



## global variable

Global variables, also known as external variables and as global external variables, refer to variables that are common across programs and which programs may access without the variable being passed as an argument.

The variables you create interactively are global variables. Even so, programs cannot access those variables without engaging in another step, and global variables can be created without your creating them interactively.

To access (and create if necessary) global external variables, you declare the variable in the body of your program:

```
function myfunction(...)
{
    external real scalar globalvar
    ...
}
```

See [Linking to external globals](#) in [M-2] **declarations**.

There are other ways of creating and accessing global variables, but the declaration method is recommended. The alternatives are `crexternal()`, `findexternal()`, and `rmexternal()` documented in [M-5] **findexternal()** and `valofexternal()` documented in [M-5] **valofexternal()**.

## hashing, hash functions, and hash tables

Hashing refers to a technique for quickly finding information corresponding to an identifier. The identifier might be a name, a Social Security number, fingerprints, or anything else on which the information is said to be indexed. The hash function returns a many-to-one mapping of identifiers onto a dense subrange of the integers. Those integers, called hashes, are then used to index a hash table. The selected element of the hash table specifies a list containing identifiers and information. The list is then searched for the particular identifier desired. The advantage is that rather than searching a single large list, one need only search one of  $K$  smaller lists. For this to be fast, the hash function must be quick to compute and produce roughly equal frequencies of hashes over the range of identifiers likely to be observed.

## Hermitian matrix

Matrix  $A$  is Hermitian if it is equal to its conjugate transpose;  $A = A'$ ; see [transpose](#). This means that each off-diagonal element  $a_{ij}$  must equal the conjugate of  $a_{ji}$ , and that the diagonal elements must be real. The following matrix is Hermitian:

$$\begin{bmatrix} 2 & 4 + 5i \\ 4 - 5i & 6 \end{bmatrix}$$

The definition  $A = A'$  is the same as the definition for a symmetric matrix, although usually the word *symmetric* is reserved for real matrices and Hermitian, for complex matrices. In this manual, we use the word *symmetric* for both; see [symmetric matrices](#).

## Hessenberg decomposition

The Hessenberg decomposition of a matrix,  $\mathbf{A}$ , can be written as

$$\mathbf{Q}'\mathbf{A}\mathbf{Q} = \mathbf{H}$$

where  $\mathbf{H}$  is in upper Hessenberg form and  $\mathbf{Q}$  is orthogonal if  $\mathbf{A}$  is real or unitary if  $\mathbf{A}$  is complex. See [M-5] [hessenbergd\(\)](#).

## Hessenberg form

A matrix,  $\mathbf{A}$ , is in upper Hessenberg form if all entries below the first subdiagonal are zero:  $A_{ij} = 0$  for all  $i > j + 1$ .

A matrix,  $\mathbf{A}$ , is in lower Hessenberg form if all entries above the first superdiagonal are zero:  $A_{ij} = 0$  for all  $j > i + 1$ .

## instance and realization

Instance and realization are synonyms for variable, as in [Mata variable](#). For instance, consider a real scalar variable  $X$ . One can equally well say that  $X$  is an instance of a real scalar or a realization of a real scalar. Authors represent a variable this way when they wish to emphasize that  $X$  is not representative of all real scalars but is just one of many real scalars. Instance is often used with structures and classes when the writer wishes to emphasize the difference between the values contained in the variable and the definition of the structure or the class. It is confusing to say that  $V$  is a class  $C$ , even though it is commonly said, because the reader might confuse the definition of  $C$  with the specific values contained in  $V$ . Thus careful authors say that  $V$  is an instance of class  $C$ .

## istmt

An *istmt* is an interactive statement, a statement typed at Mata's colon prompt.

## J(r, c, value)

$J()$  is the function that returns an  $r \times c$  matrix with all elements set to *value*; see [M-5] [J\(\)](#). Also,  $J()$  is often used in the documentation to describe the various types of *void* matrices; see [void matrix](#). Thus the documentation might say that such-and-such returns  $J(0, 0, .)$  under certain conditions. That is another way of saying that such-and-such returns a  $0 \times 0$  real matrix.

When  $r$  or  $c$  is 0, there are no elements to be filled in with *value*, but even so, *value* is used to determine the type of the matrix. Thus  $J(0, 0, 1i)$  refers to a  $0 \times 0$  complex matrix,  $J(0, 0, "")$  refers to a  $0 \times 0$  string matrix, and  $J(0, 0, NULL)$  refers to a  $0 \times 0$  *pointer* matrix.

In the documentation,  $J()$  is used for more than describing  $0 \times 0$  matrices. Sometimes, the matrices being described are  $r \times 0$  or are  $0 \times c$ . Say that a function `example(X)` is supposed to return a column vector; perhaps it returns the last column of  $X$ . Now say that  $X$  is  $0 \times 0$ . Function `example()` still should return a column vector, and so it returns a  $0 \times 1$  matrix. This would be documented by noting that `example()` returns  $J(0, 1, .)$  when  $X$  is  $0 \times 0$ .

## LAPACK

LAPACK stands for Linear Algebra PACKage and forms the basis for many of Mata's linear algebra capabilities; see [M-1] [LAPACK](#).

## left eigenvectors

A vector  $\mathbf{x}$ :  $n \times 1$  is said to be a left eigenvector of square matrix  $\mathbf{A}$ :  $n \times n$  if there is a nonzero scalar,  $\lambda$ , such that

$$\mathbf{x}\mathbf{A} = \lambda\mathbf{x}$$

## lval

*lval* stands for left-hand-side value and is defined as the property of being able to appear on the left-hand side of an equal-assignment operator. Matrices are *lvals* in Mata, and thus

```
X = ...
```

is valid. Functions are not *lvals*; thus, you cannot code

```
substr(mystr,1,3) = "abc"
```

*lvals* would be easy to describe except that *pointers* can also be *lvals*. Few people ever use pointers. See [M-2] [op\\_assignment](#) for a complete definition.

## machine precision

See [epsilon\(1\)](#), *etc.*

## .mata file

By convention, we store the Mata source code for function *function()* in file *function.mata*; see [M-1] [source](#).

## matrix

The most general organization of data, containing  $r$  rows and  $c$  columns. Vectors, column vectors, row vectors, and scalars are special cases of matrices.

## .mlib library

The object code of functions can be collected and stored in a library. Most Mata functions, in fact, are located in the official libraries provided with Stata. You can create your own libraries. See [M-3] [mata mlib](#).

## .mo file

The object code of a function can be stored in a .mo file, where it can be later reused. See [M-1] [how](#) and [M-3] [mata mosave](#).

## monadic operator

Synonym for [unary operator](#).

## NaN

NaN stands for Not a Number and is a special computer floating-point code used for results that cannot be calculated. Mata (and Stata) do not use NaNs. When NaNs arise, they are converted into . (missing value).

## norm

A norm is a real-valued function  $f(x)$  satisfying

$$\begin{aligned} f(0) &= 0 \\ f(x) &> 0 && \text{for all } x \neq 0 \\ f(cx) &= |c|f(x) \\ f(x+y) &\leq f(x) + f(y) \end{aligned}$$

The word *norm* applied to a vector  $x$  usually refers to its Euclidean norm,  $p = 2$  norm, or length: the square root of the sum of its squared elements. There are other norms, the popular ones being  $p = 1$  (the sum of the absolute values of its elements) and  $p = \text{infinity}$  (the maximum element). Norms can also be generalized to deal with matrices. See [M-5] [norm\(\)](#).

## NULL

A special value for a *pointer* that means “points to nothing”. If you list the contents of a pointer variable that contains NULL, the address will show as 0x0. See [pointer](#).

## numeric

A matrix is said to be numeric if its elements are real or complex; see [type](#), [eltype](#), and [orgtype](#).

## object code

Object code refers to the binary code that Mata produces from the source code you type as input. See [M-1] [how](#).

## object-oriented programming

Object-oriented programming is a programming concept that treats programming elements as objects and concentrates on actions affecting those objects rather than merely on lists of instructions. Object-oriented programming uses classes to describe objects. Classes are much like structures with a primary difference being that classes can contain functions (known as methods) as well as variables. Unlike structures, however, classes may inherit variables and functions from other classes, which in theory makes object-oriented programs easier to extend and modify than non-object-oriented programs.

## observations and variables

A dataset containing  $n$  observations on  $k$  variables is often stored in an  $n \times k$  matrix. An observation refers to a row of that matrix; a variable refers to a column.

## operator

An operator is  $+$ ,  $-$ , and the like. Most operators are binary (or dyadic), such as  $+$  in  $A+B$  and  $*$  in  $C*D$ . Binary operators also include logical operators such as  $\&$  and  $|$  (“and” and “or”) in  $E\&F$  and  $G|H$ . Other operators are unary (or monadic), such as  $!$  (not) in  $!J$ , or both unary and binary, such as  $-$  in  $-K$  and in  $L-M$ . When we say “operator” without specifying which, we mean binary operator. Thus colon operators are in fact colon binary operators. See [M-2] [exp](#).

## optimization

Mata compiles the code that you write. After compilation, Mata performs an *optimization* step, the purpose of which is to make the compiled code execute more quickly. You can turn off the optimization step—see [M-3] [mata set](#)—but doing so is not recommended.

## orgtype

See [type](#), [eltype](#), and [orgtype](#).

## orthogonal matrix and unitary matrix

$A$  is orthogonal if  $A$  is *square* and  $A'A=I$ . The word orthogonal is usually reserved for real matrices; if the matrix is complex, it is said to be unitary (and then transpose means conjugate-transpose). We use the word orthogonal for both real and complex matrices.

If  $A$  is orthogonal, then  $\det(A) = \pm 1$ .

## p-conformability

Matrix, vector, or scalar  $A$  is said to be  $p$ -conformable with matrix, vector, or scalar  $B$  if  $\text{rows}(A) == \text{rows}(B)$  and  $\text{cols}(A) == \text{cols}(B)$ .  $p$  stands for plus;  $p$ -conformability is one of the properties necessary to be able to add matrices together.  $p$ -conformability, however, does not imply that the matrices are of the same type. Thus  $(1,2,3)$  is  $p$ -conformable with  $(4,5,6)$  and with  $(\text{"this"}, \text{"that"}, \text{"what"})$  but not with  $(4\backslash 5\backslash 6)$ .

## permutation matrix and permutation vector

A *permutation matrix* is an  $n \times n$  matrix that is a row (or column) permutation of the identity matrix. If  $P$  is a permutation matrix, then  $P*A$  permutes the rows of  $A$  and  $A*P$  permutes the columns of  $A$ . Permutation matrices also have the property that  $P^{-1} = P'$ .

A *permutation vector* is a  $1 \times n$  or  $n \times 1$  vector that contains a permutation of the integers  $1, 2, \dots, n$ . Permutation vectors can be used with subscripting to reorder the rows or columns of a matrix. Permutation vectors are a memory-conserving way of recording permutation matrices; see [M-1] [permutation](#).

## pointer

A matrix is said to be a pointer matrix if its elements are pointers.

A pointer is the address of a *variable*. Say that variable  $X$  contains a matrix. Another variable  $p$  might contain 137,799,016 and, if 137,799,016 were the address at which  $X$  were stored, then  $p$  would be said to point to  $X$ . Addresses are seldom written in base 10, and so rather than saying  $p$  contains 137,799,016, we would be more likely to say that  $p$  contains 0x836a568, which is the way we write numbers in base 16. Regardless of how we write addresses, however,  $p$  contains a number and that number corresponds to the address of another variable.

In our program, if we refer to  $p$ , we are referring to  $p$ 's contents, the number 0x836a568. The monadic operator  $*$  is defined as “refer to the address” or “dereference”:  $*p$  means  $X$ . We could code  $Y = *p$  or  $Y = X$ , and either way, we would obtain the same result. In our program, we could refer to  $X[i, j]$  or  $(*p)[i, j]$ , and either way, we would obtain the  $i, j$  element of  $X$ .

The monadic operator  $\&$  is how we put addresses into  $p$ . To load  $p$  with the address of  $X$ , we code  $p = \&X$ .

The special address 0 (zero, written in hexadecimal as 0x0), also known as NULL, is how we record that a pointer variable points to nothing. A pointer variable contains NULL or it contains a valid address of another variable.

See [M-2] **pointers** for a complete description of pointers and their use.

## pragma

“(Pragmatic information) A standardised form of comment which has meaning to a compiler. It may use a special syntax or a specific form within the normal comment syntax. A pragma usually conveys non-essential information, often intended to help the compiler to optimise the program.” See *The Free On-line Dictionary of Computing*, <http://www.foldoc.org/>, Editor Denis Howe. For Mata, see [M-2] **pragma**.

## rank

Terms in common use are rank, row rank, and column rank. The row rank of a matrix  $A$ :  $m \times n$  is the number of rows of  $A$  that are linearly independent. The column rank is defined similarly, as the number of columns that are linearly independent. The terms *row rank* and *column rank*, however, are used merely for emphasis; the ranks are equal and the result is simply called the rank of  $A$ .

For a square matrix  $A$  (where  $m==n$ ), the matrix is invertible if and only if  $\text{rank}(A)==n$ . One often hears that  $A$  is of full rank in this case and rank deficient in the other. See [M-5] **rank()**.

## r-conformability

A set of two or more matrices, vectors, or scalars  $A, B, \dots$ , are said to be r-conformable if each is c-conformable with a matrix of  $\max(\text{rows}(A), \text{rows}(B), \dots)$  rows and  $\max(\text{cols}(A), \text{cols}(B), \dots)$  columns.

r-conformability is a more relaxed form of c-conformability in that, if two matrices are c-conformable, they are r-conformable, but not vice versa. For instance,  $A$ :  $1 \times 3$  and  $B$ :  $3 \times 1$  are r-conformable but not c-conformable. Also, c-conformability is defined with respect to a pair of matrices only; r-conformability can be applied to a set of matrices.

r-conformability is often required of the arguments for functions that would otherwise naturally be expected to require scalars. See *R-conformability* in [M-5] `normal()` for an example.

## real

A matrix is said to be a real matrix if its elements are all reals and it is stored in a `real` matrix. Real is one of the two numeric types in Mata, the other being complex. Also see *type*, *eltype*, and *orgtype*.

## row-major order

Matrices are stored as vectors. Row-major order specifies that the vector form of a matrix is created by stacking the rows. For instance,

```
: A
      1  2  3
1  1  2  3
2  4  5  6
```

1	1	2	3
2	4	5	6

is stored as

	1	2	3	4	5	6
1	1	2	3	4	5	6

in row-major order. Mata uses row-major order. The LAPACK functions use column-major order. See *column-major order*.

## rowvector

See *vector*, *colvector*, and *rowvector*.

## scalar

A special case of a *matrix* with one row and one column. A scalar may be substituted anywhere a matrix, vector, column vector, or row vector is required, but not vice versa.

## Schur decomposition

The Schur decomposition of a matrix, **A**, can be written as

$$\mathbf{Q}'\mathbf{A}\mathbf{Q} = \mathbf{T}$$

where **T** is in Schur form and **Q**, the matrix of Schur vectors, is orthogonal if **A** is real or unitary if **A** is complex. See [M-5] `schurd()`.

## Schur form

There are two Schur forms: real Schur form and complex Schur form.

A real matrix is in Schur form if it is block upper triangular with  $1 \times 1$  and  $2 \times 2$  diagonal blocks. Each  $2 \times 2$  diagonal block has equal diagonal elements and opposite sign off-diagonal elements. The real eigenvalues are on the diagonal and complex eigenvalues can be obtained from the  $2 \times 2$  diagonal blocks.

A complex square matrix is in Schur form if it is upper triangular with the eigenvalues on the diagonal.

## source code

Source code refers to the human-readable code that you type into Mata to define a function. Source code is compiled into object code, which is binary. See [M-1] [how](#).

## square matrix

A matrix is square if it has the same number of rows and columns. A  $3 \times 3$  matrix is square; a  $3 \times 4$  matrix is not.

## string

A matrix is said to be a string matrix if its elements are strings (text); see [type](#), [eltype](#), and [orgtype](#). In Mata, a string may be text or binary and may be up to 2,147,483,647 characters (bytes) long.

## structure

A structure is an [eltype](#), indicating a set of variables tied together under one name. `struct mystruct` might be

```
struct mystruct {
    real scalar    n1, n2
    real matrix    X
}
```

If variable `a` was declared a `struct mystruct scalar`, then the scalar `a` would contain three pieces: two real scalars and one real matrix. The pieces would be referred to as `a.n1`, `a.n2`, and `a.X`. If variable `b` were also declared a `struct mystruct scalar`, it too would contain three pieces, `b.n1`, `b.n2`, and `b.X`. The advantage of structures is that they can be referred to as a whole. You can code `a.n1=b.n1` to copy one piece, or you can code `a=b` if you wanted to copy all three pieces. In all ways, `a` and `b` are variables. You may pass `a` to a subroutine, for instance, which amounts to passing all three values.

Structures variables are usually scalar, but they are not limited to being so. If `A` were a `struct mystruct matrix`, then each element of `A` would contain three pieces, and one could refer, for instance, to `A[2,3].n1`, `A[2,3].n2`, and `A[2,3].X`, and even to `A[2,3].X[3,2]`.

See [M-2] [struct](#).

## subscripts

Subscripts are how you refer to an element or even a submatrix of a matrix.

Mata provides two kinds of subscripts, known as list subscripts and range subscripts.



In list subscripts,  $A[2,3]$  refers to the (2,3) element of  $A$ .  $A[(2\backslash 3), (4,6)]$  refers to the submatrix made up of the second and third rows, fourth and sixth columns, of  $A$ .

In range subscripts,  $A[|2,3|]$  also refers to the (2,3) element of  $A$ .  $A[|2,3\backslash 4,6|]$  refers to the submatrix beginning at the (2,3) element and ending at the (4,6) element.

See [M-2] [subscripts](#) for more information.

## symmetric matrices

Matrix  $A$  is symmetric if  $A = A'$ . The word *symmetric* is usually reserved for real matrices, and in that case, a symmetric matrix is a square matrix with  $a_{ij} = a_{ji}$ .

Matrix  $A$  is said to be Hermitian if  $A = A'$ , where the transpose operator is understood to mean the conjugate-transpose operator; see [Hermitian matrix](#). In Mata, the  $'$  operator is the conjugate-transpose operator, and thus, in this manual, we will use the word *symmetric* both to refer to real, symmetric matrices and to refer to complex, Hermitian matrices.

Sometimes, you will see us follow the word *symmetric* with a parenthesized Hermitian, as in, “the resulting matrix is symmetric (Hermitian)”. That is done only for emphasis.

The inverse of a symmetric (Hermitian) matrix is symmetric (Hermitian).

## symmetriconly

Symmetriconly is a word we have coined to refer to a square matrix whose corresponding off-diagonal elements are equal to each other, whether the matrix is real or complex. Symmetriconly matrices have no mathematical significance, but sometimes, in data-processing and memory-management routines, it is useful to be able to distinguish such matrices.

## time-series–operated variable

Time-series–operated variables are a Stata concept. The term refers to *op.varname* combinations such as `L.gnp` to mean the lagged value of variable `gnp`. Mata’s [M-5] [st\\_data\(\)](#) function works with time-series–operated variables just as it works with other variables, but many other Stata-interface functions do not allow *op.varname* combinations. In those cases, you must use [M-5] [st\\_tsrevar\(\)](#).

## traceback log

When a function fails—either because of a programming error or because it was used incorrectly—it produces a traceback log:

```
: myfunction(2,3)
      solve(): 3200 conformability error
      mysub(): - function returned error
myfunction(): - function returned error
      <istmt>: - function returned error
r(3200);
```

The log says that `solve()` detected the problem—arguments are not conformable—and that `solve()` was called by `mysub()` was called by `myfunction()` was called by what you typed at the keyboard. See [M-2] [errors](#) for more information.

## transmorphic

Transmorphic is an *eltype*. A scalar, vector, or matrix can be transmorphic, which indicates that its elements may be real, complex, string, pointer, or even a structure. The elements are all the same type; you are just not saying which they are. Variables that are not declared are assumed to be transmorphic, or a variable can be explicitly declared to be `transmorphic`. Transmorphic is just fancy jargon for saying that the elements of the scalar, vector, or matrix can be anything and that, from one instant to the next, the scalar, vector, or matrix might change from holding elements of one type to elements of another.

See [M-2] [declarations](#).

## transpose

The transpose operator is written different ways in different books, including  $'$ , superscript  $*$ , superscript  $T$ , and superscript  $H$ . Here we use the  $'$  notation:  $A'$  means the transpose of  $A$ ,  $A$  with its rows and columns interchanged.

In complex analysis, the transpose operator, however it is written, is usually defined to mean the conjugate transpose; that is, one interchanges the rows and columns of the matrix and then one takes the conjugate of each element, or one does it in the opposite order—it makes no difference. Conjugation simply means reversing the sign of the imaginary part of a complex number: the conjugate of  $1+2i$  is  $1-2i$ . The conjugate of a real is the number itself; the conjugate of 2 is 2.

In Mata,  $'$  is defined to mean conjugate transpose. Since the conjugate of a real is the number itself,  $A'$  is regular transposition when  $A$  is real. Similarly, we have defined  $'$  so that it performs regular transposition for string and pointer matrices. For complex matrices, however,  $'$  also performs conjugation.

If you have a complex matrix and simply want to transpose it without taking the conjugate of its elements, see [M-5] [transposeonly\(\)](#). Or code `conj(A')`. The extra `conj()` will undo the undesired conjugation performed by the transpose operator.

Usually, however, you want transposition and conjugation to go hand in hand. Most mathematical formulas, generalized to complex values, work that way.

## triangular matrix

A triangular matrix is a matrix with all elements equal to zero above the diagonal or all elements equal to zero below the diagonal.

A matrix  $A$  is *lower triangular* if all elements are zero above the diagonal, that is, if  $A[i, j] == 0, j > i$ .

A matrix  $A$  is *upper triangular* if all elements are zero below the diagonal, that is, if  $A[i, j] == 0, j < i$ .

A *diagonal matrix* is both lower and upper triangular. That is worth mentioning because any function suitable for use with triangular matrices is suitable for use with diagonal matrices.

A triangular matrix is usually *square*.

The inverse of a triangular matrix is a triangular matrix. The determinant of a triangular matrix is the product of the diagonal elements. The eigenvalues of a triangular matrix are the diagonal elements.

## type, eltype, and orgtype

The *type* of a matrix (or vector or scalar) is formally defined as the matrix's *eltype* and *orgtype*, listed one after the other—such as `real vector`—but it can also mean just one or the other—such as the *eltype* `real` or the *orgtype* `vector`.

*eltype* refers to the type of the elements. The *eltypes* are

<code>real</code>	numbers such as 1, 2, 3.4
<code>complex</code>	numbers such as 1+2i, 3+0i
<code>string</code>	strings such as "bill"
<code>pointer</code>	pointers such as <code>&amp;varname</code>
<code>struct</code>	structures
<code>numeric</code>	meaning real or complex
<code>transmorphic</code>	meaning any of the above

*orgtype* refers to the organizational type. *orgtype* specifies how the elements are organized. The *orgtypes* are

<code>matrix</code>	two-dimensional arrays
<code>vector</code>	one-dimensional arrays
<code>colvector</code>	one-dimensional column arrays
<code>rowvector</code>	one-dimensional row arrays
<code>scalar</code>	single items

The fully specified type is the element and organization types combined, as in `real vector`.

## unary operator

A unary operator is an operator applied to one argument. In `-2`, the minus sign is a unary operator. In `!(a==b | a==c)`, `!` is a unary operator.

## underscore functions

*Functions* whose names start with an underscore are called underscore functions, and when an underscore function exists, usually a function without the underscore prefix also exists. In those cases, the function is usually implemented in terms of the underscore function, and the underscore function is harder to use but is faster or provides greater control. Usually, the difference is in the handling of errors.

For instance, function `fopen()` opens a file. If the file does not exist, execution of your program is aborted. Function `_fopen()` does the same thing, but if the file cannot be opened, it returns a special value indicating failure, and it is the responsibility of your program to check the indicator and to take the appropriate action. This can be useful when the file might not exist, and if it does not, you wish to take a different action. Usually, however, if the file does not exist, you will wish to abort, and use of `fopen()` will allow you to write less code.

## unitary matrix

See [orthogonal matrix](#).

## variable

In a program, the entities that store values ( $a, b, c, \dots, x, y, z$ ) are called variables. Variables are given names of 1 to 32 characters long. To be terribly formal about it: a variable is a container; it contains a matrix, vector, or scalar and is referred to by its variable name or by another variable containing a *pointer* to it.

Also, *variable* is sometimes used to refer to columns of data matrices; see [data matrix](#).

## vector, colvector, and rowvector

A special case of a matrix with either one row or one column. A vector may be substituted anywhere a matrix is required. A matrix, however, may not be substituted for a vector.

A *colvector* is a vector with one column.

A *rowvector* is a vector with one row.

A *vector* is either a *rowvector* or *colvector*, without saying which.

## view

A view is a special type of matrix that appears to be an ordinary matrix, but in fact the values in the matrix are the values of certain or all variables and observations in the Stata dataset that is currently in memory. Its values are not just equal to the dataset's values; they are the dataset's values: if an element of the matrix is changed, the corresponding variable and observation in the Stata dataset also changes. Views are obtained by `st_view()` and are efficient; see [\[M-5\] st\\_view\(\)](#).

## void function

A function is said to be void if it returns nothing. For instance, the function [\[M-5\] printf\(\)](#) is a void function; it prints results, but it does not return anything in the sense that, say, [\[M-5\] sqrt\(\)](#) does. It would not make any sense to code `x = printf("hi there")`, but coding `x = sqrt(2)` is perfectly logical.

## void matrix

A matrix is said to be void if it is  $0 \times 0$ ,  $r \times 0$ , or  $0 \times c$ ; see [\[M-2\] void](#).

## Also see

[\[M-0\] intro](#) — Introduction to the Mata manual

[\[M-1\] intro](#) — Introduction and advice

# Subject and author index

This is the subject and author index for the *Mata Reference Manual*. Readers interested in topics other than Mata should see the [combined subject index](#) (and the [combined author index](#)) in the *Quick Reference and Index*.

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