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an73

Electronic version of the Stata Technical Bulletin is now available

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dm78

Describing variables in memory

Nicholas J. Cox, University of Durham, UK, n.j.cox@durham.ac.uk

Abstract: The `ds2` command is a variant on official Stata's `ds` that allows describing classes of variables, such as all numeric or string variables, all variables of a particular type, or variables that do or do not have value labels attached. In addition, `ds2` leaves `r(varlist)` in its wake as a macro containing a list of variable names, which may then be used in further commands.

Keywords: describing variables, variable names, data types, value labels, `ds`, data management.

Syntax

```
ds2 [varlist] [, { numeric string | type(vartype) }
      { hasval({ a | y | u | n }) | withval(value_label_name) } detail ]
```

Description

`ds2` lists variable names in a compact format (or, optionally, in the same format as `describe`). It is a variant on `ds`.

Options

`numeric` specifies that only numeric variables should be listed.

`string` specifies that only string variables should be listed.

`type(vartype)` specifies that only variables of type *vartype* should be listed. `type()` may be abbreviated down to as few as one character for byte, int, long, float and double. `string` or any abbreviation of it means string variables of any length, so that `type(str)` is equivalent to `string`. `type()` may not be combined with `numeric` or `string`.

`hasval({ a | y | u | n })` specifies that variables with or without value labels attached should be listed. `hasval(a)` or `hasval(y)` means attached and `hasval(u)` or `hasval(n)` means unattached. The variables listed will be a subset of those otherwise specified. `t(int) h(a)` means int variables with value labels attached.

`withval(value_label_name)` specifies that variables with value label *value_label_name* attached should be listed. The variables listed will be a subset of those otherwise specified.

`detail` specifies that a more detailed listing should be given. `ds2, sd` is equivalent to `describe` with all the string variables. `ds2, nd` is equivalent to `describe` with all the numeric variables.

Notice that `hasval()` and `withval()` may not be combined.

Remarks

The Stata commands `describe` and `ds` (see [R] **describe**) describe the variables in memory (or optionally, in the case of `describe`, in a Stata-format datafile on disk). Both are extremely useful, but limited in certain respects. This insert describes an alternative command, called `ds2`, for describing data in memory which offers some extra features. These features are likely to be especially useful to those managing datasets containing large numbers of variables.

Neither `describe` nor `ds` offers scope for describing classes of variables such as all numeric variables, or all byte variables, unless they are defined by the possession of similar names using some convention. With options `numeric` and `string`, `ds2` will list only numeric or string variables, respectively. Similarly, the option `type()` specifies a particular data type.

Neither `describe` nor `ds` offers scope for focusing on variables that do or do not have value labels attached, whether any set of value labels or a particular set. `hasval()` and `withval()` options meet this need.

`ds2` by default operates like `ds` by giving only a compact listing of variable names. If different options are specified, then only variables matching all criteria are listed.

Unlike `ds`, however, `ds2` has two further features. `ds2` with the `detail` option is a backdoor to `describe`, so that `ds2, detail string` is in effect `describe all string variables`. In addition, `ds2` leaves `r(varlist)` in its wake as a macro containing a list of variable names, which may then be used in further commands. Note that, like all such `r` class results, this macro is ephemeral; see [R] **Saved results**. It should thus be copied to a safe place if it may be needed later, by (for example)

```
. local varlist "`r(varlist)'"
```

Saved results

`ds2` saves in `r(varlist)` the names of the variables listed.

Examples

```
. use auto
. ds2, s
make
. ds2, s d
1. make      str18   %-18s                Make and Model
. ds2, n
price  mpg      rep78   hdroom   trunk    weight   length   turn
displ  gratio   foreign
. order `r(varlist)´
. describe
Contains data from auto.dta
obs:      74                1978 Automobile Data
vars:     12                7 Jan 1999 17:49
size:     3,478 (99.3% of memory free)
```

```
-----
1. price    int    %8.0gc          Price
2. mpg      int    %8.0g           Mileage (mpg)
3. rep78    int    %8.0g           Repair Record 1978
4. hdroom   float  %6.1f           Headroom (in.)
5. trunk    int    %8.0g           Trunk space (cu. ft.)
6. weight   int    %8.0gc          Weight (lbs.)
7. length   int    %8.0g           Length (in.)
8. turn     int    %8.0g           Turn Circle (ft.)
9. displ    int    %8.0g           Displacement (cu. in.)
10. gratio  float  %6.2f           Gear Ratio
11. foreign byte   %8.0g           origin      Car type
12. make    str18  %-18s           Make and Model
-----
```

Sorted by: foreign

```
. ds2, n
price  mpg      rep78   hdroom   trunk    weight   length   turn
displ  gratio   foreign
. su `r(varlist)´
Variable | Obs    Mean    Std. Dev.    Min    Max
-----+-----
price   | 74    6165.257  2949.496    3291   15906
mpg     | 74    21.2973  5.785503    12     41
rep78   | 69    3.405797  .9899323    1      5
hdroom  | 74    2.993243  .8459948    1.5    5
trunk   | 74    13.75676  4.277404    5      23
weight  | 74    3019.459  777.1936    1760   4840
length  | 74    187.9324  22.26634    142    233
turn    | 74    39.64865  4.399354    31     51
displ   | 74    197.2973  91.83722    79     425
gratio  | 74    3.014865  .4562871    2.19   3.89
foreign | 74    .2972973  .4601885    0      1
```

```
. ds2, t(float)
hdroom   gratio
. ds2, w(origin)
foreign
```

Acknowledgments

Jay Kaufman's suggestions led to `type()` and `withval()` options.

dm79	Yet more new matrix commands
------	------------------------------

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Abstract: Nine commands for matrix operations are introduced. Seven produce a new matrix or vector, namely a correlation matrix; a matrix that is an elementwise monadic function of another; a matrix with selected rows and/or columns of another; the `vec` or `vech` of a matrix; a vector with elements sorted by magnitude; and a matrix containing the elements of a vector. Two commands save matrices to variables: `svmat2` is an enhancement of official Stata `svmat`, while `svmatstv` saves a matrix as a single variable.

Keywords: matrices, vectors, correlation matrix, `vec`, `vech`, sorting, saving matrices, `svmat`.

Syntax

```
matcorr varlist [if exp] [in range] [weight] , matrix(matname) [ covariance ]
matewmf matrix1 matrix2 , function(f)
matselrc matrix1 matrix2 [ , row(rows) col(cols) names ]
matvec matrix vector
matvech matrix vector
matvsort vector1 vector2 [ , decrease ]
matvtom vector matrix , row(#) col(#) oorder({ row | col})
svmat2 [type] matrix [ , names({col | eqcol | matcol | stub* | namelist}) rnames(newvar) full ]
svmatstv [type] matrix , generate(newvar) [ lh pd uh ]
```

where *type* denotes a storage type for new variables, and the first-named matrix or vector supplied as argument before the comma must already exist. Any second-named matrix or vector may be new or already existing (in which case it will be overwritten).

Description

`matcorr` puts the correlations (optionally the variances and covariances) of *varlist* into matrix *matname*. This is a convenience command, given that the correlation matrix can already be obtained by

```
. matrix accum R = varlist, nocons dev
. matrix R = corr(R)
```

As with `correlate`, `matcorr` performs casewise deletion so that all correlations are calculated from those observations for which nonmissing values exist for all variables in *varlist*.

Given a matrix, say *A*, and a user-supplied monadic function *f* with Stata syntax *f*(), `matewmf` calculates a second matrix, say *B*, with typical element

$$B[i,j] = f(A[i,j])$$

provided that no *B*[*i*,*j*] would be missing. Many Stata functions have this syntax, such as `exp()` and `log()`.

Given a matrix, `matselrc` produces a matrix containing only specified rows and/or columns. The selection may include some or all of the rows and columns. The resulting matrix may be smaller or larger than the original or the same size. It may overwrite the original matrix. `matselrc` takes the order of row and column specification literally, and thus could be used to reorder the rows and/or columns of a matrix.

Given an expression defining a matrix, `matvec` calculates the `vec` of that matrix. The `vec` of an $r \times c$ matrix A is an $rc \times 1$ vector whose first r elements are the first column of A , the next r are the second column, and so on. The result is placed in the named vector.

Given an expression defining a square matrix, `matvech` calculates the `vech` of that matrix. The `vech` of an $r \times r$ matrix A is an $r(r+1)/2 \times 1$ vector defined similarly to `vec` except that only the elements of A on or below the main diagonal are placed in the vector. The result is placed in the named vector. For further details on `vec` and `vech`, see Harville (1997).

Given a row or column vector, `matvsort` sorts the elements into numeric order and places the sorted elements into another (or the same) vector. The default is increasing order; the option `decrease` specifies decreasing order. `matvsort` may make it easier to identify the smallest or largest element(s) of a vector. If a vector `b` has been sorted into increasing order, its smallest element is accessible as `b[1,1]` and its largest as `b[1,colsof(b)]` if `b` is a row vector and `b[rowsof(b),1]` if `b` is a column vector. The corresponding names are accessible as in this example:

```
. local colnames : colnames b
. local ciname : word 1 of `colnames'
```

`matvsort` typically changes the sort order of the data. You may need to resort the data.

`matvtom` converts a vector ($r \times 1$ or $1 \times c$) to a `row` \times `col` matrix so long as the numbers of elements in the vector and the matrix are equal.

`svmat2` takes a matrix and stores its columns as new variables. It is the reverse of `mkmat`. `svmat2` adds to `svmat` (in official Stata) options to save matrix row names in a new string variable and to use any desired new variable names. Matrix column names including colons `:` (always) or periods `.` (usually) cannot be used as variable names.

`svmatstv` takes a matrix and stores its values as a single new variable. Values are in row major order, i.e. `A[1,2]` precedes `A[2,1]` whenever both are included. If column major order is desired, transpose the matrix first. `svmatstv` would be useful for some analyses of correlation matrices and tables of counts (contingency tables), as when it is desired to analyze the distribution of correlations or counts. `svmatstv` requires that the number of values saved should not exceed the current number of observations. If desired, increase that by using `set obs`.

See also Weesie (1997) and Cox (1999) for other sets of matrix programs complementing Stata's built-in commands.

Options of `matcorr`

`matrix(matrix)` specifies the name of a matrix to hold the results. It is a required option.

`covariance` specifies the calculation of covariances.

Options of `matewmf`

`function(f)` specifies a monadic function and is a required option. The function must have Stata syntax `f()`, take a single finite numeric argument and produce a single finite numeric result; see [R] [functions](#).

Options of `matselrc`

`row(rows)` specifies rows. The specification should be either a numlist containing integers between 1 and the number of rows, or a list of row names, in which case each individual name should be explicit. Quotation marks `" "` should not be used in the second case.

`col(cols)` specifies columns. The specification should be either a numlist containing integers between 1 and the number of columns, or a list of column names, in which case each individual name should be explicit. Quotation marks `" "` should not be used in the second case.

`names` is for a special case, when the row or column names would look like a numlist to Stata, but they are really names. Thus `matrix rowname A = 3 2 1` is legal in Stata for a matrix with 3 rows. The `names` option is used to force Stata to treat `row(3 2 1)` as a specification of row names.

Option of `matvsort`

`decrease` specifies sorting into decreasing order, smallest last.

Options of `matvtom`

`row(#)` specifies the number of rows in the new matrix.

`col(#)` specifies the number of columns in the new matrix.

`order()` specifies whether elements are to be assigned row by row or column by column.

`order(row)` (or `order(r)`) specifies that the first row elements of *vector* be the first row of *matrix*, and so forth.

`order(col)` (or `order(c)`) specifies that the first col elements of *vector* will be the first column of *matrix*, and so forth.

All options are required.

Options of `svmat2`

`names({col | eqcol | matcol | stub* | namelist})` specifies how the new variables are to be named.

`names(col)` specifies use of column names as the names of the variables.

`names(eqcol)` specifies use of equation names prefixed to the column names.

`names(matcol)` specifies use of the matrix name prefixed to the column names.

`names(stub*)` specifies naming the variables *stub1*, *stub2*, ... The * must be given. Note: this convention differs from that in `svmat`.

`names(namelist)` specifies names for variables according to *namelist*, one new name for each column in *varlist*.

If `names()` is not specified, the variables are named *matrix1*, *matrix2*, ... where *matrix* is the name of the matrix. If necessary, names will be truncated to 8 characters; if these names are not unique, an error will be returned.

`rnames(newvar)` names a new string variable for storing matrix row names.

`full` specifies that full row names are to be used.

Options of `svmatstv`

`generate(newvar)` specifies the name of the new variable and is a required option.

For a square matrix,

`lh` specifies that the lower half matrix is to be stored. This consists of elements below the principal diagonal.

`pd` specifies that the principal diagonal (also known as the major or main diagonal, or just diagonal) is to be stored. For a matrix *A*, this is *A*[1,1], *A*[2,2], etc.

`uh` specifies that the upper half matrix is to be stored. This consists of elements above the principal diagonal.

`lh`, `pd` and `uh` may be used in any combination.

Examples

Given some variables, to produce the correlation or covariance matrix

```
. set obs 10
obs was 0, now 10
. gen x1=invnorm(uniform())
. gen x2=invnorm(uniform())
. gen x3=invnorm(uniform())
. gen x4=invnorm(uniform())
. matcorr x1 x2 x3 x4, m(rho)
(obs=10)
symmetric rho[4,4]
      x1      x2      x3      x4
x1  1.0000
x2 -0.1715  1.0000
x3  0.1387  0.1537  1.0000
x4  0.2535  0.1276 -0.2487  1.0000
```

To exponentiate each element in matrix *rho*

```
. mategmf rho erho, f(exp)
. matrix list erho
symmetric erho[4,4]
      x1      x2      x3      x4
x1  2.7182818
x2  .8423907  2.7182818
x3  1.1488146  1.1660867  2.7182818
x4  1.2885526  1.1360762  .77981106  2.7182818
```

To keep only the specified columns and/or rows, whether overwriting the original matrix or putting elements in a new matrix

```
. matselrc erho erho, c(1 3)
. matrix list erho
erho[4,2]
      x1      x3
x1  2.7182818  1.1488146
x2   .8423907  1.1660867
x3  1.1488146  2.7182818
x4  1.2885526  .77981106
```

To reverse the columns of a 4×5 matrix

```
. matrix input A = (1,2,3,4,5\6,7,8,9,10\11,12,13,14,15\16,17,18,19,20)
. matrix list A
A[4,5]
      c1  c2  c3  c4  c5
r1   1   2   3   4   5
r2   6   7   8   9  10
r3  11  12  13  14  15
r4  16  17  18  19  20
. matselrc A B, c(5/1)
. matrix list B
B[4,5]
      c5  c4  c3  c2  c1
r1   5   4   3   2   1
r2  10   9   8   7   6
r3  15  14  13  12  11
r4  20  19  18  17  16
```

To illustrate `matvec` and `matvech`, we have

```
. matrix input A=(1,2,3\2,4,5\3,5,6)
. matrix list A
symmetric A[3,3]
      c1  c2  c3
r1   1
r2   2   4
r3   3   5   6
. matvec A AVEC
. matrix list AVEC
AVEC[9,1]
      vec
r1   1
r2   2
r3   3
r1   2
r2   4
r3   5
r1   3
r2   5
r3   6
. matvech A AVECH
. matrix list AVECH
AVECH[6,1]
      vech
r1   1
r2   2
r3   3
r2   4
r3   5
r3   6
```

To illustrate sorting a vector, we have

```
. matrix input C=(5,4,8,3,2)
. matvsort C D
. matrix list D
D[1,5]
      c5  c4  c2  c1  c3
r1   2   3   4   5   8
```

We can put the elements of a 1×10 vector into a 5×2 matrix either “by rows” as in

```
. matrix input E=(1,2,3,4,5,6,7,8,9,10)
. matvtom E EMR, r(5) c(2) o(r)
. matrix list EMR

EMR[5,2]
   c1  c2
r1   1   2
r2   3   4
r3   5   6
r4   7   8
r5   9  10
```

or “by columns” as in

```
. matvtom E EMC, r(5) c(2) o(c)
. matrix list EMC

EMC[5,2]
   c1  c2
r1   1   6
r2   2   7
r3   3   8
r4   4   9
r5   5  10
```

A vector may be copied to two variables, one to contain the elements and one to contain the row names, as in the following example using Stata’s automobile data.

```
. regress mpg weight gratio foreign
. matrix c=e(b)'
. svmat2 double c, name(bvector) r(bnames)
. matrix list c

c[4,1]
           y1
weight  -.00613903
gratio   1.4571134
foreign  -2.2216815
_cons   36.101353
```

Putting a matrix into a single variable might be useful for further analysis of say a correlation matrix or a table of counts. The correlation matrix is square and symmetric and the diagonal containing correlations that are all identically 1 does not reveal any properties of the data. Hence one of the options `lh` and `uh` is appropriate. After `svmat` it is then easy to look at the distribution of correlations or counts.

Acknowledgments

Kit Baum and Vincent Wiggins gave very helpful advice.

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dm80	Changing numeric variables to string
------	--------------------------------------

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Abstract: `tostring` changes numeric variables to string. The most compact string format possible is used. Any existing string variables specified are left unchanged.

Keywords: string variables, numeric variables, data types, data management.

Syntax

```
tostring varlist [, {fformat(format) | usedisplayformat} nodecode ]
```

Description

`tostring` changes numeric variables to string. The most compact string format possible is used. Any existing string variables specified are left unchanged.

Options

`format(format)` specifies the use of a format as an argument to the `string()` function; see Remarks below, [U] **16.3.5 String functions** or [U] **15.5 Formats**.

`usedisplayformat` specifies that the current display format should be used for each individual variable. This is the format set by the `format` command. For example, this option could be useful when using social security numbers.

`format()` and `usedisplayformat` cannot be used together.

`nodecode` specifies that numeric values of each variable should always be used, even when value labels exist. The default is to decode any value labels.

Remarks

The great divide among data types in Stata is between numeric data types (`byte`, `int`, `float`, `long`, and `double`) and string data types. Conversions between different numeric types and between different string data types are essentially a matter of using data types large enough to hold values accurately, yet small enough to economize on storage. Quite often these changes are carried out automatically by Stata or easily achieved using `recast` or `compress`. There are problems needing more drastic changes of data types. In particular, `recast` does not change numeric variables to string, or vice versa, so other commands are required.

This insert describes a utility, `tostring`, for changing numeric variables to string. Suppose you have a numeric variable, and you wish to change it to a string variable. In our experience, this need arises most commonly when the numeric variable is really an identifier of some kind—it is essentially a set of labels for individuals that happens to be numeric—and you wish to carry out string manipulation on this variable using Stata's string functions.

`tostring` is almost the inverse of `destring` (Cox and Gould 1997, Cox 1999a, 1999b), which changes variables from string to numeric.

To understand what the `tostring` command does and when it is the best command to use, we need to look at existing Stata commands for numeric to string conversion. In general, precise changing of numeric data to string equivalents can be a little difficult given that Stata, like most software, holds numeric data to finite precision and in binary form; see the discussion in [U] **16.10 Precision and problems therein**. Having said that, the intent behind `tostring` is to provide a convenient command that will work well.

Comparison with `decode` and `generate`

When the problem is to create a string copy of a numeric variable, you should use `decode` or `generate`.

If the numeric variable has value labels attached, the appropriate command is `decode`; see [R] **encode**.

If the numeric variable has no value labels attached, the appropriate command is `generate` with the `string()` function; see [U] **16.3.5 String functions**:

```
. generate strtype strvar = string(numvar)
```

There are various issues that you need to think about.

First, you need to decide the string type to substitute in place of *strtype*. String variables can be any length from `str1`, holding at most 1 character, to `str80`, holding at most 80 characters. You could determine the type required by counting how many characters you need to hold the longest number in *numvar*. Often this is easy: if you know that all your numbers are 4 digits or fewer, `str4` will be necessary. Or, you could use `summarize` to show the maximum and minimum values (the minimum value may be the longest if variables take on negative values). In many cases, you might make a mistake by miscounting, forgetting about minus signs, or whatever. It may be better to let Stata work out by itself which *strtype* is needed:

```
. gen str1 strvar = ""
. replace strvar = string(numvar)
```

The principle here is that Stata will automatically promote the string variable, which is born as a `str1` with all missing values, to whatever length is needed. The time spent in typing two lines rather than one is often much less than the time spent working out the length that is necessary.

Second, you need to decide on the appropriate format. In particular, very long integers often need to be changed to strings without any loss of detail whenever each individual digit is informative, particularly if the number is the identifier of a person. The default format of the `string()` function can truncate long integers by converting them to scientific format as powers of 10. For example, the 8 digit integer 12345678 will be converted to a string representation "1.23e+07". This problem arises because, as you might imagine, the default format of `string()` was chosen as a compromise given the variety of numbers that it may meet. That compromise is not a good choice for very long integers, and so you need to supply your own format as a second argument to `string()`. You will find, for example, that a format of `%11.0g` will work as desired for 9 digit integers:

```
. gen str1 strvar = ""
. replace strvar = string(numvar, "%11.0g")
```

Third, note that a numeric missing value, represented by the period (`.`), is converted by `string()` to `."`. If you prefer instead to have empty strings, you will need to use `replace`:

```
. replace strvar = "" if strvar == "."
```

This detail is unlikely to be crucial in most applications.

You should use either `decode` or `generate` whenever you wish to have both numeric and string versions of the same information. Note that for nonintegers or very large integers, repeated changes using `tostring` and `destring` can result in loss of precision through rounding errors and are not recommended.

If you wanted to have both versions for several such variables, you would need to repeat the appropriate command for each variable. It might be convenient to do this with `for`.

Using `tostring`

If you decide that you do wish to change the variable from numeric to string, then `tostring` will be more convenient, for various reasons.

First, `tostring` can change several variables at once, and even all variables if `_all` is specified explicitly as the *varlist*. Any existing string variables are ignored and left unchanged. Deliberately, and unlike `destring`, the permitted syntax does not include a bare `tostring`; you must specify a *varlist*. The rationale is that most users have datasets in which the great majority of variables are numeric, and it is thought unlikely that users would want to change many of these to string. In any case, the loss of precision from using an inappropriate format could lead to degradation of many datasets. Thus, users are protected by this detail of syntax from accidentally changing all numeric variables. Be warned that `tostring _all` should be issued if, and only if, you are sure that all numeric variables in your dataset should be changed to string.

Second, `tostring` is smart enough to work out the appropriate string type, given a variable and a format, and to `decode` numeric variables with value labels and to use `generate` with `string()` on numeric variables without value labels. The option `nodecode` is available should you wish not to have decoding. Recall, however, the key difference between `tostring` on the one hand and `decode` and `generate` on the other: `tostring` changes existing variables, while `decode` and `generate` produce copy variables of different type containing the same information, except for possible loss of detail through inappropriate format.

Third, the default format used by `tostring` is `%12.0g`. This format is, in particular, sufficient to convert integers held as `byte`, `int`, or `long` variables to string equivalent without loss of precision. Users will, in many cases, need to specify a format themselves, especially when the numeric data have fractional parts, and for some reason, a change to string is required. The format used, whether the default or a specified alternative, is applied to all variables in the *varlist*. Thus, changing different variables using different formats would require repeated `tostring` commands.

An exception to this arises if the `usedisplayformat` option is specified, which means that the display format associated with each variable will be used. The display format is set by `format` and shown by `describe` and can differ from variable to variable. It is at the user's discretion, but many users prefer to have a display format which rounds results quite vigorously, say to 1 or 2 decimal places. Changing to string with such a display format would frequently degrade the original data, or, in the case of integers, introduce superfluous zeros and decimal points.

With `tostring`, a numeric missing value, represented by the period (`.`), is converted to the missing or empty string `."`. If you prefer to have `."`, you will need to use `replace`:

```
. replace strvar = "." if strvar == ""
```

This detail is unlikely to be crucial in most applications.

Examples

Suppose that your data include U.S. social security numbers, identifiers for people which are typically 9 digit integers. These are stored as a numeric variable in your present dataset and as a string variable in another dataset which you want to merge. `tostring` will change the numeric variable to string, making this easier. Here is a token example set. Note first that the social security numbers have been assigned a `%09.0f` format so that leading zeros are explicit:

```
. describe
Contains data from ssn.dta
  obs:      5
  vars:      1                20 Apr 2000 14:50
  size:     40 (99.9% of memory free)
-----
   1. ssn      long   %09.0f
-----
Sorted by:
. list
           ssn
  1. 129933896
  2. 070045728
  3. 103618872
  4. 075156200
  5. 140035552
```

Using commands from official Stata, the following steps convert this numeric variable to a string variable. Note the use of a format argument to the `string()` function, needed to avoid loss of information under scientific format.

```
. gen str9 ssn2 = string(ssn, "%09.0f")
. list
           ssn      ssn2
  1. 129933896 129933896
  2. 070045728 070045728
  3. 103618872 103618872
  4. 075156200 075156200
  5. 140035552 140035552
. drop ssn
. rename ssn2 ssn
. describe
Contains data from ssn.dta
  obs:      5
  vars:      1                20 Apr 2000 14:50
  size:     65 (99.9% of memory free)
-----
   1. ssn      str9   %9s
-----
Sorted by:
      Note: dataset has changed since last saved
```

The change can, however, be executed by a single `tostring` command with the `usedisplayformat` option.

```
. describe
Contains data from ssn.dta
  obs:      5
  vars:      1                20 Apr 2000 14:50
  size:     40 (99.9% of memory free)
-----
   1. ssn      long   %09.0f
-----
Sorted by:
. list
           ssn
  1. 129933896
  2. 070045728
  3. 103618872
  4. 075156200
  5. 140035552
. tostring ssn, u
ssn was long now str9
```

```

. list
      ssn
1. 129933896
2. 070045728
3. 103618872
4. 075156200
5. 140035552
. describe
Contains data from ssn.dta
  obs:          5
  vars:          1                20 Apr 2000 14:50
  size:          65 (99.8% of memory free)
-----
 1. ssn          str9          %9s
-----
Sorted by:
  Note: dataset has changed since last saved

```

Acknowledgments

Ken Higbee and Thomas Steichen made very helpful comments.

References

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- . 1999b. dm45.2: Changing string variables to numeric: correction. *Stata Technical Bulletin* 52: 2. Reprinted in *Stata Technical Bulletin Reprints*, vol. 9, p. 14.
- Cox, N. J. and W. Gould. 1997. dm45: Changing string variables to numeric. *Stata Technical Bulletin* 37: 4–6. Reprinted in *Stata Technical Bulletin Reprints*, vol. 7, pp. 34–37.

gr44

Graphing point estimate and confidence intervals

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Abstract: A program called `gorciv` is described that plots point estimates and confidence intervals following estimation commands.

Keywords: graphing estimates, point estimates, confidence intervals, graphing regression models.

Introduction

After running an estimation command, it is often desirable to graph the point estimates and confidence intervals that are produced. This is especially useful when attempting to compare the point estimates for categorical variables; it allows one to focus more on the confidence intervals and possible trends in the data than on p -values.

The program `gorciv` uses the point estimates and standard errors generated from the most recent estimation command (such as `logistic`, `glm`, `regress`, `poisson`, `stcox`, and so on). `gorciv` plots the point estimate and specified confidence intervals vertically along the x -axis for each variable the user specifies.

Syntax

```
gorciv varlist [ , eform point level(#) yscale(##) title(str) ]
```

Options

`eform` specifies that the exponential point estimate and confidence intervals will be graphed. A line is automatically graphed at $y = 0$ if `eform` is not specified, and at $y = 1$ if `eform` is specified.

`point` specifies that the value of the point estimate be labeled on the graph.

`level(#)` specifies the confidence level, in percent, for confidence intervals. The default is `level(95)` or as set by `set level`.

`yscale(##)` can be used to widen the scale of the y -axis.

`title(str)` specifies the title for the graph.

Example 1

We consider Stata's cancer data.

```
. use cancer, clear
(Patient Survival in Drug Trial)
. describe
Contains data from /usr/local/stata/cancer.dta
obs:          48          Patient Survival in Drug Trial
vars:         4          16 Nov 1998 11:49
size:        576 (99.9% of memory free)
-----
1. studytim  int    %8.0g          Months to death or end of exp.
2. died      int    %8.0g          1 if patient died
3. drug      int    %8.0g          Drug type (1=placebo)
4. age       int    %8.0g          Patient's age at start of exp.
-----
Sorted by:
```

We wish to run a Cox proportional hazards model so we first need to set the data in survival time format.

```
. stset studytim died
failure event: died ~= 0 & died ~= .
obs. time interval: (0, studytim]
exit on or before: failure
-----
48 total obs.
0 exclusions
-----
48 obs. remaining, representing
31 failures in single record/single failure data
744 total analysis time at risk, at risk from t =          0
          earliest observed entry t =          0
          last observed exit t =          39
```

Now we wish to run our survival model with the variable drug entered as a categorical variable.

```
. xi: stcox i.drug age
i.drug          Idrug_1-3    (naturally coded; Idrug_1 omitted)
      failure _d: died
      analysis time _t: studytim
Iteration 0:    log likelihood = -99.911448
Iteration 1:    log likelihood = -82.331523
Iteration 2:    log likelihood = -81.676487
Iteration 3:    log likelihood = -81.652584
Iteration 4:    log likelihood = -81.652567
Refining estimates:
Iteration 0:    log likelihood = -81.652567
Cox regression -- Breslow method for ties
No. of subjects =          48          Number of obs =          48
No. of failures =          31
Time at risk    =          744
Log likelihood  = -81.652567          LR chi2(3) =          36.52
                                          Prob > chi2 =          0.0000
-----
      _t |
      _d | Haz. Ratio  Std. Err.      z    P>|z|    [95% Conf. Interval]
-----+-----
Idrug_2 | .1805839   .0892742   -3.462  0.001   .0685292   .4758636
Idrug_3 | .0520066   .034103    -4.508  0.000   .0143843   .1880305
age     | 1.118334   .0409074    3.058  0.002   1.040963   1.201455
-----
```

Now that we have run our estimation command, we can use `gorciv` to display the hazard ratios and 90% confidence intervals graphically; see Figure 1 at the end of this insert.

```
. xi: gorciv i.drug, eform ys(0,2) level(90) ti("Point Estimates and 90% Confid
> ence Intervals")
i.drug          Idrug_1-3    (naturally coded; Idrug_1 omitted)
```

Example 2

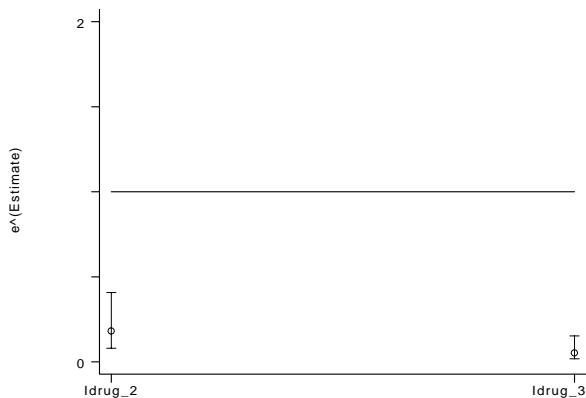
Suppose we wish to generate a graph of the coefficients and 95% C.I.'s following a linear regression command using the Stata's auto data; see Figure 2 at the end of this insert.

```
. use auto.dta
. regress mpg foreign length weight trunk turn displ hdroom gratio rep78 price
```

Source	SS	df	MS	Number of obs = 69		
Model	1674.51142	10	167.451142	F(10, 58)	=	14.59
Residual	665.691475	58	11.4774392	Prob > F	=	0.0000
				R-squared	=	0.7155
				Adj R-squared	=	0.6665
				Root MSE	=	3.3878

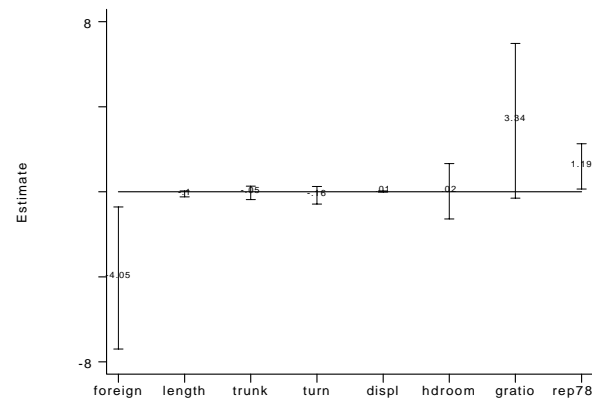
	mpg	Coef.	Std. Err.	t	P> t	[95% Conf. Interval]	
foreign		-4.051849	1.704339	-2.377	0.021	-7.463455	-.6402443
length		-.100628	.0693695	-1.451	0.152	-.2394862	.0382302
weight		-.0027956	.0025719	-1.087	0.282	-.0079438	.0023527
trunk		-.052152	.1618622	-0.322	0.748	-.3761543	.2718504
turn		-.1629084	.2099955	-0.776	0.441	-.58326	.2574432
displ		.0115954	.0143612	0.807	0.423	-.0171517	.0403425
hdroom		.022946	.6649112	0.035	0.973	-1.308018	1.35391
gratio		3.337897	1.857397	1.797	0.078	-.3800862	7.055881
rep78		1.192596	.5421194	2.200	0.032	.1074257	2.277765
price		-.0000627	.0002226	-0.282	0.779	-.0005083	.0003828
_cons		41.10397	10.25511	4.008	0.000	20.57613	61.63181

```
. gorciv foreign length trunk turn displ hdroom gratio rep78, ti("RR's and 95%
> Confidence Intervals") ys(-8,8) point
```



Point Estimates and 90% Confidence Intervals

Figure 1. Using gorciv with cancer data following stcox



RR's and 95% Confidence Intervals

Figure 2. Confidence intervals for regression coefficients using the auto data

sbe20.1

Update of galbr

Aurelio Tobias, Universidad Miguel Hernandez, Alicante, Spain, bledatobias@ctv.es

Abstract: Minor bugs in the galbr command have been fixed.

Keywords: meta-analysis.

The galbr command provides a graphical display to get a visual impression of the amount of heterogeneity from a meta-analysis. Minor bugs have been fixed on labeling axes and on display of linear regression lines.

References

Tobias, A. 1998. sbe20: Assessing heterogeneity in meta-analysis: the Galbraith plot. *Stata Technical Bulletin* 41: 15–17. Reprinted in *Stata Technical Bulletin Reprints*, vol. 7, pp. 133–136.

sbe26.1	Update of metainf
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Abstract: Minor bugs in the `metainf` command have been fixed.

Keywords: meta-analysis.

The `metainf` command introduced in Tobias (1999) investigates the influence of a single study on an overall meta-analysis estimate. A problem on sorting study labels has been solved. Now more than twenty studies can be horizontally displayed in the plot, since this update uses the `hplot` command introduced in Cox (1999), which is a more powerful graphical display. Thus, the `hplot` command must be installed, and all `hplot` options are also available using `metainf`.

References

Tobias, A. 1999. sbe26: Assessing the influence of a single study in meta-analysis. *Stata Technical Bulletin* 47: 15–17. Reprinted in *Stata Technical Bulletin Reprints*, vol. 8, pp. 108–110.

Cox, N. J. 1999. `hplot` and `hbar` for presentation graphics. Proceedings of the 5th Stata UK user's group meeting, London.

sbe28.1	Update of metap
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Abstract: Minor bugs in the `metap` command have been fixed.

Keywords: meta-analysis.

The `metap` command introduced in Tobias (1999) provides a combination of one-tailed p -values from each study. Minor bugs have been fixed on displaying results. Now different estimation methods have been obtained using the `method` option, which replaces the old `e` option. Now there are three alternatives available: `f` uses Fisher's method, `ea` uses Edgington's additive method, and `en` uses Edgington's normal approximation method. Again, by default Fisher's method is used. Finally, the following saved results are available

```

r(method)  the method used to combine the p-values
r(n)       the number of studies
r(stat)    the statistic used to combine the p-values
r(z)       the value of the statistic used
r(pvalue)  returns the combined p-value

```

References

Tobias, A. 1999. sbe28: Meta-analysis of p -values. *Stata Technical Bulletin* 49: 15–17. Reprinted in *Stata Technical Bulletin Reprints*, vol. 9, pp. 138–140.

sbe35	Menus for epidemiological statistics
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Abstract: Stata's `epitab` commands perform a number of relatively simple statistical calculations that are commonly used in epidemiology (computation of risk ratios, odds ratios, and rate ratios, with appropriate confidence intervals). Their immediate versions apply to tabular data, where subject-level information has been collapsed to the form of a two-way table. The command `calcmenu`, described in this insert, uses Stata's menu-and-dialog programming features to install two new menus. One menu provides dialog-box access to the immediate `epitab` commands, while the other is taken from the StataQuest additions to Stata, and provides access to a wide range of Stata's other commands for immediate statistical calculations.

Keywords: calculator commands, menus, epidemiologic statistics.

Introduction

Stata has long featured some very handy commands for performing a number of relatively simple statistical calculations that are commonly used in epidemiology (see [R] **epitab**). These commands calculate risk ratios, odds ratios (for unmatched and matched studies), and rate ratios, with appropriate confidence intervals and p -values (commands `cs`, `cc`, `mcc`, `ir`). The commands can be used on raw data, consisting of individual-level information on the presence or absence of "exposure" and "disease" in each subject in a study, or, in their immediate version (see [U] **Immediate commands**), on tabular data, where the subject-level

information has been collapsed to the form of a two-way table. The immediate versions of the commands provide a useful hand calculator approach to these calculations, but they are inconvenient for occasional users because of the necessity to remember (or look up in the help system) the correct order in which arguments need to be presented to the command. This insert describes a command, `calcmenu`, that uses Stata's menu-and-dialog programming features to install two new menus. One of these provides dialog-box access to the immediate `epitab` commands, while the other is taken from the StataQuest (Anagnoson and DeLeon, 1996) additions to Stata, and provides access to a wide range of Stata's other commands for immediate statistical calculations.

Syntax

```
calcmenu [on] [off]
```

Remarks

The use of this command and the menus is self-explanatory and does not seem to require an example. It should be noted that once a dialog box for one of the immediate commands is started (by selecting from the appropriate menu), it remains visible, with the command window unavailable until the cancel button is clicked. This allows the command to be run repeatedly with one or two input specifications changed for each run.

In the (unlikely) event that the results window fills before a command is completed, control passes to Stata's more system. The more condition must be cleared by using the go button on the button bar, since the command window is not available for keyboard input. Similarly, commands can only be interrupted with the break button, not with the usual keyboard alternative.

Finally, `calcmenu` cannot at present be run simultaneously with the Quest menus, and problems arise if you attempt to run `quest` with `calcmenu` on.

References

Anagnoson, J. T. and R. E. DeLeon. 1996. *StataQuest 4*. Belmont, CA: Duxbury Press.

sbe36	Summary statistics report for diagnostic tests
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Aurelio Tobias, Universidad Miguel Hernandez, Alicante, Spain, bledatobias@ctv.es

Abstract: Diagnostic tests are commonly used to classify patients into two groups according to the presence or absence of a symptom. Their results, which directly come from a 2×2 table, are summarized in terms of sensitivity, specificity, and predictive values.

Keywords: diagnostic test, sensitivity, specificity, predictive values, confidence interval, contingency tables.

Introduction

The simplest diagnostic test is one where the results of an investigation are used to classify patients into two groups according to the presence or absence of a symptom or sign. We can do this in Stata by using the `lstat` command after a logistic regression. For an epidemiologist, however, it seems to be more intuitive and easier to understand if such a result comes directly from a 2×2 table. In this insert I present a simple program named `diagtest` that summarizes in an easy way the classical results for a diagnostic test, say sensitivity, specificity, and predictive values, with their respective confidence intervals.

Methods

Sensitivity/specificity is one approach to quantifying the diagnostic ability of a test. Sensitivity is the proportion of true positives that are correctly identified by the test, while specificity is the proportion of true negatives that are correctly identified by the test. Sensitivity and specificity are proportions, so confidence intervals can be calculated for them using standard methods for proportions (Gardner and Altman 1989).

In general practice, however, the result of the diagnostic test is all that is known. The point of a diagnostic test is to use it to make a diagnosis; we need to know the probability that the test will give the correct diagnosis, and sensitivity and specificity do not give us this information. Instead, we must approach the data from the direction of the test results, using predictive values. The positive predictive value (PPV) is the proportion of patients with positive test results who are correctly diagnosed. The negative predictive value (NPV) is the proportion of patients with negative test results who are correctly diagnosed.

Test result	Pathology		
	Normal(-)	Abnormal(+)	
Normal(-)	a	b	$\implies NPV = a/(a+b)$
Abnormal(+)	c	d	$\implies PPV = d/(c+b)$
	↓	↓	
	Specificity = $a/(a+c)$	Sensitivity = $d/(b+d)$	

Table 1. Definition of sensitivity, specificity and predictive values.

The predictive values of a test in clinical practice depend critically on the prevalence of the abnormality in the patients being tested; this may well differ from the prevalence in a published study assessing the usefulness of the test. If the prevalence of the disease is very low, the positive predictive value will not be close to one even if both the sensitivity and specificity are high. Also, the prevalence can be interpreted as the probability before the test is carried out that the subject has the disease. The predictive values are the revised estimates of the same probability for those subjects who are positive and negative on the test, and are known as posterior probabilities.

The predictive values (PPV and NPV) can be calculated for any prevalence by using Bayes' theorem, as follows:

$$PPV = \frac{\text{Sensitivity} \times \text{Prevalence}}{\text{Sensitivity} \times \text{Prevalence} + (1 - \text{Sensitivity}) \times (1 - \text{Prevalence})}$$

$$NPV = \frac{\text{Specificity} \times (1 - \text{Prevalence})}{\text{Specificity} \times (1 - \text{Prevalence}) + (1 - \text{Specificity}) \times \text{Prevalence}}$$

Syntax

The `diagtest` command displays various summary statistics for diagnostic tests: sensitivity, specificity, and predictive values, from a 2×2 table.

```
diagtest testvar diagvar [weight] [if exp] [in range] [ , prev(#) level(#) tabulate_options ]
```

where *testvar* is the variable which identifies the result of the diagnostic test, and *diagvar* is the variable which contains the real status of the patient. Note that the lower category must identify the nonexposed, the negative result of the test, or the false status of the patient.

`fweights` and `pweights` are allowed.

Options

`prev(#)` requests the estimated prevalence, in percent, of the exposure to provide the positive and negative predicted values based on Bayes' theorem. The default is the estimated prevalence through the data.

`level(#)` specifies the confidence level, in percent, for confidence intervals. The default is `level(95)` or as set by `set level`.

Note that all `tabulate` command options are available.

Example

I present an example by Altman and Bland (1994a, 1994b) on the relationship between the results of a liver scan test and the correct diagnosis (Drum and Christacopoulos 1972). The proportions that were correctly diagnosed by the scan were 89.53% for normal liver scan and 62.79% for those with an abnormal scan. Also, the proportion of correct diagnoses among the patients with abnormal liver scan test was 87.83%, and among the patients with normal liver scans, the proportion was 66.67%.

```
. diagtest test diag [fw=n]
-----
diagnostic |
  test |   true diagnostic
  result |   Normal   Abnormal |   Total
-----+-----+-----
  Normal |         54         27 |         81
  Abnormal |         32        231 |        263
-----+-----+-----
  Total |         86        258 |        344
True D defined as diag ~ = 0
[95% Conf. Inter.]
-----
Sensitivity          Pr( + | D)  89.53%    86.30%    92.77%
Specificity          Pr( - | ~D)  62.79%    57.68%    67.90%
```

Positive predictive value	Pr(D +)	87.83%	84.38%	91.29%
Negative predictive value	Pr(~D -)	66.67%	61.69%	71.65%

Prevalence	Pr(D)	75.00%	70.42%	79.58%

In the liver scan study, the percentage of abnormality was 75%. If the same test were used in a different clinical setting where the prevalence of abnormality was 0.25, we would have a positive predictive value of 44.51% and a negative predictive value of 94.74%.

```
. diagtest test diag [fw=n], prev(25)
```

diagnostic test result	true diagnostic		Total
	Normal	Abnormal	
Normal	54	27	81
Abnormal	32	231	263
Total	86	258	344

True D defined as diag ~ = 0 [95% Conf. Inter.]

Sensitivity	Pr(+ D)	89.53%	86.30%	92.77%
Specificity	Pr(- ~D)	62.79%	57.68%	67.90%
Positive predictive value	Pr(D +)	44.51%	.%	.%
Negative predictive value	Pr(~D -)	94.74%	.%	.%

Prevalence	Pr(D)	25.00%	.%	.%

References

- Altman, D. G. and J. M. Bland. 1994a. Diagnostic tests 1: sensitivity and specificity. *British Medical Journal* 308: 1552.
- . 1994b. Diagnostic tests 2: predictive values. *British Medical Journal* 309: 102.
- Drum, D. E. and J. S. Christacapoulos. 1972. Hepatic scintigraphy in clinical decision making. *Journal of Nuclear Medicine* 13: 908–915.
- Gardner, M. J. and D. G. Altman. 1989. Calculating confidence intervals for proportions and their differences. In *Statistics with Confidence*, ed. M. J. Gardner and D. G. Altman. London: BMJ Publishing Group.

sbe37

Special restrictions in multinomial logistic regression

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Abstract: Two Stata programs, `mclgen` and `mclest`, can be used to impose special restrictions on multinomial logistic models. MCL models (Multinomial Conditional Logit, Breen 1994) use a conditional logit program to estimate a multinomial logistic model, providing greater flexibility for restrictions on the dependent variable. `mclgen` transforms the data to the format required, `mclest` calls `clogit` to estimate the model. In addition, `mclest` can estimate stereotyped ordered regression models (Anderson 1984, DiPrete 1990) and the row and columns model 2 (Goodman 1979), which contain both linear and multiplicative terms.

Keywords: multinomial logistic, stereotyped ordered regression, row and columns model 2, mobility models.

Introduction

This insert describes two Stata programs, `mclgen` and `mclest`, for imposing special restrictions on multinomial logistic models. MCL stands for Multinomial Conditional Logit, a term coined by Breen (1994). An MCL model uses a conditional logit program to estimate a multinomial logistic model. This produces the same log likelihood, estimates, and standard errors, but allows greater flexibility in imposing constraints. The MCL approach makes it possible to impose different restrictions on the response variable for different independent variables. For example, linear logits could be imposed for certain independent variables and an unordered response for others. One specific application is to include models for the analysis of square tables, e.g., quasi-independence, uniform association, symmetric association, into a multinomial logistic model (Logan 1983, Breen 1994).

`mclest` can also estimate two types of models with both linear and multiplicative terms. The Stereotyped Ordered Regression model (SOR) estimates a metric for the dependent variable and a single parameter for each independent variable (Anderson 1984, DiPrete 1990). It is more flexible than `ologit` because it does not assume ordered categories, although it does assume that the response categories can be scaled on a single dimension. This makes it useful for "semi-ordered" variables such as occupation, where the rank of categories such as farmers is not altogether clear.

A second special model that can be estimated by `mclest` is the Row and Columns model 2 (Goodman 1979). This model, originally developed for loglinear analysis, estimates a metric for a categorical independent variable as well as the response variable. The effect of the independent variable can therefore be expressed through a single parameter. The SOR and RC2 models are estimated by iteratively running MCL models, taking first one element of the multiplicative terms as given, then the other.

Multinomial conditional logit models

Multinomial logistic models and conditional logit models are very similar. Any model that can be estimated by `mlogit` can also be estimated by `clogit`, but this involves extra steps that are unnecessary for typical multinomial models. In order to estimate the model with `clogit`, the data must first be transformed into a person/choice file, the format for McFadden's choice model. In a person/choice file, each respondent has a separate record for each category of the response variable. A stratifying variable indexes respondents, the response variable indexes response options, and a dichotomous variable indicates which response option is the respondent's actual choice. This dichotomous variable is entered as the dependent variable in `clogit` and the stratifying variable is specified in the `group` option. The response variable, which in a standard multinomial program would be the dependent variable, is now entered as an independent variable. Its main effects, using one of the categories as reference, correspond with the intercept term of a multinomial model. Interactions of the response variable with explanatory variables correspond with the effects of these variables.

The following example shows how the data can be transformed into a person/choice file and how an MNL model can be estimated using `clogit` (see for example, the Stata Version 3 Reference Manual). The data here are taken from the 1972–78 GSS data used by Logan (1983, 332–333) and contain 838 cases. The response variable is `occ` (occupational class) with 5 categories; 1: farm occupations, 2: operatives, service, and laborers, 3: craftsmen and kindred workers, 4: sales and clerical, 5: professional, technical, and managerial. There are two explanatory variables: `educ` (education in years) and `black` (race; 1:black, 0:non-black).

```
. use logan
. gen strata=_n
. expand 5
. sort strata
. gen respfact=mod(_n-1,5)+1
. gen didep=(occ==respfact)
. quietly replace occ=respfact
. xi: clogit didep i.occ i.occ|black i.occ|educ, strata(strata)
```

The first step in creating the person/choice file is to define the stratifying variable `strata` using the current case numbers. Next, `expand` is used to create a copy of each record for each of the response options. The data are then sorted so that each respondent's records are grouped together. The variable `respfact` is constructed with values 1 to 5 within each stratum in order to index response options. The variable `didep` is then created to indicate which record corresponds with the respondent's actual choice. Once this has been done, the response variable `occ` is no longer needed and its contents can be replaced by those of `respfact`. Of course, `respfact` could be used in the following instead, but this procedure using `occ` has the advantage that variable and value labels assigned to `occ` will be used in the output.

Once the person/choice file has been created, the multinomial logistic model can be estimated in `clogit`. `didep` is specified as the dependent variable and `strata` is entered in the `strata` option. The main effects of `occ` using the first category as reference correspond with the intercept term and interactions of `occ` with `educ` and `black` correspond with the effects of these two variables. `xi` will also include main effects of `educ` and `black` but these will be dropped by `clogit` due to the fact that they are constant within strata. Alternatively, `desmat` (Hendrickx 1999) can be used here to generate the design matrix. `desmat` provides greater flexibility in specifying interactions and contrasts and the companion program `desrep` can summarize the output using informative labels.

Using `mclgen` and `mclest`

The programs `mclgen` and `mclest` automate the above procedure. `mclgen` transforms the data into a person/choice file and `mclest` enters the dichotomous dependent variable and stratifying variable, then estimates the model. The necessary steps are now reduced to

```
. mclgen occ
. xi: mclest i.occ i.occ|black i.occ|educ
```

This provides the following output:

```
. mclgen occ
(3352 observations created)
Your response factor is occ with 5 categories.
Its main effects form the intercept of a multinomial logistic model,
interactions with independent variables form their effects.
```

```

. xi: mclest i.occ i.occ|black i.occ|educ
i.occ          Iocc_1-5      (naturally coded; Iocc_1 omitted)
i.occ|black    IoXbla_#     (coded as above)
i.occ|educ     IoXedu_#     (coded as above)
Note: educ omitted due to no within-group variance.
Note: black omitted due to no within-group variance.
Iteration 0:   log likelihood = -1223.0058
Iteration 1:   log likelihood = -1025.7296
Iteration 2:   log likelihood = -1009.3479
Iteration 3:   log likelihood = -1007.2919
Iteration 4:   log likelihood = -1007.1621
Iteration 5:   log likelihood = -1007.1614
Iteration 6:   log likelihood = -1007.1614
Conditional (fixed-effects) logistic regression   Number of obs   =       4190
                                                    LR chi2(12)     =       683.10
                                                    Prob > chi2     =       0.0000
                                                    Pseudo R2      =       0.2532
Log likelihood = -1007.1614
-----+-----
  _didep |          Coef.   Std. Err.      z    P>|z|     [95% Conf. Interval]
-----+-----
  Iocc_2 |    2.913547    1.373878     2.121  0.034     .2207963    5.606298
  Iocc_3 |    1.843265    1.381555     1.334  0.182    -1.8645331   4.551063
  Iocc_4 |   -3.138894    1.47574     -2.127  0.033    -6.031291   -1.2464979
  Iocc_5 |   -6.131355    1.441328    -4.254  0.000    -8.956306   -3.306405
IoXbla_2 |    1.305156    1.043259     1.251  0.211    -1.7395935   3.349906
IoXbla_3 |    .628363    1.055375     0.595  0.552    -1.440134    2.69686
IoXbla_4 |    .3326202    1.103486     0.301  0.763    -1.830173    2.495413
IoXbla_5 |   -.2258867    1.093162    -0.207  0.836    -2.368446    1.916672
IoXedu_2 |   -.0505162    .1108293    -0.456  0.649    -1.2677376    .1667052
IoXedu_3 |    .0386727    .1111411     0.348  0.728    -1.1791599    .2565054
IoXedu_4 |    .3692091    .1163028     3.175  0.002     .1412598    .5971583
IoXedu_5 |    .6439505    .1135536     5.671  0.000     .4213896    .8665115
-----+-----

```

The parameters, standard errors, and log likelihood are the same as those of a model estimated using `mlogit occ educ black, base(1)`. The number of observations reported is 5 times the sample size since the data have been transformed into a person/choice file. Note that the LR `chi2` value is the chi-squared improvement relative to a model where all response options are equally likely, not an intercept-only multinomial model. Likewise, the pseudo R2 value uses an equiprobability model as its baseline, not an intercept model.

For a standard multinomial model, the MCL approach is only cumbersome. The advantage of using it lies in the ability to impose different restrictions on the response variable for different independent variables, something that cannot be easily done using `mlogit`. One application of this is to specify models for square tables, such as quasi-independence or uniform association, in a multinomial model with continuous covariates (Logan 1983, Breen 1994). These models were developed as loglinear models with special restrictions on the interaction between the row and column variable. They can be recast as multinomial logistic models where the restrictions on the response (column) variable depend on the category of the row variable.

Specifying a square table design as an MCL model follows the same procedure as for a loglinear model. One creates interactions between dummy variables for the response variable and dummy variables for the categorical independent using nonstandard restrictions. This can be illustrated in the following example.

```

. use logan
. mclgen occ
. gen d1=(focc==1)*(occ==1)
. gen d2=(focc==2)*(occ==2)
. gen d3=(focc==3)*(occ==3)
. gen d4=(focc==4)*(occ==4)
. gen d5=(focc==5)*(occ==5)
. gen u=focc*occ
. xi: mclest i.occ d* u i.occ|black i.occ|educ

```

This model specifies a quasi-uniform association model between father's occupation `focc` and the response variable `occ`, with `educ` and `black` as covariates. The dummies `d1` through `d5` measure the likelihood that father and son have the same occupation. They can be seen as interactions of a dummy for `focc==j` and a single dummy for the response variable corresponding with the logit `focc==j` versus `focc~=j`. The variable `u` for uniform association can be seen as an interaction of `occ` using a linear logits restriction and treating `focc` as a continuous rather than a categorical variable. The linear logits restriction means that for this effect, a unit's change in `focc` will result in a constant change in the logit between any two adjacent categories of `occ`.

In general, any type of restriction can be applied to the response variable and the restriction type can be varied at will

for each independent variable. By applying the difference contrast to the response variable (Hendrickx 1999), an adjacent logits model could be estimated. Another application could be to apply linear logits for some independent variables, and standard logits for others.

Stereotyped ordered regression

`mclest` can also estimate two special designs incorporating both linear and multiplicative effects. One is the Stereotyped Ordered Regression model (Anderson 1984, DiPrete 1990). The SOR model is an alternative to the proportional odds model used in `ologit`. The SOR model estimates a scaling metric for the response factor based on the effects of independent variables. The model has a standard multinomial intercept with $J - 1$ parameters for a response variable with J categories. It estimates $J - 2$ independent scale values ϕ_j for the response factor and a single scaled beta parameter for each independent variable. This means that the SOR model is less parsimonious than the proportional odds model, since it has an extra $J - 2$ parameters for the scaling metric. On the other hand, the SOR model does not assume that the response categories are ordered, although it does assume that they can be ordered. This makes it particularly useful when the rank of one or more categories is not altogether clear.

The SOR model can be specified as

$$\log \left(\frac{P(Y = q)}{P(Y = r)} \right) = \text{logit} \left(\frac{q}{r} \right) = \alpha_q - \alpha_r + (\phi_q - \phi_r)(\beta_1 X_1 + \cdots + \beta_K X_K)$$

where Y is the response variable with categories $j = 1$ to J , q and r are categories of Y , α_j represents the intercept parameters with suitable restrictions, ϕ_j represents the scaling metric with suitable restrictions, X_1, \dots, X_K represent K independent variables, and the β_k 's represent parameters of the independent variables. Two restrictions must be placed on the scaling metric ϕ_j in order to identify the model. `mclest` sets the first value to 0 and the last value to 1 while estimating the model. For the final estimates, the scaling metric is also normalized, with a mean of zero and a sum of squares of one.

Compare the SOR model to a standard multinomial model:

$$\text{logit} \left(\frac{q}{r} \right) = \alpha_q - \alpha_r + (\beta_{q1} - \beta_{r1})X_1 + (\beta_{q2} - \beta_{r2})X_2 + \cdots + (\beta_{qK} - \beta_{rK})X_K$$

In a multinomial model, the difference between β_{qk} and β_{rk} shows how the logit of q/r is affected by X_k . In the SOR model, this effect is equal to $(\phi_q - \phi_r)\beta_k X_k$. The effect on the logit of any two outcomes in the SOR model is proportional for all independent variables. Differences between scale values indicate how strongly the logit for two options is affected by the independent variables. The greater the difference between scale values, the more the logit between two outcomes is affected by the independent variables. The β_k parameters show how independent variable X_k affects the logit of higher versus lower scores, where “higher” and “lower” are defined by the ϕ_j metric.

An SOR model can be requested by specifying a *varlist* in the `sort` option. An SOR model with only one X_k variable would be trivial and equivalent to standard multinomial model since it contains the same number of parameters. A simple SOR model with two variables could be specified as

```
. use logan
. mclgen occ
. xi: mclest i.occ, sort(educ black)
```

This model will contain 9 parameters: 4 intercept parameters, 3 independent ϕ_j parameters, and 2 β_k parameters. This is only 3 less than for an unrestricted multinomial model. However, the parsimony of a SOR model does increase as the number of X_k variables increases.

The SOR model contains both linear and multiplicative elements. To estimate it, `mclest` iteratively estimates MCL models, first taking the ϕ_j scaling metric as given and estimating the β_k parameters, then taking the β_k parameters as given and estimating the ϕ_j parameters. This continues until the change in log likelihood between successive MCL models is less than the value specified in the `sortol` option (default .0001) or the maximum number of iterations specified in the `sortiter` option is exceeded (default 10). The standard errors for effects are conditional, given the scaling metric ϕ_j . A likelihood ratio test is therefore advisable before drawing any definite conclusions on the significance of effects.

Row and columns model 2

A second special model that can be estimated by `mclest` is Goodman's (1979) Row and Columns model 2 (RC2). Originally developed for frequency tables, the RC2 model estimates scaling metrics for both the dependent variable and one of the independent variables. The association between the two variables can then be expressed through a single parameter μ . The scaling metric for the dependent variable is ϕ_j as in the SOR model and the scaling metric for the independent variable is σ_v . Two restrictions

must be imposed on ϕ_j and σ_v in order to identify the model. During estimation, `mclest` sets $\phi_1 = \sigma_1 = 0$ and $\phi_J = \sigma_V = 1$. The final estimates are also given for normalized scale values, with $\text{mean}(\phi_j) = \text{mean}(\sigma_v) = 0$ and $\text{SS}(\phi_j) = \text{SS}(\sigma_v) = 1$.

A model containing an RC2 effect could be specified as

$$\text{logit}\left(\frac{q}{r}\right) = \alpha_q - \alpha_r + (\phi_q - \phi_r)\mu\sigma_v$$

This model can be seen as the SOR effects of the categorical variable scaled by $\mu\sigma_v$. In fact, entering dummies for the categorical variable in a SOR model results in an equivalent model. Using the RC2 specification has the advantages that it expresses the effect of the categorical variable through a single parameter μ and allows a comparison between the scale for the response variable and that of the categorical independent.

A variation of the RC2 model is the Equal Row and Columns model 2 (EQRC2). As the name suggests, this model uses the same metric for both the response variable and the categorical independent.

$$\text{logit}\left(\frac{q}{r}\right) = \alpha_q - \alpha_r + (\phi_q - \phi_r)\mu\phi_v$$

This is basically the same model, except that the effects of the categorical variable are scaled by ϕ_v instead of σ_v thus saving $J - 2$ degrees of freedom.

Another variation implemented in `mclest` allows the association μ between the dependent and independent variable to vary by one or more other independent variables.

$$\text{logit}\left(\frac{q}{r}\right) = \alpha_q - \alpha_r + (\phi_q - \phi_r)(\mu_0 + \mu_t X_t)\mu\sigma_v$$

An overall association parameter μ_0 is estimated, together with μ_t parameters indicating how the association changes for each independent variable X_t , for t from 1 to T . This is basically a parsimonious interaction effect of the categorical variable and the X_t variables.

An RC2 model is requested by specifying a varname in the `rc2` option. At present, only one variable can be used for the RC2 effect. Similarly, an EQRC2 model can be requested by specifying a varname in the `eqrc2` option. The `rc2` and `eqrc2` options are mutually exclusive. To let the overall association vary by one or more independent variables, specify a varlist in the `muby` option.

Models containing RC2 or EQRC2 effects are estimated by iteratively running MCL models, as is the case for SOR models. Convergence criterion and maximum iterations are determined by the `sortol` and `sortiter` options. The standard errors for effects are conditional, given the scaling metrics ϕ_j and σ_v and ϕ_v . A likelihood-ratio test is therefore advisable before drawing any definite conclusions on the significance of effects.

The following example estimates a quasi-RC2 model for father's occupation, including effects for the likelihood of having the same occupation as father (`diag`) and an RC2 effect. The overall association μ between father's occupation and respondent's occupation is allowed to vary by race. Furthermore, race and education are included in the model as covariates using a SOR effect.

```
. use logan
. mclgen occ
. gen diag=(focc==occ)*focc
. xi: mclest i.occ i.diag, sor(educ black) rc2(focc) muby(black)
```

This produces the following results:

```
. xi: mclest i.occ i.diag, sor(educ black) rc2(focc) muby(black)
i.occ          Iocc_1-5      (naturally coded; Iocc_1 omitted)
i.diag         Idiag_0-5     (naturally coded; Idiag_0 omitted)
Estimating Stereotype Ordered Regression for educ black
Estimating rc2 effects for focc
mu varies by black

iteration      log likelihood      sub-changes      main changes
-----
1.1            -985.5668            -985.5668            -985.5668
1.2            -971.8872             13.6796            -971.8872
1.3            -971.4130              0.4742            -971.4130
2.1            -970.7999              0.6131             14.7669
```

2.2	-970.9284	-0.1285	0.9588
2.3	-970.7092	0.2192	0.7038
3.1	-970.7035	0.0057	0.0964
3.2	-970.6918	0.0117	0.2366
3.3	-970.6900	0.0018	0.0192
4.1	-970.6896	0.0004	0.0140
4.2	-970.6895	0.0001	0.0023
4.3	-970.6894	0.0002	0.0007
5.1	-970.6893	0.0000	0.0003
5.2	-970.6893	0.0000	0.0002
5.3	-970.6893	0.0000	0.0001

 Convergence criterion .0001 reached in 5 iterations

When the `muby` option is used in conjunction with an RC2 model, `mc1est` uses three subiterations per iteration. First ϕ_j is taken as given and β_k and the product $\sigma_v(\mu_0 + \mu_1 X_1)$ are estimated. Next, β_k and $\sigma_v(\mu_0 + \mu_1 X_1)$ are taken as given and ϕ_j is estimated. In the third subiteration, ϕ_j and σ_v are taken as given and β_k , μ_0 , and μ_1 are estimated. If the `muby` option is not used or if it is used in conjunction with an EQRC2 model, only two subiterations are used. The subchanges and main changes indicate the change in log likelihood between subiterations and main iterations respectively. The change between main iterations is the criterion for determining whether the model has converged.

Phi scale for occ

	Coef.

phi(1): Farm	0
phi(2): Operatives	-0.4818
phi(3): Craftsmen	-0.2938
phi(4): Sales	0.3653
phi(5): Professional	1

The ϕ_j scale uses the restriction that the first category is fixed to 0 and the last category is fixed to 1. Differences between ϕ_j scale values show how the logit of one occupation versus another is affected by education, race, and scaled father's occupation (controlling for the likelihood of having the same occupation as father). The presence of negative values show that respondents with a higher education, who are nonblack, who have a "well-placed" father, are more likely to become a farmer than either an operative or a craftsman. The greatest impact of the independent variables on a logit between adjacent categories is for professionals versus sales, where the difference between the scale values is .635. The smallest impact is on craftsmen versus operatives, a difference of only .188.

Sigma scale for focc:

	Coef.

sig(1): Farm	0
sig(2): Operatives	0.4401
sig(3): Craftsmen	0.3165
sig(4): Sales	0.6886
sig(5): Professional	1

The σ_v metric defines a "well-placed" father, in the context of the model and given the data. As a resource for obtaining a good position and controlling for the likelihood of having the same occupation as one's father, a farm occupation scores the lowest. An operative rather than a craftsman as father increases the likelihood of the respondent getting a better occupation, but the difference is rather small. A father who is a craftsman rather than a farmer, or a professional rather than a salesman, has the greatest impact of any two adjacent scale values on the respondent's occupation.

Mu scaled association between occ and focc:

	Coef.	Std. Err.	z	P> z

MU: RC2 association between occ and	1.0241	0.3551	2.8840	0.0039
MU by black: race	1.1291	1.0633	1.0619	0.2883

The μ_0 and μ_1 parameters indicate the magnitude of the effect of father's occupation on respondent's occupation. These standard errors given here are conditional on ϕ_j and σ_v . The estimates indicate that there is a strong association between father's

occupation and respondent's occupation but that this is not significantly different for blacks and nonblacks. The impact of having a father who is a professional rather than a salesman on the logit of becoming a professional rather than a salesman is $(1 - .689)1.024(1 - .365) = .202$ for nonblacks and $(1 - .689)(1.024 + 1.129)(1 - .365) = .426$ for blacks (noting that μ_1 is large but also has a large standard error and is not significant).

Beta parameters:

	Coef.	Std. Err.	z	P> z
SOR effect for educ: education in ye	0.4406	0.0343	12.8461	0.0000
SOR effect for black: race	-1.4045	0.5878	-2.3896	0.0169

These are the SOR effects of education and race. The impact of one year of education on the logit of becoming a professional versus a salesman is $.441(1 - .365) = .280$. The impact of being black rather than nonblack is $-1.405(1 - .365) = -.891$.

Full parameter listing:

Conditional (fixed-effects) logistic regression	Number of obs	=	4190
	LR chi2(13)	=	756.04
	Prob > chi2	=	0.0000
Log likelihood = -970.6893	Pseudo R2	=	0.2803

__didep	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]
Iocc_2	6.528278	.5511616	11.845	0.000	5.448021 7.608535
Iocc_3	5.436115	.5270407	10.314	0.000	4.403134 6.469096
Iocc_4	.8334484	.5414884	1.539	0.124	-.2278494 1.894746
Iocc_5	-2.480348	.7093216	-3.497	0.000	-3.870593 -1.090103
Idiag_1	3.442951	.5782425	5.954	0.000	2.309617 4.576285
Idiag_2	.5049115	.1881059	2.684	0.007	.1362307 .8735924
Idiag_3	.4020211	.1897849	2.118	0.034	.0300496 .7739927
Idiag_4	-.4116733	.3947225	-1.043	0.297	-1.185315 .3619686
Idiag_5	-.5496774	.2975123	-1.848	0.065	-1.132791 .0334361
__beta1	.4406277	.0343005	12.846	0.000	.3734001 .5078554
__beta2	-1.404548	.5877656	-2.390	0.017	-2.556548 -.252549
__mu	1.024126	.3551078	2.884	0.004	.3281276 1.720125
__muby1	1.129056	1.06328	1.062	0.288	-.9549348 3.213046

These are the estimates from the final subiteration. Standard errors are conditional, given ϕ_j and σ_v . The first four parameters are standard multinomial logistic intercepts. The *Idiag_** parameters measure the likelihood of the respondent having the same occupation as his father. Having a father who is a farmer has a tremendous impact on the logit for becoming a farmer versus any other occupation. The *Idiag_4* and *Idiag_5* parameters for sales and professional occupations are negative, reducing the impact of the RC2 effect if father and son both have either of these occupations. The remaining parameters have been treated above.

Normalized Solution:

Normalized phi scale for occ:

	Coef.
phi(1): Farm	-0.1003
phi(2): Operatives	-0.5101
phi(3): Craftsmen	-0.3502
phi(4): Sales	0.2104
phi(5): Professional	0.7502

Normalized Sigma scale for focc:

	Coef.
sig(1): Farm	-0.6465
sig(2): Operatives	-0.0647
sig(3): Craftsmen	-0.2281
sig(4): Sales	0.2638
sig(5): Professional	0.6755

Unless the `nonorm` option is used, `mclgen` also produces a normalized solution. The scaling metrics now have a mean of zero and a sum of squares of 1.

```

Iteration 0:  log likelihood = -1218.2183
Iteration 1:  log likelihood = -1000.3561
Iteration 2:  log likelihood = -976.65487
Iteration 3:  log likelihood = -971.5582
Iteration 4:  log likelihood = -970.73319
Iteration 5:  log likelihood = -970.6895
Iteration 6:  log likelihood = -970.6893

Conditional (fixed-effects) logistic regression   Number of obs   =       4190
                                                    LR chi2(13)     =       756.04
                                                    Prob > chi2     =       0.0000
                                                    Pseudo R2      =       0.2803

Log likelihood = -970.6893
-----+-----
  __didep |      Coef.   Std. Err.      z    P>|z|      [95% Conf. Interval]
-----+-----
   Iocc_2 |   6.286966   .5487816    11.456  0.000    5.211373   7.362558
   Iocc_3 |   5.288965   .5245917    10.082  0.000    4.260784   6.317145
   Iocc_4 |   1.016421   .5376269     1.891  0.059   - .0373089   2.07015
   Iocc_5 |  -1.979513   .7071723    -2.799  0.005   -3.365545  - .5934809
  Idiag_1 |   3.442951   .5782421     5.954  0.000    2.309617   4.576285
  Idiag_2 |   .5049115   .1881059     2.684  0.007    .1362307   .8735924
  Idiag_3 |   .4020211   .1897849     2.118  0.034    .0300496   .7739927
  Idiag_4 |  - .4116732   .3947225    -1.043  0.297   -1.185315   .3619686
  Idiag_5 |  - .5496775   .2975123    -1.848  0.065   -1.132791   .033436
  __beta1 |   .5180692   .0403289    12.846  0.000    .4390261   .5971123
  __beta2 |  -1.002211   .3355206    -2.987  0.003   -1.659819  - .3446024
  __mu    |   .9108703   .3158372     2.884  0.004    .2918408   1.5299
  __muby1 |   1.004196   .9456943     1.062  0.288   - .8493306   2.857723
-----+-----

```

These are the parameter estimates, given the normalized ϕ_j and σ_v scaling metrics. Which set of parameters is used is basically a matter of personal preference.

Syntax for `mclgen`

```
mclgen depvar
```

The argument `depvar`, a categorical response factor with a maximum of 12 levels, must be specified for use in the MCL model. Note that the `mclgen` program will modify the data and that the data should therefore be saved before running it. Besides transforming the data into a person/choice file, `mclgen` adds `__didep` and `__strata`, the dichotomous dependent variable and stratifying variable used by `clogit`. In addition, it defines the global macros `ncat` containing the number of categories of the response variable and `respfact`, the name of the response factor.

Syntax for `mclest`

```
mclest varlist [weight] [if exp] [in range] [, sor(varlist) soriter(#) sortol(#) rc2(varname)
           eqrc2(varname) muby(varlist) nonorm debug ]
```

`fweights` and `iweights` are allowed.

`mclest` is used to estimate a model after transforming the data to a “person/choice” file using `mclgen`. The `varlist` should contain dummies based on the response factor specified in `mclgen` and interactions of these dummies with independent variables. This design matrix can be specified using `xi` or `desmat` (Hendrickx 1999).

`mclest` passes the `weight`, `if`, and `in` arguments to `clogit` unaltered. See the Stata documentation on `clogit` for further details.

Options

The following options are used to request the special nonlinear models Stereotyped Ordered Regression (SOR) and/or the Row and Columns model 2 (RC2).

`sor(varlist)` specifies a list of variables for which the SOR constraint should be used. Note that at least two variables should be specified, unless either the `rc2` or `eqrc2` option is being used.

`soriter(#)` specifies the maximum number of iterations for estimating a SOR or RC2 model. The default value is 10.

`sortol(#)` specifies the convergence criterion for estimating a SOR or RC2 model. The default value is .0001.

`rc2(varname)` specifies a categorical independent variable for which the RC2 model is to be used. The `eqrc2` option will be ignored if the `rc2` option is specified.

`eqrc2(varname)` specifies a categorical independent variable for which the EQRC2 model is to be used. The `rc2` option may not be used together with the `eqrc2` option.

`muby(varlist)` specifies one or more variables that affect the association between the `rc2` or `eqrc2` variable and the dependent variable. This option is ignored if not used in conjunction with the `rc2` or `eqrc2` option.

`nonorm` prevents the `mclrest` program from estimating a normalized solution if a SOR and/or RC2 model has been requested.

`debug` prints intermediate results of `clogit`. This can be used to determine the source of error if something goes wrong.

Recovering data

`mclgen` and `mclrest` create a number of variables for internal use when estimating models. To transform the person/choice file back to its original form, specify

```
. keep if __didep==1
. drop __*
```

References

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sg143	Cronbach's alpha one-sided confidence interval
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Abstract: Cronbach's alpha has been commonly used to assess the reliability for summative rating scales. However, Cronbach's alpha is a lower bound of reliability, thus a one-sided confidence interval can be calculated. The `cialpha` command computes the Cronbach's alpha one-sided confidence interval.

Keywords: Cronbach's alpha, reliability, rating scales, confidence interval.

Introduction

Cronbach's alpha (Cronbach 1951) assesses the reliability of a summative rating scale composed of the items specified. This can be easily computed in Stata using the `alpha` command. It is also demonstrated that Cronbach's alpha is always a lower bound of reliability (Guttman 1953). Thus, a one-sided confidence interval can be easily computed (Kristoff 1963, Feldt 1965). Although this result is well known and confidence intervals are commonly used in statistics, it is rarely applied by researchers evaluating instruments yielding a continuous measure (Bravo and Potvin 1991). In this way, Stata usually provides confidence intervals with most parameter estimates, but a confidence interval for Cronbach's alpha is not provided by the `alpha` command. In this insert we provide the command `cialpha`, which computes the Cronbach's alpha one-sided confidence interval.

Method

Kristoff (1963) and Feldt (1965) independently derived a transformation of the sample Cronbach's alpha coefficient, which they showed has an F distribution. From their sampling theory, a one-sided confidence interval can be shown to be given by

$$\alpha_X \geq 1 - \left[(1 - \hat{\alpha}_X) F_{a, n-1, (k-1)(n-1)} \right] = \alpha_L$$

This value enables one to determine whether the value achieved is significantly greater than some minimum value.

Syntax

```
cialpha [ , level(#) ]
```

`cialpha` can only be run after the `alpha` command, see [R] [alpha](#).

Options

`level(#)` specifies the confidence level, in percent, for confidence intervals. The default is `level(95)` or as set by `set level`.

Example

To illustrate the `cialpha` command, we consider Stata's automobile data. The scale derived from the arbitrarily chosen automobile items appears rather good, since `alpha` gives a value of 0.8975.

```
. alpha price hroom rep78 trunk weight length turn displ, std
Test scale = mean(standardized items)
Reversed item: rep78
Average interitem correlation:      0.5225
Number of items in the scale:      8
Scale Reliability Coefficient:      0.8975
```

Then, the 95% one-sided confidence interval for Cronbach's alpha has lower limit 0.8651.

```
. cialpha
Cronbach's alpha one-sided confidence interval
-----
Items | alpha          [95% Conf.Interval]
-----+-----
Test  | .89746938  >=    .86513734
-----
```

This means that there is a 95% chance that the Cronbach's alpha will be higher than this value.

Saved results

`cialpha` saves in `r()`

<code>r(alpha)</code>	scale reliability coefficient	<code>r(ci)</code>	one-sided confidence interval
<code>r(k)</code>	number of items in the table	<code>r(level)</code>	specifies the confidence level
<code>r(n)</code>	number of observations		

References

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Marginal effects of the tobit model

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Abstract: This article describes the `dtobit` command which provides a table of four forms of marginal effects for a tobit model. The marginal effects are computed for the latent dependent variable, the expected value of the dependent variable conditional on being uncensored, the unconditional expected value of the dependent variable, and the probability of being uncensored. The article also includes a brief discussion of the McDonald and Moffitt decomposition. Several examples are used to illustrate how this command can be run and how to interpret the output.

Keywords: tobit model, marginal effects of the tobit model, McDonald and Moffitt decomposition.

Syntax

```
dtobit [ , at(matname) censor brief nodiscrete level(#) ]
```

`dtobit` is for use after estimation by `tobit`; see [R] [tobit](#).

Description

`dtobit` provides a table of marginal effects evaluated at the means of the independent variables by default, or at the observed censoring rate of the dependent variable if `censor` is specified, or at user specified points if `at(matname)` is specified. The marginal effects are computed for the latent dependent variable, the expected value of the dependent variable conditional on being uncensored, the unconditional expected value of the dependent variable, and the probability of being uncensored. For dummy variables, `dtobit` will report the discrete changes unless the `nodiscrete` option is specified.

Options

`at(matname)` specifies the points around which the marginal effects are to be estimated. The default is to estimate the effects around the means of the independent variables.

`censor` specifies that the marginal effects should be evaluated at the observed censoring rate of the dependent variable. This option is not allowed with the two-tailed censoring case. `at(matname)` and `censor` cannot be specified at the same time.

`brief` will simplify the output. Standard deviations of the estimates and the confidence intervals will be excluded from the table.

`nodiscrete` treats dummy variables as continuous. If `nodiscrete` is not specified, the marginal effect of a dummy variable is calculated as the discrete change in the expected value of the dependent variable as the dummy variable changes from 0 to 1.

`level(#)` specifies the confidence level, in percent, for confidence intervals. The default is `level(95)` or as set by `set level`; see [U] [23.5 Specifying the width of confidence intervals](#).

Marginal effects of the tobit model

Let $y = \mathbf{X}\beta + \epsilon$ be the latent regression model. y represents continuous outcomes either observed or not observed. Our model assumes that $\epsilon \sim N(\mathbf{0}, \sigma^2 \mathbf{I})$.

Let a be the lower censored limit, and b be the upper censored limit. The tobit model can be expressed by the following relationship:

$$y_i^* = \begin{cases} y_i & \text{if } a < y_i < b \\ a & \text{if } y_i \leq a \\ b & \text{if } y_i \geq b \end{cases}$$

For the tobit model, there are four forms of marginal effects that are of great interest:

1. the β coefficients themselves are the changes in the mean of the latent dependent variable: $\beta = \partial E(y_i) / \partial x_i$
2. the changes in the unconditional expected value of the observed dependent variable: $\partial E(y_i^*) / \partial x_i$
3. the changes in the conditional expected value of the dependent variable: $\partial E(y_i^* \mid a < y_i^* < b) / \partial x_i$
4. the changes in the probability of being uncensored: $\partial P(a < y_i^* < b) / \partial x_i$

Example

We will demonstrate `dtobit` using an admittedly contrived example with the auto dataset. First, we run `tobit`. Assume that our data is censored in the sense that we cannot observe a mileage rating below 17 mpg, that is, if the true mpg is 17 or less, all we know is that the mpg is less than or equal to 17.

```
. use auto, clear
. tobit mpg trunk weight, ll(17)
Tobit estimates
```

Number of obs	=	74
LR chi2(2)	=	73.86
Prob > chi2	=	0.0000
Pseudo R2	=	0.1840

```
Log likelihood = -163.74787
```

mpg	Coef.	Std. Err.	t	P> t	[95% Conf. Interval]	
trunk	-.1487203	.1477163	-1.007	0.317	-.4431875	.1457468
weight	-.0063328	.0008709	-7.271	0.000	-.008069	-.0045966
_cons	41.90603	2.094632	20.006	0.000	37.73045	46.0816
_se	3.822488	.3637359	(Ancillary parameter)			

Obs. summary: 18 left-censored observations at mpg<=17
 56 uncensored observations

tobit reports the β coefficients of the latent regression model. dtobit, by default, reports all four forms of marginal effects at the means of the independent variables. (This includes the β coefficients which are the marginal effects on the latent dependent variable.)

```
. dtobit
```

Marginal Effects: Latent Variable

variable	dF/dx	Std. Err.	z	P> z	X_at	[95% C.I.]	
trunk	-.1487203	.1477163	-1.01	0.314	13.7568	-.438239	.140798
weight	-.0063328	.0008709	-7.27	0.000	3019.46	-.00804	-.004626
_cons	41.90603	2.094632	20.01	0.000	1.00000	37.8006	46.0114

Marginal Effects: Unconditional Expected Value

variable	dF/dx	Std. Err.	z	P> z	X_at	[95% C.I.]	
trunk	-.1243259	.1234866	-1.01	0.314	13.7568	-.366355	.117703
weight	-.005294	.0007281	-7.27	0.000	3019.46	-.006721	-.003867
_cons	35.03223	1.751052	20.01	0.000	1.00000	31.6002	38.4642

Marginal Effects: Conditional on being Uncensored

variable	dF/dx	Std. Err.	z	P> z	X_at	[95% C.I.]	
trunk	-.0926812	.0920555	-1.01	0.314	13.7568	-.273107	.087744
weight	-.0039465	.0005428	-7.27	0.000	3019.46	-.00501	-.002883
_cons	26.11546	1.305356	20.01	0.000	1.00000	23.557	28.6739

Marginal Effects: Probability Uncensored

variable	dF/dx	Std. Err.	z	P> z	X_at	[95% C.I.]	
trunk	-.009621	.0095561	-1.01	0.314	13.7568	-.028351	.009109
weight	-.0004097	.0000563	-7.27	0.000	3019.46	-.00052	-.000299
_cons	2.710991	.1355063	20.01	0.000	1.00000	2.4454	2.97658

The brief option will simplify the output:

```
. dtobit, brief
```

Name	Marginal Effects at Means			
	Latent Variable	Unconditional Expected Value	Conditional on being Uncensored	Probability Uncensored
trunk	-.14872035	-.12432593	-.09268119	-.00962104
weight	-.00633279	-.00529403	-.00394654	-.00040968
_cons	41.906025	35.032232	26.115461	2.7109914

To obtain marginal effects at the observed censoring rate, i.e., the fraction of uncensored samples, specify the `tensor` option. Note that the `tensor` option is not allowed with the two-tailed censoring case; see *Methods and formulas* for the underlying reason.

```
. dtobit, censor brief
```

Marginal Effects at Observed Censoring Rate				
Name	Latent Variable	Unconditional Expected Value	Conditional on being Uncensored	Probability Uncensored
trunk	-.14872035	-.11254513	-.08042806	-.01218351
weight	-.00633279	-.00479238	-.00342478	-.0005188
_cons	41.906025	31.712668	22.662807	3.4330384

To obtain marginal effects at particular points of the independent variables, specify the `at()` option. For example, suppose we would like to obtain marginal effects at `trunk = 12` and `weight = 2500`:

```
. mat M=(12, 2500, 1)
. dtobit, at(M) brief
```

Marginal Effects at e(at)				
Name	Latent Variable	Unconditional Expected Value	Conditional on being Uncensored	Probability Uncensored
trunk	-.14872035	-.14451729	-.12916254	-.00251919
weight	-.00633279	-.00615381	-.00549998	-.00010727
_cons	41.906025	40.721699	36.395078	.70985111

Example

Dummy variables are treated differently by `dtobit`. With dummy variables (or indicators), we are typically interested in the net effect on a form of the dependent variable when the indicator changes from 0 to 1; the marginal effect of a discrete change in the indicator variable. This is what `dtobit` calculates by default. If the instantaneous rate of change for a one-unit change in the indicator is desired, that is to say, if we want to treat the dummy variable like a continuous variable, we can specify the `nodiscrete` option.

In the `auto` dataset, `foreign` is a dummy variable. Assume that we are interested in its effect on mileage rating. What would be the difference in mileage between a foreign car (`foreign = 1`) and a domestic car (`foreign = 0`) given the same weight and trunk space?

```
. tobit mpg trunk weight for , ll(17)
```

```
Tobit estimates                               Number of obs =      74
                                                LR chi2(3)       =    77.09
                                                Prob > chi2     =     0.0000
Log likelihood = -162.13558                    Pseudo R2       =     0.1921
```

mpg	Coef.	Std. Err.	t	P> t	[95% Conf. Interval]	
trunk	-.1288913	.1458352	-0.884	0.380	-.4196784	.1618959
weight	-.0072948	.0010278	-7.097	0.000	-.0093442	-.0052454
foreign	-2.249467	1.2558	-1.791	0.078	-4.753461	.2545274
_cons	45.15628	2.794429	16.159	0.000	39.58434	50.72821
_se	3.763147	.3574121	(Ancillary parameter)			

```
Obs. summary:      18 left-censored observations at mpg<=17
                   56 uncensored observations
```

```
. dtobit, brief
```

Marginal Effects at Means				
Name	Latent Variable	Unconditional Expected Value	Conditional on being Uncensored	Probability Uncensored
trunk	-.12889128	-.10781396	-.08040115	-.00845294
weight	-.00729477	-.00610187	-.00455041	-.00047841
foreign*	-2.2494667	-2.171794	-1.8423142	-.16425745
_cons	45.156276	37.771965	28.168056	2.9614379

(*) dF/dx is for discrete change of dummy variable from 0 to 1

The results above indicate that we would expect a higher mileage rating in domestic cars than foreign cars given that `weight` and `trunk` space are taken at their means. A foreign car with mean weight and trunk space would have a 2.17 decrease in unconditional expected mileage and a 1.84 decrease in conditional expected mileage. Moreover, a typical domestic car has a .16 higher probability of being uncensored than that of a foreign car.

McDonald and Moffitt decomposition

A simple case of the tobit model, and probably the most useful one, is

$$y_i = \mathbf{X}\beta + \epsilon$$

$$y_i^* = \begin{cases} y_i & \text{if } y_i > 0 \\ 0 & \text{if } y_i \leq 0 \end{cases}$$

A very interesting feature of this simplified tobit model was discovered by McDonald and Moffitt (1980). Consider the expectation of y^* :

$$E(y^*) = P(y^* > 0)E(y^*|y^* > 0)$$

The effect of a change in the j^{th} continuous variable of \mathbf{X} on $E(y^*)$ can be expressed as

$$\partial E(y^*)/\partial x_j = P(y^* > 0)\partial E(y^*|y^* > 0)/\partial x_j + E(y^*|y^* > 0)\partial P(y^* > 0)/\partial x_j$$

Thus the total change in the unconditional expected value of y^* can be decomposed into two very intuitive parts:

1. the change in the expected value of y^* of those above zero, weighted by the probability of being above zero;
2. the change in the probability of being above zero, weighted by the conditional expected value of y^* .

Example

To illustrate the decomposition, we need to modify our auto dataset.

```
. replace mpg = mpg - 17
. tobit mpg trunk weight, ll(0)
Tobit estimates
Log likelihood = -163.74787
```

	Number of obs	=	74
	LR chi2(2)	=	73.86
	Prob > chi2	=	0.0000
	Pseudo R2	=	0.1840

```
-----+-----
```

mpg	Coef.	Std. Err.	t	P> t	[95% Conf. Interval]	
trunk	-.1487203	.1477163	-1.007	0.317	-.4431875	.1457468
weight	-.0063328	.0008709	-7.271	0.000	-.008069	-.0045966
_cons	24.90603	2.094632	11.890	0.000	20.73045	29.0816

```
-----+-----
      _se | 3.822488 .3637359 (Ancillary parameter)
-----+-----
```

```
Obs. summary:      18 left-censored observations at mpg<=0
                  56 uncensored observations
. dtobit, brief
```

```
-----+-----
```

Marginal Effects at Means				
Name	Latent Variable	Unconditional Expected Value	Conditional on being Uncensored	Probability Uncensored
trunk	-.14872035	-.12432593	-.09268119	-.00962104
weight	-.00633279	-.00529403	-.00394654	-.00040968
_cons	24.906025	20.820721	15.521213	1.6112246

```
-----+-----
```

In order to numerically show that the decomposition holds, we will use `predict, e(0,.)` to get the conditional expected value at the means of trunk and weight, and `predict, p(0,.)` to get the probability of being above zero. We will use an added observation at the end of the data to create a record of mean values, the point where we have evaluated the marginal effects.

```

. mat list e(at)
e(at)[1,3]
      trunk      weight      _cons
_cons 13.756757 3019.4595      1
. set obs 75
obs was 74, now 75
. replace trunk = 13.756757 in 75
trunk was int now float
(1 real change made)
. replace weight = 3019.4595 in 75
weight was int now float
(1 real change made)
. predict e, e(0,.)
. predict p, p(0,.)
. list e in 75 /* the conditional expected value */
      e
75. 4.869236
. list p in 75 /* the probability for above zero */
      p
75. .8359712

```

Now let's see if the decomposition holds:

```

. display -.09268119*.8359712 + (-.00962104)*4.869236
-.12432592

```

As we expected, the decomposition recovers the change in the unconditional expected value of mileage.

Technical notes

The application of the decomposition was well illustrated in McDonald and Moffitt's paper. There is no doubt that the decomposition is useful. However, the decomposition will only hold when the limit is zero, and when the variable is continuous. Let's consider both of these cases.

Decomposition with dummy variables

For dummy variables, the decomposition no longer holds. However, we can show a similar decomposition. Consider the marginal effects of dummy variable X_j :

$$\begin{aligned}
 E_1(y^*) - E_0(y^*) &= P_1(y^* > 0)E_1(y^*|y^* > 0) - P_0(y^* > 0)E_0(y^*|y^* > 0) \\
 &= P_1(y^* > 0) [E_1(y^*|y^* > 0) - E_0(y^*|y^* > 0)] + E_0(y^*|y^* > 0) [P_1(y^* > 0) - P_0(y^* > 0)] \\
 &= P_0(y^* > 0) [E_1(y^*|y^* > 0) - E_0(y^*|y^* > 0)] + E_1(y^*|y^* > 0) [P_1(y^* > 0) - P_0(y^* > 0)]
 \end{aligned}$$

where

P_1	is the probability evaluated at $x_j = 1$
P_0	is the probability evaluated at $x_j = 0$
E_1	is the expected value evaluated at $x_j = 1$
E_0	is the expected value evaluated at $x_j = 0$

Thus the total change in the unconditional expected value of y^* can also be decomposed into two parts:

1. the change in the expected value of y^* of those above zero, weighted by the probability of being above zero given $x_j = 1$ (or $x_j = 0$);
2. the change in the probability of being above zero, weighted by the conditional expected value of y^* given $x_j = 0$ (or $x_j = 1$).

Decomposition with specified censoring points

For the more general tobit model where the lower limit is a and the upper limit is b , the expected value of y^* is

$$E(y^*) = P(y^* > 0)E(y^*|y^* > 0) + aP(y^* < a) + bP(y^* > b)$$

Thus, besides the change in conditional expected y^* and the change in the probability of being uncensored, we need also consider the change in the probability of being below a and the change in the probability of being above b .

Saved results

In addition to the standard saved results from `tobit`, `dtobit` saves in `e()`:

Scalars

`e(rate)` uncensored rate

Macros

`e(dummy)` 0 indicates continuous, 1 indicates dummy

Matrices

`e(b)` marginal effects for the latent variable

`e(se_c)` standard deviations of `e(df dx_c)`

`e(se_b)` standard deviations of `e(b)`

`e(df dx_p)` marginal effects for the uncensored probability

`e(df dx_u)` marginal effects for the unconditional expected value

`e(se_p)` standard deviations of `e(df dx_p)`

`e(se_u)` standard deviations of `e(df dx_u)`

`e(at)` points at which marginal effects were computed

`e(df dx_c)` marginal effects for the conditional expected value

Methods and formulas

`dtobit` is implemented as an ado-file.

Let $\mathbf{y} = \mathbf{X}\beta + \epsilon$ be the model. \mathbf{y} represents continuous outcomes either observed or not observed. Our model assumes that $\epsilon \sim N(\mathbf{0}, \sigma^2 \mathbf{I})$.

Let a be the lower limit, and b be the upper limit.

$$y_i^* = \begin{cases} y_i & \text{if } a < y_i < b \\ a & \text{if } y_i \leq a \\ b & \text{if } y_i \geq b \end{cases}$$

For the simple case where $a = 0$ and $b = +\infty$, let $z = \mathbf{X}\beta/\sigma$, $f(z)$ be the standard normal density, and $F(z)$ be the cumulative normal distribution function, then

$$\begin{aligned} E(y^*) &= \mathbf{X}\beta F(z) + \sigma f(z) \\ E(y^* | y^* > 0) &= \mathbf{X}\beta + \sigma f(z)/F(z) \\ P(y^* > 0) &= F(z) \end{aligned}$$

The marginal effects for continuous variable \mathbf{x}_j are

$$\begin{aligned} \partial E(y^*)/\partial x_j &= F(z)\beta_j && \text{(unconditional expected value)} \\ \partial E(y^* | y^* > 0)/\partial x_j &= \beta_j \left[1 - z f(z)/F(z) - f(z)^2/F(z)^2 \right] && \text{(conditional expected value)} \\ \partial P(y^* > 0)/\partial x_j &= f(z)\beta_j/\sigma && \text{(probability uncensored)} \end{aligned}$$

For the more general case, let $\alpha_a = (a - \mathbf{X}\beta)/\sigma$, $\alpha_b = (b - \mathbf{X}\beta)/\sigma$, and $f_j = f(j)$ $F_j = F(j)$. Then

$$\begin{aligned} E(y^*) &= aF_a + b(1 - F_b) + (F_b - F_a)\mathbf{X}\beta + \sigma(f_a - f_b) \\ E(y^* | a < y^* < b) &= \mathbf{X}\beta + \sigma(f_a - f_b)/(F_b - F_a) \\ P(a < y^* < b) &= F_b - F_a \end{aligned}$$

The marginal effects for continuous variable \mathbf{x}_j are

$$\begin{aligned} \partial E(y^*)/\partial x_j &= (F_b - F_a)\beta_j && \text{(unconditional expected value)} \\ \partial E(y^* | a < y^* < b)/\partial x_j &= \beta_j \left[1 - \frac{\alpha_a f_a - \alpha_b f_b}{F_a - F_b} - \left(\frac{f_a - f_b}{F_a - F_b} \right)^2 \right] && \text{(conditional expected value)} \\ \partial P(a < y^* < b)/\partial x_j &= \beta_j(f_a - f_b)/\sigma && \text{(probability censored)} \end{aligned}$$

`dtobit` by default, calculates marginal effects at the means of the independent variables; it also allows the user to specify points where marginal effects are to be evaluated. A special feature of `dtobit` is that the user can ask `dtobit` to calculate marginal effects at the uncensoring rate. The uncensoring rate is defined as the proportion of uncensored observations of the whole dataset. If censoring is one tailed, the value of $\mathbf{X}\beta$ can be recovered from the uncensoring rate (for example, if the lower limit is a , then $\mathbf{X}\beta = \sigma * \text{invnorm}(\text{uncensoring rate}) + a$), and is used for further calculation. However, when censoring is two tailed, the value of $\mathbf{X}\beta$ cannot be recovered; thus, the `sensor` option is not allowed with these cases.

For dummy variables, the marginal effects are calculated as the discrete change of the expected values (probability of uncensoring) as dummy variables change from 0 to 1.

Reference

McDonald, J. F. and R. A. Moffitt. 1980. The uses of tobit analysis. *The Review of Economics and Statistics* 62: 318–387.

sg145	Scalar measures of fit for regression models
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Abstract: `fitstat` is a post-estimation command that computes a variety of scalar measures of fit for assessing regression models for categorical, limited, and count dependent variables. It also computes the AIC and BIC measures which can be used for selecting among non-nested models. By using the `save` and `dif` options, it is simple to compare measures of fit across models.

Keywords: scalar measures of fit, AIC, BIC, pseudo- R^2 s, regression models.

Overview

Many scalar measures have been developed to summarize the overall goodness of fit for regression models of continuous, count, or categorical dependent variables. The post-estimation command `fitstat` calculates a large number of fit statistics for the estimation commands `clogit`, `cnreg`, `cloglog`, `gologit`, `intreg`, `logistic`, `logit`, `mlogit`, `nbreg`, `ologit`, `oprobit`, `omodel`, `poisson`, `probit`, `regress`, `zinb`, and `zip`. With its `saving()` and `using()` options, the command also allows the comparison of fit measures across two models. While `fitstat` duplicates some measures computed by other commands (e.g., the pseudo- R^2 in standard Stata output of `lfit`), `fitstat` adds many more measures and makes it convenient to compare measures across models. Details on the measures that are discussed below can be found in Long (1997) which cites the original sources for each measure and provides further details on their derivation.

Before proceeding, a word of caution regarding the use of these measures. A scalar measure of fit can be useful in comparing competing models and ultimately in selecting a final model. Within a substantive area of research, measures of fit can provide a *rough* index of whether a model is adequate. However, there is no convincing evidence that selecting a model that maximizes the value of a given measure results in a model that is optimal in any sense other than the model having a larger (or smaller) value of that measure. While measures of fit provide some information, it is only partial information that must be assessed within the context of the theory motivating the analysis, past research, and the estimated parameters of the model being considered.

Syntax

```
fitstat [ , saving(name) save using(name) dif bic force ]
```

While many measures of fit are based on values returned by the estimation command, for some measures it is necessary to compute additional statistics from the estimation sample. While `fitstat` does not include `if` and `in` options, analysis is based on the sample defined by `e(sample)` from the last model estimated. Accordingly, `fitstat` is appropriate for models estimated using `if` and `in` restrictions in the original model. `fitstat` can also be used when models are estimated with weighted data. Here there are two limitations. First, some measures cannot be computed with some types of weights. Second, with `pweights` we use values of the “pseudo-likelihoods” to compute our measures of fit. Given the heuristic nature of the various measures of fit, we see no reason why the resulting measures would be appropriate. Also, `fitstat` ends with an error if the last estimation command does not return a value for the log-likelihood equation with only an intercept (that is, if `e(ll_0)` returns a missing value). This will occur, for example, if the `noconstant` option is used with the estimation command.

Options

`saving(name)` saves the computed measures in a matrix for subsequent comparisons. *name* cannot be longer than 4 characters.

`save` is equivalent to `saving(0)`.

`using(name)` compares the fit measures for the current model with those of the model saved as *name*. *name* cannot be longer than 4 characters.

`dif` is equivalent to `using(0)`.

`bic` presents only BIC and other information measures. In comparing two models, `fitstat` reports the guidelines in Raftery (1996) for assessing the strength of one model over another.

`force` allows comparison of two models even when the number of observations or the estimation method varies between the two models.

Models and measures

Details on the measures of fit are given below. Here we only summarize which measures are computed for which models. Y indicates a measure is computed; N indicates the measure is not computed.

	regress	logistic logit probit	cloglog	ologit oprobit	clogit mlogit	cnreg intreg tobit	gologit nbreg poisson zinb zip
Log-likelihood	Y	Y	Y ¹	Y	Y	Y	Y ²
Deviance & LR chi-square	Y	Y	Y	Y	Y	Y	Y
AIC, AIC×n, BIC, BIC'	Y	Y	Y	Y	Y	Y	Y
R ² & Adjusted R ²	Y	N	N	N	N	N	N
Efron's R ²	N	Y	Y	N	N	N	N
McFadden's, ML, C&U's R ²	N	Y	Y	Y	Y	Y	Y
Count & Adjusted Count R ²	N	Y	Y	Y	Y ³	N	N
Var(e), Var(y*) and M&Z's R ²	N	Y	N	Y	N	Y	N

1. For `cloglog` the log likelihood for the intercept-only model does not correspond to the first step in the iterations.
2. For `zip` and `zinb`, the log likelihood for the intercepts-only model is calculated by estimating `zip|zinb lhs-variable, inf(_cons)`.
3. The adjusted count R² is not defined for `clogit`.

Example

To compute fit statistics for a single model:

```
. use mroz, clear
(PSID 1976 from T. Mroz)
. * compute fit statistics for a single model
. logit lfp k5 k618 age wc hc lwg inc
Iteration 0:  log likelihood = -514.8732
Iteration 1:  log likelihood = -454.32339
Iteration 2:  log likelihood = -452.64187
Iteration 3:  log likelihood = -452.63296
Iteration 4:  log likelihood = -452.63296

Logit estimates                               Number of obs =      753
LR chi2(7) = 124.48
Prob > chi2 = 0.0000
Pseudo R2 = 0.1209

Log likelihood = -452.63296
```

lfp	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
k5	-1.462913	.1970006	-7.426	0.000	-1.849027	-1.076799
k618	-.0645707	.0680008	-0.950	0.342	-.1978499	.0687085
age	-.0628706	.0127831	-4.918	0.000	-.0879249	-.0378162
wc	.8072738	.2299799	3.510	0.000	.3565215	1.258026
hc	.1117336	.2060397	0.542	0.588	-.2920969	.515564
lwg	.6046931	.1508176	4.009	0.000	.3090961	.9002901
inc	-.0344464	.0082084	-4.196	0.000	-.0505346	-.0183583
_cons	3.18214	.6443751	4.938	0.000	1.919188	4.445092

```
. fitstat
Measures of Fit for logit of lfp
Log-Lik Intercept Only:   -514.873   Log-Lik Full Model:   -452.633
D(745):                   905.266   LR(7):                124.480
                          Prob > LR:         0.000
McFadden's R2:           0.121   McFadden's Adj R2:   0.105
Maximum Likelihood R2:   0.152   Cragg & Uhler's R2:  0.204
McKelvey and Zavoina's R2: 0.217   Efron's R2:          0.155
Variance of y*:          4.203   Variance of error:   3.290
Count R2:                 0.693   Adj Count R2:        0.289
AIC:                      1.223   AIC*n:               921.266
BIC:                      -4029.663  BIC':                 -78.112
```

To compute and save fit measures:

```
. logit lfp k5 k618 age wc hc lwg inc
  (output same as above)
. fitstat, saving(mod1)
  (output same as above)
  (Indices saved in matrix fs_mod1)
```

To compare the saved model to the current model:

```
. logit lfp k5 age age2 wc inc
  (output omitted)
. fitstat, using(mod1)
Measures of Fit for logit of lfp
```

	Current	Saved	Difference
Model:	logit	logit	
N:	753	753	0
Log-Lik Intercept Only:	-514.873	-514.873	0.000
Log-Lik Full Model:	-461.653	-452.633	-9.020
D:	923.306(747)	905.266(745)	18.040(2)
LR:	106.441(5)	124.480(7)	-18.040(-2)
Prob > LR:	0.000	0.000	0.000
McFadden's R2:	0.103	0.121	-0.018
McFadden's Adj R2:	0.092	0.105	-0.014
Maximum Likelihood R2:	0.132	0.152	-0.021
Cragg & Uhler's R2:	0.177	0.204	-0.028
McKelvey and Zavoina's R2:	0.182	0.217	-0.035
Efron's R2:	0.135	0.155	-0.020
Variance of y*:	4.023	4.203	-0.180
Variance of error:	3.290	3.290	0.000
Count R2:	0.677	0.693	-0.016
Adj Count R2:	0.252	0.289	-0.037
AIC:	1.242	1.223	0.019
AIC*n:	935.306	921.266	14.040
BIC:	-4024.871	-4029.663	4.791
BIC':	-73.321	-78.112	4.791

Difference of 4.791 in BIC' provides positive support for saved model.

Saved results

fitstat saves in r() whichever of the following are computed for a particular model:

Scalars

r(aic)	AIC	r(n_rhs)	number of right hand side variables
r(aic_n)	AIC × n	r(r2)	R ² for linear regression model
r(bic)	BIC	r(r2_adj)	adjusted R ² for linear regression model
r(bic_p)	BIC'	r(r2_ct)	count R ²
r(dev)	deviance	r(r2_ctadj)	adjusted count R ²
r(dev_df)	degrees of freedom for deviance	r(r2_cu)	Cragg & Uhler's R ²
r(ll)	log-likelihood for full model	r(r2_ef)	Efron's R ²
r(ll_0)	log-likelihood for model with only intercept	r(r2_mz)	McKelvey and Zavoina's R ²
r(lrx2)	likelihood ratio chi-square	r(r2_mf)	McFadden's R ²
r(lrx2_df)	degrees of freedom for likelihood ratio chi-square	r(r2_mfadj)	McFadden's adjusted R ²
r(lrx2_p)	probability level of chi-square test	r(r2_ml)	maximum likelihood R ²
r(N)	number of observations	r(v_error)	variance of error term
r(n_parm)	number of parameters	r(v_ystar)	variance of y*

When the `saving(name)` option is specified, computed measures are also saved in matrix `fs_name`. The column names of the matrix correspond to the names of the measures listed above. The row name is the command used to estimate the saved model. Values of `-9999` in the matrix indicate that a measure is not appropriate for the given model. The row name is the name of the estimation procedure.

Extending `fitstat` to other models and measures

`fitstat` can be extended to other models and measures of fit. When doing this, there are several things to keep in mind. First, the program depends on values returned by `eclass` estimation commands (for example, `e(sample)`). Programs such as `ocratio` which do not use `eclass` returns cannot be incorporated into `fitstat` without major changes to the structure of the program. Second, not all measures of fit are appropriate for all models. The programmer must be careful to ensure that `fitstat` does not automatically compute inappropriate fit statistics. Third, the way in which values such as the number of parameters and the number of right-hand-side variables are computed differs across models. Consequently, additional code may be needed for these computations.

Methods and formulas

In this section we provide brief descriptions of each measure computed by `fitstat`. Full details along with citations to original sources are found in Long (1997). The measures are listed in the same order as the output illustrated above.

Log-likelihood based measures

Stata begins maximum likelihood iterations by computing the log likelihood of the model with all parameters except the intercept(s) constrained to zero, referred to as $L(M_{\text{Intercept}})$ below. The log likelihood upon convergence, referred to as M_{Full} below, is also listed. In Stata this information is usually presented as the first step of the iterations and in the header for the estimation results. Note that in `cloglog`, the value at iteration 0 is not the log likelihood with only the intercept. For `zip` and `zinb`, the intercept-only model can be defined in different ways. These commands return as `e(l1_0)` the value of the log likelihood with the binary portion of the model unrestricted while only the intercept is free for the Poisson or negative binomial portion of the model. Alternatively, `fitstat` returns the value of the log likelihood from the model with only an intercept in both the binary and count portion of the model.

Chi-square test of all coefficients

A likelihood-ratio test of the hypothesis that all coefficients except the intercept(s) can be computed by comparing the log likelihoods.

$$LR = 2 \ln L(M_{\text{Full}}) - 2 \ln L(M_{\text{Intercept}})$$

This statistic is sometimes designated as G^2 . LR is reported by Stata as: `LR chi2(7) = 124.48` where the degrees of freedom, (7), are the number of constrained parameters. `fitstat` reports this statistic as: `LR(7): 124.48`. For `zip` and `zinb`, LR tests that the coefficients in the count portion (not the binary portion) of the model are zero.

Deviance

The *deviance* compares a given model to a model that has one parameter for each observation and can reproduce perfectly the observed data. The deviance is defined as $D = -2 \ln L(M_{\text{Full}})$, where the degrees of freedom equals N minus the number of parameters. Note that D does not have a chi-square distribution.

Coefficient of determination in the LRM

For `regress`, `fitstat` reports the standard coefficient of determination which can be defined variously as

$$R^2 = 1 - \frac{\sum_{i=1}^N (y_i - \hat{y}_i)^2}{\sum_{i=1}^N (y_i - \bar{y})^2} = \frac{\widehat{\text{Var}}(\hat{y})}{\widehat{\text{Var}}(\hat{y}) + \widehat{\text{Var}}(\hat{\epsilon})} = 1 - \left[\frac{L(M_{\text{Intercept}})}{L(M_{\text{Full}})} \right]^{2/N}$$

The adjusted R^2 is defined as

$$\bar{R}^2 = \left(R^2 - \frac{K}{N-1} \right) \left(\frac{N-1}{N-K-1} \right)$$

where K is the number of independent variables.

Pseudo R-squared

While each of the definitions of R^2 above give the same numeric value in the LRM, they give different answers and thus provide different measures of fit when applied to the other models evaluated by `fitstat`.

McFadden's R-squared

The McFadden R^2 , also known as the “likelihood ratio index,” compares a model with just the intercept to a model with all parameters. It is defined as

$$R_{\text{McF}}^2 = 1 - \frac{\ln \widehat{L}(M_{\text{Full}})}{\ln \widehat{L}(M_{\text{Intercept}})}$$

If model $M_{\text{Full}} = M_{\text{Intercept}}$, R_{McF}^2 equals 0, but R_{McF}^2 can never exactly equal one. This measure, which is computed by Stata as `Pseudo R2 = 0.1209`, is listed in `fitstat` as: `McFadden's R2: 0.121`. Since R_{McF}^2 always increases as new variables are added, an adjusted version is also available

$$\bar{R}_{\text{McF}}^2 = 1 - \frac{\ln \widehat{L}(M_{\text{Full}}) - K^*}{\ln \widehat{L}(M_{\text{Intercept}})}$$

where K^* is the number of parameters (not independent variables).

Maximum likelihood R-squared

Another analogy to R^2 in the LRM was suggested by Maddala.

$$R_{\text{ML}}^2 = 1 - \left[\frac{L(M_{\text{Intercept}})}{L(M_{\text{Full}})} \right]^{2/N} = 1 - \exp(-G^2/N)$$

Cragg & Uhler's R-squared

Since R_{ML}^2 only reaches a maximum of $1 - L(M_{\text{Intercept}})^{2/N}$, Cragg and Uhler suggested a normed measure.

$$R_{\text{C\&U}}^2 = \frac{R_{\text{ML}}^2}{\max R_{\text{ML}}^2} = \frac{1 - [L(M_{\text{Intercept}})/L(M_{\text{Full}})]^{2/N}}{1 - L(M_{\text{Intercept}})^{2/N}}$$

Efron's R-squared

For binary outcomes, Efron's pseudo- R^2 defines $\widehat{y} = \widehat{\pi} = \widehat{\text{Pr}}(y = 1|x)$ and equals

$$R_{\text{Efron}}^2 = 1 - \frac{\sum_{i=1}^N (y_i - \widehat{\pi}_i)^2}{\sum_{i=1}^N (y_i - \widehat{y})^2}$$

McKelvey and Zavoina's R-squared

Some models can be defined in terms of a latent variable y^* . This includes the models for binary or ordinal outcomes; `logit`, `probit`, `ologit` and `oprobit`, as well as some models with censoring: `tobit`, `cnreg`, and `intreg`. Each model is defined in terms of a regression on a latent variable y^* :

$$y^* = x\beta + \epsilon$$

Using $\widehat{\text{Var}}(\widehat{y}^*) = \widehat{\beta}' \widehat{\text{Var}}(x) \widehat{\beta}$, McKelvey and Zavoina proposed

$$R_{\text{M\&Z}}^2 = \frac{\widehat{\text{Var}}(\widehat{y}^*)}{\widehat{\text{Var}}(y^*)} = \frac{\widehat{\text{Var}}(\widehat{y}^*)}{\widehat{\text{Var}}(\widehat{y}^*) + \text{Var}(\epsilon)}$$

In models for categorical outcomes, $\text{Var}(\epsilon)$ is assumed to identify the model; in models with censoring, it can be estimated.

Count and adjusted count R^2

Observed and predicted values can be used in models with categorical outcomes to compute what is known as the count R^2 . Consider the binary case where the observed y is 0 or 1 and $\pi_i = \Pr(y = 1|x_i)$. Define the expected outcome \hat{y} as

$$\hat{y}_i = \begin{cases} 0 & \text{if } \hat{\pi}_i \leq 0.5 \\ 1 & \text{if } \hat{\pi}_i > 0.5 \end{cases}$$

This allows us to construct a table of observed and predicted values, such as produced by the Stata command `lstat`:

Classified	True		Total
	D	~D	
+	342	145	487
-	86	180	266
Total	428	325	753

A seemingly appealing measure is the proportion of correct predictions, referred to as the *count* R^2 :

$$R^2_{\text{Count}} = \frac{1}{N} \sum_j n_{jj}$$

where the n_{jj} 's are the number of correct predictions for outcome j . The count R^2 can give the faulty impression that the model is predicting very well. In a binary model without knowledge about the independent variables, it is possible to correctly predict at least 50 percent of the cases by choosing the outcome category with the largest percentage of observed cases. To adjust for the largest row marginal, we have

$$R^2_{\text{AdjCount}} = \frac{\sum_j n_{jj} - \max_r(n_{r+})}{N - \max_r(n_{r+})}$$

where n_{r+} is the marginal for row r . The adjusted count R^2 is the proportion of correct guesses beyond the number that would be correctly guessed by choosing the largest marginal.

Information measures

This class of measures can be used to compare models across different samples or to compare nonnested models.

AIC

Akaike's information criteria is defined as

$$\text{AIC} = \frac{-2 \ln \hat{L}(M_k) + 2P}{N}$$

where $\hat{L}(M_k)$ is the likelihood of the model and P is the number of parameters in the model (for example, $K + 1$ in the binary regression model where K is the number of regressors). All else being equal, the model with the smaller AIC is considered the better fitting model. Some authors define AIC as being N times the value we report. We report this quantity as `AIC*n`.

BIC and BIC'

The Bayesian information criterion has been proposed by Raftery (1996 and the literature cited therein) as a measure of overall fit and a means to compare nested and nonnested models. Consider the model M_k with deviance $D(M_k)$. BIC is defined as:

$$\text{BIC}_k = D(M_k) - df_k \ln N$$

where df_k is the degrees of freedom associated with the deviance. The more negative the BIC_k is, the better the fit. A second version of BIC is based on the LR chi-square with df'_k equal to the number of regressors (not parameters) in the model. Then

$$\text{BIC}'_k = -G^2(M_k) + df'_k \ln N$$

The more negative the BIC'_k , the better the fit. The difference in the BICs from two models indicates which model is more likely to have generated the observed data. Since $\text{BIC}_1 - \text{BIC}_2 = \text{BIC}'_1 - \text{BIC}'_2$, the choice of which BIC measure to use is a matter of convenience. If $\text{BIC}_1 - \text{BIC}_2 < 0$, then the first model is preferred. If $\text{BIC}_1 - \text{BIC}_2 > 0$, the second model is preferred. Raftery

suggested guidelines for the strength of evidence favoring M_2 against M_1 based on a difference in BIC or BIC' as given in the table below.

Absolute	
Difference	Evidence
0–2	Weak
2–6	Positive
6–10	Strong
> 10	Very strong

Acknowledgments

We thank David M. Drukker, Senior Statistician at Stata Corporation for his helpful suggestions.

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sg146	Parameter estimation for the generalized extreme value distribution
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Abstract: In the analysis of extreme value data, such as wind speeds, air pollution levels, and river heights the generalized extreme value distribution (GEVD) plays a basic role. We present a program to fit the GEVD using the method of probability-weighted moments and the hybrid estimator proposed by Dupuis and Tsao.

Keywords: generalized extreme value distribution, extreme values, estimation, parameter, environmental statistics.

Introduction

In Scotto (2000) we considered the Gumbel distribution and its applications to environmental statistics. In this paper we turn our attention to the generalized extreme value distribution (GEVD).

Suppose we are interested in making inferences about extremal values of some random variables, for which we have a dataset available, in such a way that it is reasonable to identify it with X_1, \dots, X_n , an independent, identically distributed sample with underlying model F , and denote by M_r the sample maximum. According to classical extreme value theory, the distribution of linear functions of M_r can be approximated by taking limits as $n \rightarrow \infty$ and approximating for large but finite n , by the GEVD

$$G_{k,\zeta,\alpha}(x) = \exp \left\{ - \left[1 - \frac{k(x - \zeta)}{\alpha} \right]^{1/k} \right\}$$

where x is bounded by $\zeta + \alpha/k$ from above if $k > 0$ and from below if $k < 0$. As usual, the case $k = 0$ corresponds to the Gumbel distribution

$$G_{0,\zeta,\alpha} = \exp \left\{ - \exp \left\{ - \frac{x - \zeta}{\alpha} \right\} \right\}$$

The parameter k is a shape parameter which determines the weight of the tail of G , while ζ and α are location and scale parameters, respectively. The GEVD was introduced by Jenkinson (1955), and it has been successfully applied in modeling annual maximum river heights, sea levels, and air pollutants.

Our main concern here is to obtain point estimates for the parameters k , ζ , and α . Commonly, the method of probability-weighted moments (PWM) (Greenwood et al. 1975) has been used for this purpose. However, as is known, this method frequently leads to nonfeasible estimates in the sense that the supports inferred from the estimates fail to contain all observations. Dupuis and Tsao (1998) proposed the hybrid estimator which is derived by incorporating an auxiliary constraint on feasibility into the estimates from PWM to produce feasible estimates. Both methods have been implemented in the `gevd` command described below.

Method of probability-weighted moments

The method of probability-weighted moments consists of equating model moments based on $G_{k,\zeta,\alpha}$ to the corresponding empirical moments based on the data. The PWM method was developed by Greenwood et al. (1975). An overview of this method is as follows. The probability-weighted moments of a random variable X with distribution function F are defined as

$$M_{p,r,s} = E[X^p \{F(X)\}^r \{1 - F(X)\}^s]$$

where p , r and s are real numbers. To obtain PWM estimates for k , ζ , and α , it is convenient to fix $p = 1$ and $s = 0$ in the above expression for $M_{p,r,s}$ and use the quantities

$$\beta_s = E[X \{F(X)\}^s] = \frac{\zeta + \frac{\alpha}{k} \left\{ 1 - \frac{\Gamma(1+k)}{(r+1)^k} \right\}}{r+1}$$

which exist provided that $k > -1$. From this expression we obtain

$$\beta_0 = \zeta + \frac{\alpha}{k} \{1 - \Gamma(1+k)\} \quad (1)$$

$$2\beta_1 - \beta_0 = \frac{\alpha}{k} \Gamma(1+k)(1 - 2^{-k}) \quad (2)$$

$$\frac{3\beta_2 - \beta_1}{2\beta_1 - \beta_2} = \frac{1 - 3^{-k}}{1 - 2^{-k}} \quad (3)$$

The method consists in matching β_0 , β_1 , and β_2 with appropriate sample equivalents b_0 , b_1 , and b_2 in order to obtain the PWM estimators \hat{k} , $\hat{\zeta}$, and $\hat{\alpha}$. In general, b_i may be taken as

$$b_i = n^{-1} \sum_{j=1}^n w_j X_{j:n}$$

where $X_{1:n} \leq X_{2:n} \leq \dots \leq X_{n:n}$ are the ordered sample. The weights w_j are defined as

$$w_j = \frac{(j-1)(j-2)\cdots(j-i)}{(n-1)(n-2)\cdots(n-i)}$$

Clearly, equation (3) only depends on \hat{k} . The exact solution requires iterative methods, but since the function is almost linear over the range of interest ($-0.5 < k < 0.5$), which is usually encountered in practice, the following approximate low-order polynomial estimator (Hosking et al. 1985) is used

$$\hat{k} = 7.859c + 2.955c^2, \quad c = \frac{2b_1 - b_0}{3b_2 - b_0} - \frac{\ln 2}{\ln 3} \quad (4)$$

The error due to using (4) instead of (3) is less than 0.0009 throughout the range (Hosking et al. 1985). Given \hat{k} , the scale and location parameters can be estimated successively from (1) and (2) as

$$\hat{\alpha} = \frac{(2b_1 - b_0)\hat{k}}{\Gamma(1+\hat{k})(1 - 2^{-\hat{k}})} \hat{\zeta} = b_0 + \frac{\hat{\alpha}}{\hat{k}} \left\{ \Gamma(1+\hat{k}) - 1 \right\}$$

The parameter estimate of the bound is thus $\hat{\zeta} + \hat{\alpha}/\hat{k}$. The PWM estimates are not feasible when the following auxiliary constraints are violated

$$\begin{aligned} X_{n:n} &\leq \hat{\zeta} + \frac{\hat{\alpha}}{\hat{k}} > 0 \\ X_{1:n} &\geq \hat{\zeta} + \frac{\hat{\alpha}}{\hat{k}} < 0 \end{aligned} \quad (5)$$

(Dupuis and Tsao 1998). For further details of the method see Hosking et al. (1985). See also Dupuis (1996) for an exhaustive study of the percentage of nonfeasibility of the PWM estimators of the GEVD.

Hybrid estimator based on PWM

The hybrid estimator, proposed by Dupuis and Tsao (1998), and based on PWM and always satisfying (5) gives $\zeta_h = \hat{\zeta}$, $\alpha_h = \hat{\alpha}$, and

$$k_h = \begin{cases} \hat{k}\hat{\zeta} + \frac{\hat{\alpha}}{\hat{k}} > X_{n:n} & \text{and } \hat{k} > 0 \\ \hat{k}\hat{\zeta} + \frac{\hat{\alpha}}{\hat{k}} < X_{n:n} & \text{and } \hat{k} < 0 \\ \tilde{k} & \text{otherwise} \end{cases}$$

where

$$\tilde{k} = \begin{cases} \frac{\hat{\alpha}}{(X_{n:n} - \hat{\zeta})} & \hat{k} > 0 \\ \frac{\hat{\alpha}}{(X_{1:n} - \hat{\zeta})} & \hat{k} < 0 \end{cases}$$

Further results about the comparison between the hybrid estimator and the original PWM estimators in terms of bias and RMSE can be seen in Dupuis and Tsao (1998).

Syntax

```
gevd varname [if exp] [in range] [, centile(#) hybrid ]
```

Options

`centile(k)` provides the k estimated percentiles dividing the variable into k parts. For example, `centile(3)` return the tertile values, `centile(4)` the quartile values, and so on.

`hybrid` specifies that the estimation of the parameters of the GEVD is done by using the hybrid estimator.

Example

We fitted the GEVD to 22 annual maximum December temperatures at Fair Isle Weather Station, UK for 1974 through 1995, reported by Dupuis and Tsao (also available from the website <http://www.zetnet.co.uk/sigs/weather>).

```
. use weather
. describe
Contains data from weather.dta
obs:          22
vars:         1          19 Sep 1998 11:17
size:        176 (99.9% of memory free)
-----
1. temp      float %9.0g
-----
Sorted by:
. summarize temp, detail
-----+-----
                temp
-----+-----
Percentiles    Smallest
1%             7.5      7.5
5%             9        9
10%            9.2      9.2   Obs          22
25%            9.4      9.3   Sum of Wgt.  22
50%            10.2     Std. Dev.    9.977273
                    Largest  Std. Dev.    .8211095
75%            10.4     10.6
90%            10.7     10.7   Variance    .6742207
95%            10.9     10.9   Skewness    -1.060649
99%            11.5     11.5   Kurtosis    5.056575
```

Graphically, we can estimate the density of these annual maximum temperature levels using Stata's `kdensity` command giving the result in Figure 1.

```
. kdensity temp, xlab ylab
```

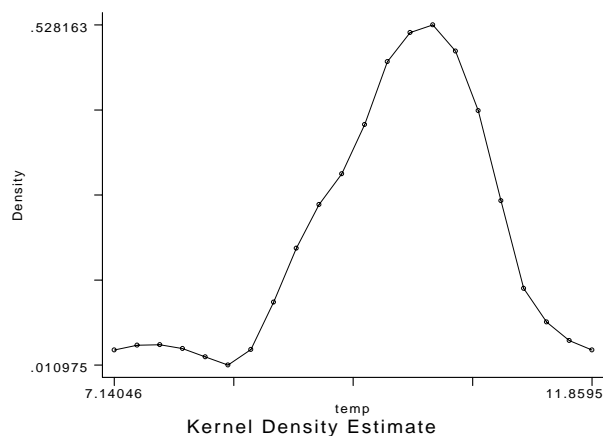


Figure 1: Density estimate of annual maximum temperatures.

Dupuis and Tsao fitted the GEVD, obtaining $\hat{\alpha} = 0.874$, $\hat{k} = 0.635$, and $\hat{\zeta} = 9.837$ as PWM estimates. They obtained $\hat{k} = 0.525$ when they used the hybrid estimator based on PWM.

Using `gevd` we obtained the following PWM estimates

```
. gevd temp
Parameter estimation for GEVD distribution by using
Probability Weighted Moments (PWM)
Variable | alpha      k      Psi
-----+-----
temp     | .8739451   .63474808  9.8366951
```

and, optionally, for the hybrid estimator we found

```
. gevd temp, hybrid
Parameter estimation for GEVD distribution by using
Hybrid estimator based on PWM
Variable | alpha      k      Psi
-----+-----
temp     | .8739451   .52542688  9.8366951
```

Estimates using the `gevd` command strongly agree with values obtained by Dupuis and Tsao for both methods.

Saved results

`gevd` saves the estimates of k , ζ and α in the global macros `S_k`, `S_psi`, and `S_alpha`, respectively.

References

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sg147

Hill estimator for the index of regular variation

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Abstract: The Hill estimator for the index of regular variation is presented. We also pay attention to the Hill plot since the statistical analysis based on the Hill estimator is usually summarized graphically.

Keywords: index of regular variation, Hill estimator, Hill plot.

Introduction

Assume we have iid random variables with distribution function F which belongs to the maximum domain of attraction of the Fréchet distribution, that is,

$$P(X > x) = x^{-\alpha}L(x), \quad x > 0$$

where α is the index of regular variation, and L is some slowly varying function. Distributions with such tails form the prime examples for modeling heavy-tailed phenomena. For many applications the knowledge of the index α is of major importance. We use the well-known Hill estimator (Hill 1975) to obtain an estimate of α .

The Hill estimator takes the form

$$\hat{\alpha}_{k,n} = \left[\frac{1}{n} \sum_{j=1}^k \ln X_{j,n} - \ln X_{k,n} \right]^{-1}$$

where $X_{j,n}$ is the j th upper order statistic and $k = k(n) \rightarrow \infty$ in an appropriate way. Note that k is the number of upper order statistics used in the estimation. The rough idea behind using only k upper order statistics is that you should have only sampled from that part of the distribution which looks most Fréchet-like.

The theoretical properties of the Hill estimator are given in Resnick (1997). The main difficulty when using the Hill estimator is that it involves a sequence of upper order statistics increasing with n . But the question arising is, what value of k do you use? Unfortunately, the optimal choice of k is a delicate point for which no uniformly best solution exists. In practice, an analysis based on the Hill estimator is usually summarized graphically. We graph $\{(k, \hat{\alpha}_{k,n})\}$ in order to allow for a choice depending on k . One should choose $\hat{\alpha}_{k,n}$ from such a k -region where the plot is roughly horizontal.

Syntax

```
hillp varname [if exp] [in range]
```

Saved results

hillp saves the values of k and $\hat{\alpha}_{k,n}$ in the file `points.dta` as variables `values` and `y`.

Example

We applied the Hill plot to a set of 3000 simulated observations from a Pareto distribution with $\alpha = 1$ with the result given in Figure 1.

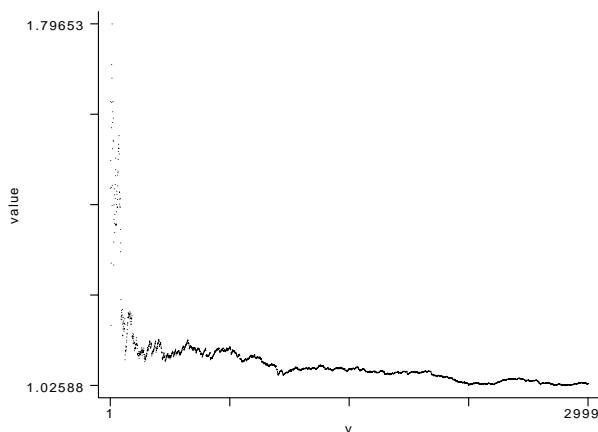


Figure 1. The Hill plot for a simulated sample of 3000 Pareto observations.

The Hill plot becomes stable after settling down and is in a tight neighborhood, and it seems to identify $\alpha = 1$ correctly.

References

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sg148

Profile likelihood confidence intervals for explanatory variables in logistic regression

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Abstract: Use of confidence intervals based on the Wald test and p -values for statistical significance based on the likelihood-ratio test can occasionally produce inconsistent results. One method used to avoid this problem is to estimate confidence intervals using the profile likelihood. This paper describes the command `logprof` written to estimate profile-likelihood based confidence intervals after fitting a logistic regression model. An example is given of how to use `logprof` using the automobile data available with Stata.

Keywords: profile likelihood, logistic regression.

Syntax

```
logprof varname [ , level(#) nograph or { inc(#) | incfrac(#) } from(#) to(#) saving(filename) ]
```

Description

Logistic regression analyses often produce (and report) p -values based on the likelihood-ratio test, while the standard confidence intervals produced in Stata are based on the Wald test. This can occasionally produce inconsistent results where the p -value suggests statistical significance, but the confidence interval does not. This problem occurred during the analysis of a recently published epidemiological study (Parker et al. 1999), and measures were taken to overcome it, leading to the creation of `logprof`.

One method of ensuring consistency between the confidence intervals and p -values is to estimate confidence intervals using the profile likelihood. `logprof` calculates such profile-likelihood based confidence intervals for independent variables after fitting a logistic regression model.

Options

`level(#)` specifies the confidence level, in percent, for confidence intervals. The default is `level(95)` or as set by `set level`.

`no`graph suppresses the graphing of the profile likelihood function. If not specified, the profile log-likelihood function is graphed and a line placed on the graph. The intercepts of this line with the function correspond to the confidence limits.

`or` requests confidence intervals to be displayed in terms of an odds ratio. If `or` is not specified, the beta coefficient and corresponding confidence interval are displayed.

`inc(#)` specifies the interval between each iterated value of the beta coefficient. The smaller the increment, the more accurate the results (and the longer the computer time taken). The default value of `inc` is 0.01. If the confidence interval contains a blank value for either the upper limit or the lower limit, the increment may have been set too large and the command must be rerun using a smaller increment.

`incfrac(#)` is an alternative to `inc`, where the increment is based on the value of the estimated beta coefficient. For example, using `incfrac(100)` would correspond to an increment of one-hundredth of the estimated beta coefficient.

`f`rom(#) specifies the value of the beta coefficient from which to start iterations.

`t`o(#) specifies the value of the beta coefficient at which to stop iterations.

Both `f`rom(#) and `t`o(#) have default values based on the maximum likelihood estimate of beta (and its confidence interval) from the previous logistic regression model. If the user-defined range does not produce a confidence interval, then the range is increased until it does.

`s`aving(filename) specifies that if a graph is produced, then it should be saved in the file having name *filename*.

Remarks

`logprof` was written to estimate 95% confidence intervals for an independent variable D , based on profile likelihood estimation, a method described in detail by Clayton and Hills (1993).

Briefly, the likelihood L for a logistic regression model can be written as

$$L = \prod_{i=1}^s \hat{p}_i \prod_{i=s+1}^{n-s} (1 - \hat{p}_i)$$

where \hat{p}_i is the predicted probability, given the final logistic regression model, of a stillbirth, s is the observed number of cases, and n is the total number of observations. The log likelihood for a particular value, β_D , of the coefficient for the effect of D

is calculated by fitting a logistic regression model to estimate the effects of the other covariates, but constraining the effect of D to be equal to β_D . Applying this to a range of values for β_D gives a range of values for L , a profile likelihood function. This function is then used to obtain a confidence interval based on the profile likelihood and corresponding to the value of β_D which gave a log likelihood equal to $L_{\max} - 0.5Z_\alpha$, where L_{\max} is the log likelihood from the original logistic regression model giving the maximum likelihood estimates of β , and Z_α is the critical value from a chi-squared distribution at the α level of significance.

`logprof` sets up a range for β , and obtains the profile likelihood function and the values needed for the confidence interval. This value of the log likelihood may not correspond exactly to a value of β in the range; hence, linear interpolation is used to obtain the confidence interval. The accuracy of the confidence interval depends on the difference between each increasing value of β_D , that is, that specified by the `inc` option.

Example

The use of `logprof` is now demonstrated using the `auto` dataset distributed with Stata. The dataset contains details on fuel consumption, weight, whether the car is foreign, and other characteristics of 74 cars in the year 1978.

The variable `foreign` takes on two unique values: 0 (domestic car) and 1 (foreign cars). If, as in the *Stata Reference Manual*, we wish to estimate the model

$$\Pr(\text{foreign} = 1) = F(\beta_0 + \beta_1 \times \text{weight} + \beta_2 \times \text{mpg})$$

where $F(z) = e^z / (1 + e^z)$ is the cumulative logistic distribution, we use

```
. use auto.dta, clear
(1978 Automobile Data)

. logit foreign weight mpg

Iteration 0:  log likelihood = -45.03321
Iteration 1:  log likelihood = -29.898968
Iteration 2:  log likelihood = -27.495771
Iteration 3:  log likelihood = -27.184006
Iteration 4:  log likelihood = -27.175166
Iteration 5:  log likelihood = -27.175156

Logit estimates                                Number of obs =          74
                                                LR chi2(2)         =        35.72
                                                Prob > chi2        =         0.0000
Log likelihood = -27.175156                    Pseudo R2         =         0.3966
```

foreign	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
weight	-.0039067	.0010116	-3.862	0.000	-.0058894	-.001924
mpg	-.1685869	.0919174	-1.834	0.067	-.3487418	.011568
_cons	13.70837	4.518707	3.034	0.002	4.851864	22.56487

This gives a 95% confidence interval based on the Wald test, which is suitable for many analyses. However, we may wish to estimate the confidence interval for `mpg` using a profile likelihood approach.

To do this, we can use `logprof`:

```
. logprof mpg, f(-1) t(1)
Maximum Profile Log-Likelihood = -27.175156
```

foreign	Coef.	[95% Conf. Interval]	
mpg	-.1685869	-.36799022	.00178716

As we haven't used the `nograph` option, we can also see what the profile log-likelihood looks like over the range of values for β considered, as illustrated in Figure 1.

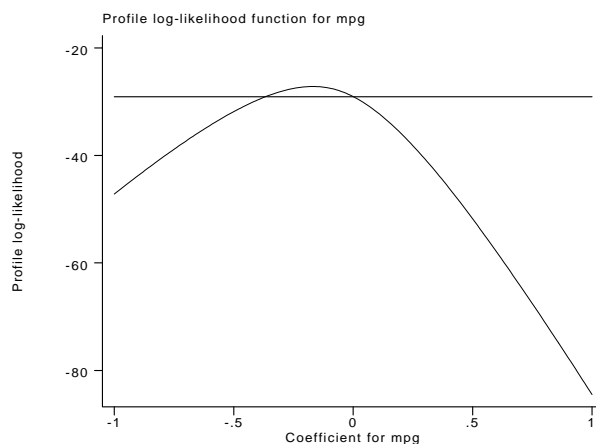


Figure 1. The profile log-likelihood for Example 1.

Saved results

The results and estimated coefficients and their standard errors from the most recently used estimation class model prior to the use of `logprof` are available in the usual way; see [U] **20.5 Accessing coefficients and standard errors**.

Acknowledgments

I thank Dr. Heather Dickinson and Professor Murray Aitkin, University of Newcastle and Ronna Cong, Stata Corporation for their useful comments and help in the formulation of `logprof`.

References

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sg149

Tests for seasonal data via the Edwards and Walter & Elwood tests

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Abstract: Many different methods have been proposed to assess the seasonality of binary outcomes, most often using monthly frequencies. This paper presents the command `seast` which will use either the Edwards test, for which population data is not required, or the test proposed by Walter and Elwood which takes into account the underlying population at risk. Examples are given of both tests using Canadian data on the monthly frequency of cases of anencephaly.

Keywords: seasonality, Edwards test, Walter and Elwoods test.

Syntax

```
seast outcomevar [popvar] [, gen(varname) sector(varname) edwards exact notab length(varname) ]
```

Description

`seast` performs a statistical significance test to detect seasonality of binary outcomes by fitting a simple harmonic curve to the observed data.

The data should be grouped into sectors, for example, number of events per month, with one record per sector.

`seast` uses either the Edwards test for seasonality (Edwards 1961), for which population data is not needed, or the test proposed by Walter and Elwood (1975), which was designed to take account of any seasonal variation in the underlying population at risk. Both methods are commonly used in epidemiology (Rothwell et al. 1996; Westerbeek et al. 1998) and can have uses in other fields. For a full discussion of both of these tests, see Walter (1994).

Options

The default for this command is that there are 12 sectors, the sector variable is called `month`, each month is the same length, and the Walter and Elwood test is to be used, which requires population data.

`gen(newvar)` generates a new variable containing the expected number of events for each month of birth. This may be useful for graphing the data, or for further calculations.

`sector(varname)` declares the name of the variable denoting the sector. This should be numbered sequentially from 1, . . . , n , where n is the total number of sectors. If $n = 12$, the program assumes that the sectors 1 through 12 correspond to January through December.

`edwards` specifies that the Edwards test is to be used. When using this option, the population variable does not need to be included as the test does not utilize this information. The default Edwards test makes no adjustment for variable sector length.

`exact` forces the test to adjust for variable month length. This is used when the sector has not been specified as the data consists of 12 months. This option sets each month length to be as used by the Gregorian calendar. `exact` does not need to be specified when the `length` option is used.

`notab` suppresses the displaying of a table of observed and expected number of events by sector.

`length(varname)` declares the name of the variable containing the length of each sector. For example, if the sectors were the first 6 months of the year, then the length variable should contain the number of days in each of the months. If `length` is specified, the test will take account of the length of the sectors varying.

Remarks

The methodology behind these tests has been discussed in detail elsewhere (Edwards 1961, Walter and Elwood 1975, Walter 1994). Both tests used in `seast` consider the data to be in the form of a circle divided into k sectors, corresponding to time intervals. In the case of months, the circle is divided into 12 sectors. In a situation where no seasonality exists, the expected center of the circle will be the actual center of the circle. Seasonality will cause a shift in the expected center to a distance from the actual center known as the amplitude. The direction of this shift will indicate the maximum and minimum points of the cyclical trend.

The Walter and Elwood test tests for seasonality of events while allowing for a seasonally variable population. The test statistic is based on the deviation of the observed number of cases from the expected number for each sector and compared with a chi-squared distribution on two degrees of freedom, to estimate a significance level.

The test also yields an estimate of the time at which the maximum occurs, the amplitude of seasonal variation, and a goodness-of-fit statistic which determines how well the differences between observed and expected cases in each sector can be fitted to a sine curve. Where fewer than 100 cases are observed, the test is known to perform poorly, as the asymptotic approximations on which it is based are not valid (St. Leger 1976, Roger 1977).

If the underlying population is unknown, then the Edwards test can be used, but with the assumption of a constant underlying population.

Example

Analyses of seasonal data are often carried out using monthly frequencies. One such analysis was used as an example by Walter and Elwood (1975) when proposing their test. The use of `seast` is illustrated with the data from their example. The listing below gives the monthly frequencies of cases of anencephalus and total births for Canada between 1954 and 1962 (Elwood 1975).

```
. use walter
(Canadian anencephaly data from Elwood (1975))
. list
      births   month   cases
1.    340797         1     468
2.    318319         2     399
3.    363626         3     471
4.    359689         4     437
5.    373878         5     376
6.    361290         6     410
7.    368867         7     399
8.    358531         8     472
9.    363551         9     418
10.   352173        10     448
11.   331964        11     409
12.   336894        12     397
```

The initial set of results are for the Walter and Elwood test, adjusting for variable month length and generating a variable `expected`, containing the expected number of events per month.


```

. seast cases births, exact gen(expected) notab
Walter & Elwood Test (using exact month lengths)
Total Number of Cases = 5104
Seasonality Test                                Goodness of fit test
chi2(2) = 12.4321                               chi2(2) = 23.7629
Prob > chi2 = 0.0020                             Prob > chi2 = 0.0138
-----
Parameter | Estimate
-----+-----
Amplitude of cyclic variation | .06968022
Angle of maximum rate        | -8.2767909
-----

```

The χ^2 statistic (12.43) indicates there is significant evidence of seasonality of birth for anencephalus cases ($P=0.002$), with the maximum rate estimated to be -8.3 degrees (late December) and amplitude of 0.07. However, we should interpret these results with caution because the model is a poor fit ($P=0.014$).

Removing the adjustment for variable month length makes little difference in this example:

```

. seast cases births, notab
Walter & Elwood Test
Total Number of Cases = 5104
Seasonality Test                                Goodness of fit test
chi2(2) = 12.4756                               chi2(2) = 23.7261
Prob > chi2 = 0.0020                             Prob > chi2 = 0.0139
-----
Parameter | Estimate
-----+-----
Amplitude of cyclic variation | .06993671
Angle of maximum rate        | -7.3950716
-----

```

Finally, we perform the Edwards test, which takes no account of the varying population at risk.

```

. seast cases births, notab edwards
Edwards Test
Total Number of Cases = 5104
Seasonality Test                                Goodness of fit test
chi2(2) = 0.7958                               chi2(2) = 27.2396
Prob > chi2 = 0.6717                             Prob > chi2 = 0.0042
-----
Parameter | Estimate
-----+-----
Amplitude of cyclic variation | .01765885
Angle of maximum rate        | -18.799564
-----

```

The use of the Edwards test on these data has removed the significance of the seasonality, and the seasonality is also less well described by a simple harmonic curve.

References

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STB categories and insert codes

Inserts in the STB are presently categorized as follows:

General Categories:

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<i>dt</i>	datasets	<i>tt</i>	teaching
<i>gr</i>	graphics	<i>zz</i>	not elsewhere classified
<i>in</i>	instruction		

Statistical Categories:

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<i>sed</i>	exploratory data analysis	<i>ssi</i>	simulation & random numbers
<i>sg</i>	general statistics	<i>sss</i>	social science & psychometrics
<i>smv</i>	multivariate analysis	<i>sts</i>	time-series, econometrics
<i>snp</i>	nonparametric methods	<i>svy</i>	survey sampling
<i>sqc</i>	quality control	<i>sxd</i>	experimental design
<i>sqv</i>	analysis of qualitative variables	<i>szz</i>	not elsewhere classified
<i>srd</i>	robust methods & statistical diagnostics		

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2. Any ado-files, .exe files, or other software that accompanies the submission.
3. A help file for each ado-file included in the submission. See any recent STB diskette for the structure a help file. If you have questions, fill in as much of the information as possible and we will take care of the details.
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5. Files containing the graphs to be included in the insert. If you have used STAGE to edit the graphs in your submission, be sure to include the .gph files. Do not add titles (e.g., "Figure 1: ...") to your graphs as we will have to strip them off.

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```
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