The *Stata Journal* publishes reviewed papers together with shorter notes or comments, regular columns, book reviews, and other material of interest to Stata users. Examples of the types of papers include 1) expository papers that link the use of Stata commands or programs to associated principles, such as those that will serve as tutorials for users first encountering a new field of statistics or a major new technique; 2) papers that go “beyond the Stata manual” in explaining key features or uses of Stata that are of interest to intermediate or advanced users of Stata; 3) papers that discuss new commands or Stata programs of interest either to a wide spectrum of users (e.g., in data management or graphics) or to some large segment of Stata users (e.g., in survey statistics, survival analysis, panel analysis, or limited dependent variable modeling); 4) papers analyzing the statistical properties of new or existing estimators and tests in Stata; 5) papers that could be of interest or usefulness to researchers, especially in fields that are of practical importance but are not often included in texts or other journals, such as the use of Stata in managing datasets, especially large datasets, with advice from hard-won experience; and 6) papers of interest to those who teach, including Stata with topics such as extended examples of techniques and interpretation of results, simulations of statistical concepts, and overviews of subject areas.

For more information on the *Stata Journal*, including information for authors, see the webpage

http://www.stata-journal.com

The *Stata Journal* is indexed and abstracted in the following:

- CompuMath Citation Index®
- Current Contents/Social and Behavioral Sciences®
- RePEc: Research Papers in Economics
- Science Citation Index Expanded (also known as SciSearch®)
- Scopus™
- Social Sciences Citation Index®

**Copyright Statement:** The *Stata Journal* and the contents of the supporting files (programs, datasets, and help files) are copyright © by StataCorp LP. The contents of the supporting files (programs, datasets, and help files) may be copied or reproduced by any means whatsoever, in whole or in part, as long as any copy or reproduction includes attribution to both (1) the author and (2) the *Stata Journal*.

The articles appearing in the *Stata Journal* may be copied or reproduced as printed copies, in whole or in part, as long as any copy or reproduction includes attribution to both (1) the author and (2) the *Stata Journal*.

Written permission must be obtained from StataCorp if you wish to make electronic copies of the insertions. This precludes placing electronic copies of the *Stata Journal*, in whole or in part, on publicly accessible websites, file servers, or other locations where the copy may be accessed by anyone other than the subscriber.

Users of any of the software, ideas, data, or other materials published in the *Stata Journal* or the supporting files understand that such use is made without warranty of any kind, by either the *Stata Journal*, the author, or StataCorp. In particular, there is no warranty of fitness of purpose or merchantability, nor for special, incidental, or consequential damages such as loss of profits. The purpose of the *Stata Journal* is to promote free communication among Stata users.

The *Stata Journal* electronic version (ISSN 1536-8734) is a publication of Stata Press. Stata, *Stata*, Mata, and NetCourse are registered trademarks of StataCorp LP.
From resultssets to resultstables in Stata

Roger B. Newson
National Heart and Lung Institute
Imperial College London
London, UK
r.newson@imperial.ac.uk

Abstract. The listtab package supersedes the listtex package. It inputs a list of variables and outputs them as a table in one of several formats, including \TeX, \LaTeX, HTML, Microsoft Rich Text Format, or possibly future XML-based formats. It works with a team of four other packages: sdecode, chardef, xrewide, and ingap, which are downloaded from the Statistical Software Components archive. The sdecode package is an extension of decode; it allows the user to add prefixes or suffixes and to output exponents as \TeX, \HTML, or Rich Text Format superscripts. It is called by another package, bmjcip, to convert estimates, confidence limits, and \textit{p}-values from numeric to string. The chardef package is an extension of \texttt{char define}; it allows the user to define a characteristic for a list of variables, adding prefixes or suffixes. The xrewide package is an extension of \texttt{reshape wide}; it allows the user to store additional results in variable characteristics or local macros. The ingap package inserts gap observations into a dataset to form gap rows when the dataset is converted to a table. Together, these packages form a general toolkit to convert Stata datasets (or resultssets) to tables, complete with row labels and column labels. Examples are demonstrated using \LaTeX and also using the rtfutil package to create Rich Text Format documents. This article uses data from the Avon Longitudinal Study of Parents and Children, based at Bristol University, UK.

Keywords: st0254, listtab, listtex, sdecode, chardef, xrewide, ingap, bmjcip, rtfutil, keyby, addinby, table, resultsset, \TeX, \LaTeX, HTML, XML, RTF

1 Introduction

Statisticians and other scientists routinely produce tables of descriptive and inferential statistics. Unsurprisingly, Stata users have produced several add-on packages to input the various Stata data and output structures and to output tables of statistics in a variety of formats, such as \TeX, \LaTeX, Hypertext Markup Language (HTML), Extensible Markup Language (XML), Microsoft Rich Text Format (RTF), or delimited text tables for menu-driven manual incorporation into word processor documents.

The packages for this purpose that are most frequently downloaded from the Statistical Software Components (SSC) archive are Roy Wada’s outreg2, Ben Jann’s estout, John Luke Gallup’s outreg, and Ian Watson’s tabout. Some very inspiring presentations were given at the 2009 UK Stata Users Group meeting by Ben Jann (2009), who gave a tutorial on estout, and by Adam Jacobs (2009), who described some of his own programs to create XML-based documents for Open Document Format packages.
From resultssets to resultstables in Stata

My own method for table production was based initially on Stata’s listtex package, which has since been superseded by listtab for users of Stata 10 or higher. The latest versions of both packages are still downloadable from SSC. Both of these packages input a list of variables from the Stata dataset in memory and output a text table (in the Stata log or in a file) with one row per observation. The table is derived by outputting the values of these variables, preceded by a begin() string option, separated by a delimiter() string option, and suffixed by an end() string option. Optionally, the user may specify a list of string header rows to be output before the observation rows by using the headlines() option or a list of string footer rows to be output after the observation rows by using the footlines() option.

This general format includes, as special cases, most table formats known to computer science and probably a few yet to be invented. To save the user the trouble of specifying a long list of options, there is an rstyle() option to specify one of a collection of row style names, such as rstyle(tabular) for \texttt{L\LaTeX} tabular environments (Lamport 1994). A row style is defined as a vector of four string values—namely, a begin() string, a delimiter() string, an end() string, and a missnum() string—to be output in the place of numeric missing values. All of these options default to the empty string for listtab. However, the delimiter() option defaults to the ampersand (\&) for listtex, because I was thinking of \TeX tables as the norm when I wrote listtex. The missnum() option is probably not often used with either package, because numeric variables should usually be converted to string variables before being output to tables.

The listtab or listtex method is simple and versatile, if only the user already has a dataset in memory with one observation per table row. Fortunately, StataCorp and other Stata users have already developed a variety of programs, such as official Stata’s \texttt{statsby} and the SSC package \texttt{parmest}, that could be used to produce such datasets (or resultssets). Some of these programs, and their use with listtex, are described in Newson (2003). Since that article was written, resultsset processing has developed further and additional Stata packages have become available.

In this article, I aim to give an update of techniques for converting resultssets (and other datasets) to tables (or resultstables). The methods used can be summarized as “decode, label, and list” (in the simplest cases) or as “decode, label, reshape, insert gaps, and list” (in the most complicated cases).

1.1 Example resultsset: Genotype frequencies in ALSPAC

The resultsset that we will use to demonstrate these techniques was produced using data from the Avon Longitudinal Study of Parents and Children (ALSPAC). For further information about this birth cohort study, refer to http://www.alspac.bris.ac.uk (the study website). Analyses of some of these data are discussed in Shaheen et al. (2010) and Henderson et al. (2010).

In the ALSPAC birth cohort study, 14,060 children born in and around Bristol, UK, in 1991 and 1992 have been observed through progression to adulthood. Subsets of these children, and of their mothers, gave blood or other tissue samples, which were
used to measure their genotypes with respect to 24 biallelic polymorphisms, using assay methods developed by molecular geneticists. An allele is a version of a gene, and a biallelic polymorphism is a set of two possible alleles for a gene. The genotype of a study subject with respect to a polymorphism is the total set of alleles present in the two copies of the human genome present in each cell of that study subject; one copy comes from the subject’s mother’s egg, while the other copy comes from the subject’s father’s sperm. Genotypes with respect to a biallelic polymorphism may have values of the form $GG$, $Gg$, or $gg$, where $G$ is the first allele and $g$ is the second allele.

Before measuring associations of the two alleles of a biallelic polymorphism with a disease, geneticists usually measure the frequencies of the three possible genotypes in the sample of subjects. An important population parameter of this frequency distribution is the geometric mean homozygote or heterozygote ratio, defined as $\sqrt{P_{GG}P_{gg}/P_{Gg}}$, where $P_{GG}$, $P_{Gg}$, and $P_{gg}$ are the proportions of subjects in the population with genotypes $GG$, $Gg$, and $gg$, respectively. The sample estimate of this ratio is $\sqrt{N_{GG}N_{gg}/N_{Gg}}$, and the standard error of its log is $\sqrt{1/(4N_{GG})+1/(4N_{gg})+1/N_{Gg}}$, where $N_{GG}$, $N_{Gg}$, and $N_{gg}$ are the numbers of subjects in the sample with genotypes $GG$, $Gg$, and $gg$, respectively. If the geometric mean homozygote or heterozygote ratio is 0.5, as would be the case under random mating (where the maternal egg allele and the paternal sperm allele are statistically independent), then the frequency distribution of the genotypes is said to be in Hardy–Weinberg equilibrium. The alternative possibilities are a tendency to inbreeding (where the ratio is greater than 0.5, indicating a tendency for mothers and fathers to contribute the same allele) and a tendency to outbreeding (where the ratio is less than 0.5, indicating a tendency for mothers and fathers to contribute different alleles). Justifications of these formulas can be found in Newson (2008a).

The example resultsset to be used in this article contains one observation per polymorphism per DNA source (“Child” or “Maternal”) for each of 24 polymorphisms assayed. The resultsset also contains data on the number of subjects (children or mothers) whose genotypes were assessed for the polymorphism, and the numbers and percentages with each genotype. This is together with estimates and confidence intervals for the geometric mean homozygote or heterozygote ratios for 24 biallelic polymorphisms, and $p$-values for testing the hypothesis of Hardy–Weinberg equilibrium.

The observations in the resultsset are sorted (and uniquely identified) by a key of three value-labeled numeric variables: source (indicating DNA source), polygp (indicating a polymorphism group corresponding to a particular protein specified by a particular gene), and poly (indicating the polymorphism). The resultsset was created using the SSC packages xcontract, dsconcat, and factmerg, together with the parmcip command of the SSC package parmest. The methods used to create this resultsset are discussed in Newson (2008b), which also gives a plot of some of the confidence intervals. Methods for production of resultssets are discussed more generally in Newson (2003) and also in many more recent references accessible by hypertext from the online help for the parmest package.

In this article, I will demonstrate how to convert these results into tables in L\TeX (which Stata programmers frequently like to use) and in RTF (which produces Microsoft-
From resultssets to resultstables in Stata

ready documents, which their colleagues frequently demand). In section 2, I explain how a table can be viewed as a special case of a Stata dataset. In sections 3–5, I demonstrate increasingly advanced examples of table production in \LaTeX. In section 6, I describe the creation of tables in RTF documents. Finally, in section 7, I outline possible further extensions of the methods presented in this article. The example resultsset, together with the Stata do-files used to output the tables, can be downloaded (using Stata) as online supplementary material to this article.

2 Tables as Stata datasets

To create a table using \texttt{listtab}, we must first have a suitable Stata dataset in the memory, with one observation per table row.

In general, a well-formed Stata dataset should be designed to have one observation per \textit{thing} and data on \textit{attributes of things}. For instance, the example resultsset has one observation per combination of DNA source (“Child” or “Maternal”) and polymorphism, data on genotype frequencies for that polymorphism in that DNA source, and data on statistics derived from those frequencies. Specifically, a well-formed Stata dataset should be a table, as defined by the relational model of Date (1986). This model (in its essentials) assumes that a table (or dataset) is a mathematical function whose domain is the set of existing combinations of key variable values (specifying the \textit{things}) and whose range is the set of all possible combinations of non-key variable values (specifying the \textit{attributes of things}). For a Stata dataset to conform to this model, it should be keyed (that is to say, its observations should be sorted and uniquely identified) by the values of a list of key variables. In the case of the example resultsset, these key variables are \texttt{source}, \texttt{polygp}, and \texttt{poly}.

Not all Stata datasets conform to the relational model. Some are not sorted or are sorted by variables that do not uniquely identify the observations. Similarly, not all keyed Stata datasets are ideal for input to \texttt{listtab}. I would argue (from experience) that a Stata dataset input to \texttt{listtab} should be keyed, with key variables identifying the table rows in order of appearance on the page, possibly preceded in the key by key variables identifying the pages of a multipage table. The dataset should also have the following features:

1. The variable list input to \texttt{listtab} should contain only string variables.

2. The first variable in the list should be a row label variable whose value, in each observation, is determined from the values of the key variables in that observation.

3. The variables input to \texttt{listtab} should have one or more characteristics (see \texttt{[P] char}) containing the corresponding column labels for the output table.

The first of these features is a good idea because numeric variables nearly always have to be reformatted before inclusion in tables in ways that are not provided by any official Stata format. The other two features are a good idea because tables in
documents have row labels and column labels, both of which can contain justification, font specification, or other formatting information.

2.1 Making resultstables by breaking resultssets

A typical resultsset produced by parmust, contract, collapse, or statsby contains mostly numeric variables containing frequencies or sample statistics. Formatting resultssets for input to listtab is usually a destructive process, because these numeric variables are converted irreversibly to string variables. The process is even more destructive when the dataset is also subset, reshaped, or provided with gap rows to make the table more readable. It is therefore usually a good idea to program this conversion process as a sequence of commands in a do-file and to enclose these commands between a preserve statement and a restore statement (see [P] preserve).

Stata programmers are frequently urged to avoid the use of preserve and restore, because they involve file processing and therefore consume computing resources. Unfortunately, in Stata 1 to 12, the user may only have access to one dataset in memory at a time and therefore frequently has no choice but to use preserve and restore. Fortunately, if a resultsset is small enough to be converted to a table or even to a multipage document of multipage tables, then this resource intensity is not likely to be an important issue.

I usually follow a policy of writing a do-file to produce a keyed resultsset on disk, and then writing a second do-file to input this resultsset and to produce the tables (and plots), which frequently have to be revised iteratively before they have exactly the right look and feel.

2.2 Enforcing the relational model with keyby and addinby

Two useful packages to enforce and use the relational model in Stata are keyby and addinby, both downloadable from SSC. The command keyby is an extension of sort (see [D] sort); it sorts the dataset by a list of variables and then checks that the dataset is keyed by these variables. The keyby package also contains the command keybygen, which sorts the data by a list of variables and adds an additional variable containing the preexisting order of an observation within its by-group to complete the key. Both keyby and keybygen reorder the key variables to be the first in the dataset unless the user specifies a noorder option.

The addinby package is an extension of merge (see [D] merge); it adds extra variables to the dataset in memory from a second dataset by using a list of existing variables as a foreign key, which is assumed to be the primary key of the second dataset. Both keyby and addinby fail if the list of variables specified does not key the dataset that it is supposed to key. They also fail if these key variables contain missing values (unless the user specifies a missing option). Both commands restore the original dataset in the event of failure (unless the user specifies a fast option).
The `keyby` and `addinby` packages are frequently used to produce a keyed resultssset to be stored on disk before that resultssset is subset, decoded, reshaped, or supplied with gap rows for input to `listtab`.

### 3 Simple tables: Decode, label, list

Our first `listtab` examples are the creation of tables 1 and 2. Table 1, a minimal example, is a modified version of table 1 of Newson (2003), with one row for each Volkswagen car model in `auto.dta`. Table 2, a more typical example, has one row per polymorphism in the example resultssset, labeled by the polymorphism group and the name of the individual polymorphism; data on the total number of child subjects; data on the numbers of subjects with genotypes $GG$, $Gg$, and $gg$ (where $G$ is the first allele and $g$ is the second allele); and the estimates, confidence limits, and $p$-values for the geometric mean homozygote or heterozygote ratios.

#### Table 1. Volkswagen cars in `auto.dta`

<table>
<thead>
<tr>
<th>Make and Model</th>
<th>Mileage (mpg)</th>
<th>Weight (lbs.)</th>
<th>Price</th>
</tr>
</thead>
<tbody>
<tr>
<td>VW Dasher</td>
<td>23</td>
<td>2,160</td>
<td>$7,140</td>
</tr>
<tr>
<td>VW Diesel</td>
<td>41</td>
<td>2,040</td>
<td>$5,397</td>
</tr>
<tr>
<td>VW Rabbit</td>
<td>25</td>
<td>1,930</td>
<td>$4,697</td>
</tr>
<tr>
<td>VW Scirocco</td>
<td>25</td>
<td>1,990</td>
<td>$6,850</td>
</tr>
</tbody>
</table>
Table 2. Child genotype frequencies and homozygote or heterozygote ratios

<table>
<thead>
<tr>
<th>Polymorphism (by group)</th>
<th>N</th>
<th>GG</th>
<th>Gg</th>
<th>gg</th>
<th>Ratio (95% CI)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>AHR, rs2066853</td>
<td>8704</td>
<td>6811</td>
<td>1788</td>
<td>105</td>
<td>0.47 (0.43, 0.53)</td>
<td>.3</td>
</tr>
<tr>
<td>CYP1A1, rs1048943</td>
<td>8764</td>
<td>8206</td>
<td>544</td>
<td>15</td>
<td>0.64 (0.49, 0.84)</td>
<td>.062</td>
</tr>
<tr>
<td>CYP1A1, rs4646903</td>
<td>8587</td>
<td>6899</td>
<td>1567</td>
<td>121</td>
<td>0.58 (0.53, 0.65)</td>
<td>.0033</td>
</tr>
<tr>
<td>CYP2A6, rs1801272</td>
<td>8666</td>
<td>8203</td>
<td>454</td>
<td>9</td>
<td>0.60 (0.43, 0.84)</td>
<td>.3</td>
</tr>
<tr>
<td>CYP2A6, rs28399433</td>
<td>8691</td>
<td>7716</td>
<td>936</td>
<td>39</td>
<td>0.59 (0.49, 0.69)</td>
<td>.067</td>
</tr>
<tr>
<td>GCL, rs17883901</td>
<td>9389</td>
<td>7925</td>
<td>1401</td>
<td>63</td>
<td>0.50 (0.44, 0.58)</td>
<td>.9</td>
</tr>
<tr>
<td>GPX, rs713041</td>
<td>8611</td>
<td>2733</td>
<td>4168</td>
<td>1710</td>
<td>0.52 (0.50, 0.54)</td>
<td>.093</td>
</tr>
<tr>
<td>GSTM1, GSTM1 CNV</td>
<td>8399</td>
<td>585</td>
<td>3163</td>
<td>4651</td>
<td>0.52 (0.49, 0.55)</td>
<td>.14</td>
</tr>
<tr>
<td>GSTP1, rs947894</td>
<td>8692</td>
<td>3695</td>
<td>3937</td>
<td>1060</td>
<td>0.50 (0.48, 0.53)</td>
<td>.82</td>
</tr>
<tr>
<td>GSTT1, GSTT1 CNV</td>
<td>7591</td>
<td>2448</td>
<td>3653</td>
<td>1490</td>
<td>0.52 (0.50, 0.55)</td>
<td>.056</td>
</tr>
<tr>
<td>IGF, rs35767</td>
<td>9383</td>
<td>6579</td>
<td>2555</td>
<td>249</td>
<td>0.50 (0.47, 0.54)</td>
<td>.96</td>
</tr>
<tr>
<td>IGF, rs2854744</td>
<td>8489</td>
<td>2456</td>
<td>4202</td>
<td>1831</td>
<td>0.50 (0.48, 0.53)</td>
<td>.67</td>
</tr>
<tr>
<td>LTA, rs909253</td>
<td>8664</td>
<td>3397</td>
<td>4089</td>
<td>1178</td>
<td>0.49 (0.47, 0.51)</td>
<td>.34</td>
</tr>
<tr>
<td>MIF, rs755622</td>
<td>9500</td>
<td>6454</td>
<td>2757</td>
<td>289</td>
<td>0.50 (0.46, 0.53)</td>
<td>.79</td>
</tr>
<tr>
<td>MTNR1B, rs10830963</td>
<td>8584</td>
<td>4570</td>
<td>3357</td>
<td>657</td>
<td>0.52 (0.49, 0.54)</td>
<td>.24</td>
</tr>
<tr>
<td>Nr2f, rs1806649</td>
<td>8606</td>
<td>4889</td>
<td>3199</td>
<td>518</td>
<td>0.50 (0.47, 0.53)</td>
<td>.86</td>
</tr>
<tr>
<td>Nr2f, rs1962142</td>
<td>8763</td>
<td>7154</td>
<td>1528</td>
<td>81</td>
<td>0.50 (0.44, 0.56)</td>
<td>.95</td>
</tr>
<tr>
<td>Nr2f, rs2364723</td>
<td>8725</td>
<td>4047</td>
<td>3760</td>
<td>918</td>
<td>0.51 (0.49, 0.54)</td>
<td>.31</td>
</tr>
<tr>
<td>Nr2f, rs6706649</td>
<td>8731</td>
<td>6696</td>
<td>1920</td>
<td>115</td>
<td>0.46 (0.41, 0.51)</td>
<td>.086</td>
</tr>
<tr>
<td>Nr2f, rs6721961</td>
<td>9419</td>
<td>7534</td>
<td>1775</td>
<td>110</td>
<td>0.51 (0.46, 0.57)</td>
<td>.64</td>
</tr>
<tr>
<td>Nr2f, rs6726395</td>
<td>8692</td>
<td>2580</td>
<td>4323</td>
<td>1789</td>
<td>0.50 (0.48, 0.52)</td>
<td>.78</td>
</tr>
<tr>
<td>Nr2f, rs10183914</td>
<td>8671</td>
<td>3617</td>
<td>3975</td>
<td>1079</td>
<td>0.50 (0.47, 0.52)</td>
<td>.8</td>
</tr>
<tr>
<td>TNF, rs1800629</td>
<td>8685</td>
<td>5679</td>
<td>2699</td>
<td>307</td>
<td>0.49 (0.46, 0.52)</td>
<td>.53</td>
</tr>
<tr>
<td>UGB, rs3741240</td>
<td>8671</td>
<td>3602</td>
<td>4031</td>
<td>1038</td>
<td>0.48 (0.46, 0.50)</td>
<td>.079</td>
</tr>
</tbody>
</table>

In both of these cases, we start with a dataset (auto.dta for table 1 and the example resultsset for table 2) and simply decode, label, and list the variables we want to tabulate. The tools used to do this are sdecode (to decode the numbers and label the rows), chardef (to label the columns), and listtab (including the listtab.vars command) to output the table as a \LaTeX\ tabular environment to the Stata log or the Results window or an output file, from which it was cut and pasted into the \LaTeX\ version of this article.

### 3.1 Numeric to string conversion using sdecode

The decoding and row labeling is done by the sdecode package, which is a “super” version of decode (see [D] decode). It has the companion package sencode, a “super” version of encode (see [D] encode). Both packages are downloadable from SSC. The sencode package converts a string variable to a value-labeled numeric variable and is used extensively to produce graphs, because a string variable must usually be converted to numeric before it can be used as an axis variable on a graph. By contrast, the sdecode package converts a numeric variable (value-labeled or unlabeled) to a string
variable and is used extensively in producing tables. Numeric variables in tables are usually formatted to a specified number of decimal places or significant figures and often are formatted to scientific notation with superscripts. They may have added parentheses (if they are confidence limits) or stars (if they are $p$-values), and they may also have additional prefixes or suffixes indicating justification (left, right, or center) or fonts (bold, italic, or nonstandard size).

To cater for all these possibilities, `sdecode` has some extra options that are not available for `decode`:

1. The user can include a `generate()` option to specify that a new string output variable will be generated and can include a `replace` option to specify that the string output variable will replace the numeric input variable, inheriting its position in the dataset and its characteristics (see [P] `char`).

2. Unlabeled numeric values can be converted to string by using formats, possibly specified with a `format()` option, as with the `tostring` command (see [D] `destring`).

3. The `esub()` option allows the user to convert format-generated string values containing embedded $e$- or $e+$ substrings, indicating scientific notation, to a number of exponentiation formats. These include `esub(texsuper)` (converting exponents to $\TeX$ superscripts), `esub(htmlsuper)` (converting exponents to HTML superscripts), and `esub(rtfsuper)` (converting exponents to RTF superscripts).

4. The `ftrim` option allows the user to trim format-generated string values, removing spaces on the left and on the right.

5. The `xmlsub` option replaces output substrings &lt; and &gt; with the XML escape sequences &amp;, &lt;, and &gt; respectively.

6. The `prefix()` and `suffix()` options allow the user to add a prefix and a suffix, respectively, to the output string values.

These extra options allow numeric variables to be converted to string equivalents suited to a wide variety of document formats. However, to make life easier for users, `sdecode` has been extended further by the provision of the `bmjcip` package, also downloadable from SSC, which is a front end for `sdecode`. `bmjcip` inputs up to four numeric variables, which it interprets as a $p$-value, an estimate with a $p$-value, an estimate with two confidence limits, or an estimate with two confidence limits and a $p$-value, depending on whether the user supplies one, two, three, or four input variables, respectively. It decodes these variables, replacing them with their decoded string versions, prefixed and suffixed appropriately.

### 3.2 Characteristic assignment using `chardef`

As stated previously, users are advised to store to `listtab` column labels for tables in characteristics assigned to the corresponding variable’s output (see [P] `char`). To
reduce the number and complexity of char define statements required, the chardef package, downloadable from SSC, has been provided to semiautomate this assignment. chardef assigns a user-specified characteristic for a list of variables, assigning a list of corresponding values that may be prefixed or suffixed using the prefix() option or the suffix() option. The characteristics may be cleared using the charundef command of the chardef package.

The aim of this mass production (and mass destruction) of variable characteristics is to enable the user to create header lines in local macros to be input to the headlines() option of listtab. The listtab package includes the command listtab vars, which inputs a list of variables (with a row style) that will later be output using listtab and outputs a header line to a local macro. This header line contains a list of the variable names, the variable labels, the values of a named variable characteristic, or another attribute of the variables, separated by the delimiter() string of the row style, and prefixed and suffixed by the begin() and end() strings of the row style, respectively. This local macro can then be input to listtab as part of the headlines() string list option, to provide one or more rows of column labels in the output table.

3.3 The code to create table 1

Having introduced the packages to be used, I can now present the Stata code that combines them to produce table 1. If we assume that auto.dta is in the memory, then this code (and its output to the Stata log) is as follows:

```
. preserve
  . keep if word(make,1) == "VW"  
  (70 observations deleted)
  . sdecode mpg, replace
  . sdecode weight, replace
  . sdecode price, replace prefix("\$")
  . chardef make mpg weight price, char(varname) prefix("\textsl{") suffix("}\")
     values("Make and Model" "Mileage (mpg)" "Weight (lbs.)" "Price")
  . listtab_vars make mpg weight price, substitute(char varname) rstyle(tabular)
     > local(h1)
  . listtab make mpg weight price, type rstyle(tabular)
     > headlines("\begin{tabular}{rrrr} " 
     > \hline
     > \textsl{Make and Model} & \textsl{Mileage (mpg)} & \textsl{Weight (lbs.)} & \textsl{Price} \\
     > \hline
     > VW Dasher & 23 & 2,160 & \$7,140 \\
     > VW Diesel & 41 & 2,040 & \$5,397 \\
     > VW Rabbit & 25 & 1,930 & \$4,697 \\
     > VW Scirocco & 25 & 1,990 & \$6,850 \\
     > \hline
     > \end{tabular}"
  . restore
```
We start by using `preserve` to save a copy of the original `auto.dta`, which in this case is not a resultset. We then discard the data on all non-Volkswagen car models by using `keep`, and we use `sdecode` to convert the numeric variables `mpg`, `weight`, and `price` to string variables, prefixed in the case of `price` by a \texttt{LaTeX} dollar sign. (Note that we do not need to create a row label variable because the existing variable `make` will play that role and the role of a key for the dataset.) We then specify column labels by using `chardef`, which assigns the characteristic `varname` of the variables `make`, `mpg`, `weight`, and `price` with a list of values specified by the `values()` option, prefixed by \texttt{\textsl{and suffix by}} {}, so the column labels will be slanted in a \texttt{LaTeX} environment.

Then we use the `listtab_vars` command to combine the `varname` characteristics of the variables `make`, `mpg`, `weight`, and `price` into an output local macro `h1` containing a header row for the table. We do this by using the option `rstyle(tabular)` to specify that the header row be in the \texttt{LaTeX} \texttt{tabular} row style and by using the option `substitute(char varname)` to specify that the cells in the header row be taken from the variable characteristic `varname` instead of from the variable names.

After this, we use `listtab` with a `headlines()` option to specify table headlines including the local macro `h1` and a `footlines()` options to specify table footlines. The `listtab` command outputs the variables `make`, `mpg`, `weight`, and `price` to the Stata log in the form of some alien-looking output, which readers may recognize as a \texttt{LaTeX} \texttt{tabular} environment, ready to be cut and pasted into a \texttt{LaTeX} document to produce table 1. Finally, the `restore` command restores the original `auto.dta`.

### 3.4 The code to create table 2

As a less trivial example, I can now present the Stata code to produce table 2. If we assume that the example resultset is in the memory, then this code (and its output to the Stata log) is as follows:

```
. preserve
. keep if source==1
(24 observations deleted)
. keep source polygp poly N _freq* homhet min* max* p
. format homhet min* max* %8.2f
. bmjcip homhet min* max* p, esub(texsuper) prefix($) suffix($)
. foreach Y of var N _freq* {
   2. sdecode `Y´, replace esub(texsuper) prefix($) suffix($)
   3. }
. sdecode polygp, generate(S_1)
. sdecode poly, generate(S_2)
. generate row=S_1", "S_2
. chardef row N _freq* homhet min* max* p,
   > char(varname) prefix("\textsl{") suffix("})
   > values("Polymorphism (by group)" N GG Gg gg Ratio "(95\% CI)" P)
. listtab_vars row N _freq* homhet min* max* p, rstyle(tabular)
   > substitute(char varname) local(h1)
```
We start by using *preserve* to save a copy of the original resultsset (with data on genotype frequencies in children and in mothers). We keep only the subset *source==1*, because the variable *source* is 1 for child results and 2 for maternal results. We then reset the formats of the homozygote or heterozygote ratios and their confidence limits to *%8.2f* (implying two decimal places), and we use *bmjcip* to decode the ratios, their confidence limits, and their *p*-values from numeric variables to string variables.

We then use a *foreach* loop to decode the variables *N*, *freq0*, *freq1*, and *freq2* to string variables. For all of these decoded variables, we use the options *prefix($) and suffix($)* to output the numbers in *\LaTeX* math mode; this was not really necessary but might have been necessary if there had been any exponentiated numbers (such as very small *p*-values or astronomical sample numbers). We label the table rows, using *sdecode* to decode the key variables *polygp* and *poly* to new variables *S*1 and *S*2, respectively, and combining their values to produce a new row label variable *row*. We label the table columns—which are the variable *row* and the decoded statistical variables—using *chardef*, which assigns the characteristic *varname* to contain slanted table header cells, in the way earlier demonstrated in the creation of table 1.

We can now use *listtab_vars* to input the table column variables and output a header row to the local macro *h1*, again in the way demonstrated in the creation of table 1. Having created this header row, we can use *listtab*, which again uses the local macro *h1* as part of its *headlines()* option. The *listtab* command creates a *\LaTeX* *tabular* environment, even more alien-looking than the one for table 1, and (fortunately) does not type it to the Stata log but writes it to the file *texdemo1_2.tex*. This file may be cut and pasted into a *\LaTeX* document or included using \input to produce table 2. Finally, the *restore* command restores the original resultsset, with the original numeric statistical variables and observations for child and maternal results, ready to be used to produce further resultstables.

4 Parallel tables: Decode, label, reshape, list

In our next example, we demonstrate the production of table 3. Unlike table 2, table 3 does not include the individual genotype frequencies or the *p*-values. However, it does include sample numbers, homozygote or heterozygote ratios, and confidence limits both for child genotypes and for maternal genotypes. The child and maternal results are displayed side by side in the same table rows. This can be done by using the official Stata *reshape* command (see [D] *reshape*) or the SSC package *xrewide*. 
From resultssets to resultstables in Stata

Table 3. Child and maternal homozygote or heterozygote ratios

<table>
<thead>
<tr>
<th>Polymorphism</th>
<th>Child:</th>
<th>Maternal:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Ratio (95% CI)</td>
</tr>
<tr>
<td>AHR, rs2066853</td>
<td>8704</td>
<td>0.47 (0.43, 0.53)</td>
</tr>
<tr>
<td>CYP1A1, rs1048943</td>
<td>8704</td>
<td>0.64 (0.49, 0.84)</td>
</tr>
<tr>
<td>CYP1A1, rs4646903</td>
<td>8587</td>
<td>0.58 (0.53, 0.65)</td>
</tr>
<tr>
<td>CYP2A6, rs1801272</td>
<td>8666</td>
<td>0.60 (0.43, 0.84)</td>
</tr>
<tr>
<td>CYP2A6, rs28399433</td>
<td>8691</td>
<td>0.59 (0.49, 0.69)</td>
</tr>
<tr>
<td>GCL, rs17883901</td>
<td>9389</td>
<td>0.50 (0.44, 0.58)</td>
</tr>
<tr>
<td>GPX, rs713041</td>
<td>8611</td>
<td>0.52 (0.50, 0.54)</td>
</tr>
<tr>
<td>GSTM1, GSTM1 CNV</td>
<td>8399</td>
<td>0.52 (0.49, 0.55)</td>
</tr>
<tr>
<td>GSTP1, rs947894</td>
<td>8692</td>
<td>0.50 (0.48, 0.53)</td>
</tr>
<tr>
<td>GSTT1, GSTT1 CNV</td>
<td>7591</td>
<td>0.52 (0.50, 0.55)</td>
</tr>
<tr>
<td>IGF, rs25767</td>
<td>9383</td>
<td>0.50 (0.47, 0.54)</td>
</tr>
<tr>
<td>IGF, rs2854744</td>
<td>8489</td>
<td>0.50 (0.48, 0.53)</td>
</tr>
<tr>
<td>LTA, rs909253</td>
<td>8664</td>
<td>0.49 (0.47, 0.51)</td>
</tr>
<tr>
<td>MIF, rs755622</td>
<td>9500</td>
<td>0.50 (0.46, 0.53)</td>
</tr>
<tr>
<td>MNTR1B, rs1083963</td>
<td>8584</td>
<td>0.52 (0.49, 0.54)</td>
</tr>
<tr>
<td>Nrf2, rs1806649</td>
<td>8606</td>
<td>0.50 (0.47, 0.53)</td>
</tr>
<tr>
<td>Nrf2, rs1962142</td>
<td>8703</td>
<td>0.50 (0.44, 0.56)</td>
</tr>
<tr>
<td>Nrf2, rs2364723</td>
<td>8725</td>
<td>0.51 (0.49, 0.54)</td>
</tr>
<tr>
<td>Nrf2, rs706649</td>
<td>8731</td>
<td>0.46 (0.41, 0.51)</td>
</tr>
<tr>
<td>Nrf2, rs721961</td>
<td>9419</td>
<td>0.51 (0.46, 0.57)</td>
</tr>
<tr>
<td>Nrf2, rs726295</td>
<td>8692</td>
<td>0.50 (0.48, 0.52)</td>
</tr>
<tr>
<td>Nrf2, rs10183914</td>
<td>8671</td>
<td>0.50 (0.47, 0.52)</td>
</tr>
<tr>
<td>TNF, rs1800629</td>
<td>8685</td>
<td>0.49 (0.46, 0.52)</td>
</tr>
<tr>
<td>UGB, rs3741240</td>
<td>8671</td>
<td>0.48 (0.46, 0.50)</td>
</tr>
</tbody>
</table>

4.1 Reshaping tables using reshape and xrewide

The **reshape** command reshapes a dataset in memory from a long form to a wide form (using its **reshape wide** syntax) or from a wide form to a long form (using its **reshape long** syntax). If we think of tables as Stata datasets, then we can think of using **reshape wide** to convert two versions of table 2 (the child and maternal versions) into table 3. Alternatively, we can think of using **reshape long** to convert table 2 into a table form similar to some of those created by the **esttab** package (Jann 2007), in which the sample numbers, ratios, confidence limits, and p-values are stacked vertically. (This might be done using an integer-valued j() variable named stat, with value labels like “N”, “Ratio”, “95% CI”, and “P”.)

The package **xrewide** is an extended version of **reshape wide** that stores additional results in variable characteristics or local macros. These additional results can be useful if the dataset is output to a table by using **listtab**. To store these results, **xrewide** has the following options, in addition to those available with **reshape wide**: 
1. The `cjvalue()` and `cjlabel()` options specify the names of variable characteristics to be assigned to the generated reshaped output variables and to contain the corresponding values and value labels, respectively, of the j() variable.

2. The `vjvalue()` and `vjlabel()` options specify the names of subsets of the input variables to be reshaped, whose corresponding output variables will be assigned the variable characteristics named by `cjvalue()` and `cjlabel()`, respectively. (By default, all output variables are assigned these characteristics.)

3. The `pjvalue()` and `sjvalue()` options specify a prefix and a suffix, respectively, to be added to the variable characteristic named by `cjvalue()`. The `pjlabel()` and `sjlabel()` options specify a prefix and a suffix, respectively, for the `cjlabel` variable characteristic.

4. The `xmlsub` option specifies that XML escape sequence substitutions be performed for the characters &< and > in the `cjlabel()` variable characteristic.

5. Finally, the `lxjk()` and `lxkj()` options specify the names of local macros, to be assigned with full lists of the generated reshaped output variables. In the case of the `lxjk()` macro, these lists are sorted primarily by the corresponding j() value and secondarily by the corresponding input variable; in the case of the `lxkj()` macro, these lists are sorted primarily by the corresponding input variable and secondarily by the corresponding j() value.

These extra saved results, stored in variable characteristics or local macros, can then be passed to `listtab` and `listtab_vars` to save the user the trouble of manually typing column header rows or variable lists. The saved results can save a lot of work if a colleague suddenly decides that three columns of confidence intervals were wanted instead of two or that the columns need to be relabeled.

### 4.2 The code to create table 3

This code and its Stata log output are as follows:

```
. preserve
. keep source polygp poly N homhet min* max*
. format homhet min* max* %8.2f
. bmjcip homhet min* max*, esub(texsuper) prefix($) suffix($)
. sdecode N, replace esub(texsuper) prefix($) suffix($) supported
. sdecode polygp, generate(S_1)
. sdecode poly, generate(S_2)
. generate row=S_1+, "S_2"
. chardef row N homhet min* max*, char(varname) prefix("\textsl{") suffix("")"
. chardef row N homhet min* max*, char(justify) values(l r r r r)
```

From resultssets to resultstables in Stata

```
. xrewide N homhet min* max*, i(polygp poly) j(source)
   > cjlabel(varname2) vjlabel(N) pjlabel("\textsl{") sjlabel(:")
   > lxjk(nonrowvars)
```

Data summary:

<table>
<thead>
<tr>
<th></th>
<th>long -&gt; wide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of obs.</td>
<td>48 -&gt; 24</td>
</tr>
<tr>
<td>Number of variables</td>
<td>10 -&gt; 13</td>
</tr>
<tr>
<td>j variable (2 values)</td>
<td>source -&gt; (dropped)</td>
</tr>
<tr>
<td>xij variables:</td>
<td>N -&gt; N1 N2</td>
</tr>
<tr>
<td></td>
<td>homhet -&gt; homhet1 homhet2</td>
</tr>
<tr>
<td></td>
<td>min95 -&gt; min951 min952</td>
</tr>
<tr>
<td></td>
<td>max95 -&gt; max951 max952</td>
</tr>
</tbody>
</table>

We start as before by preserving the input resultset, keeping only the key variables and the variables to be tabulated, and formatting the estimates and confidence limits to two decimal places. We then use `bmjcip` as before to decode the estimates and confidence limits (this time without $p$-values). We use `sdecode` to create the row label variable `row` as before. Note that, at this point, the dataset is keyed by the variables `source`, `polygp`, and `poly`, and the two observations for each polymorphism with different values of `source` (containing child and maternal statistics, respectively) will have the same value of `row`.

We then use `chardef` to assign values to two variable characteristics, whose names are `varname` and `justify`, containing, respectively, the variable header cell and the variable justification of the variable in the table (l for left-justified, r for right-justified). We can then use `xrewide` to reshape the dataset to a wide format, keyed by the variables `polygp` and `poly`. The wide format also has two sets of statistical variables, the first containing the sample numbers, estimates, and confidence limits for child genotypes, and the second containing the sample numbers, estimates, and confidence limits for maternal genotypes. Note that the characteristics `varname` and `justify`, assigned to the input statistical variables to be reshaped, are inherited automatically by the corresponding generated reshaped output variables. The output variables N1 and N2 also have the second characteristic `varname2`, as specified by the `cjlabel()` option, containing the value labels of the j() variable `source`, prefixed and suffixed by the `pjlabel()` and `sjlabel()` options, to be slanted and terminated with colons in \LaTeX.

The option `lxjk(nonrowvars)` assigns the local macro `nonrowvars` with the variable list

```
N1 homhet1 min951 max951 N2 homhet2 min952 max952
```
containing the reshaped statistical variables in the order in which they are to be output. We then use `listtab_vars` with the option `rstyle(tabular)` to create header rows in local macros `h1` and `h2` containing the variable characteristics `varname` and `varname2`, respectively. We use `listtab_vars` with no `rstyle()` option to create an unprefixed, unsuffixed, and unsynchronized list of `justify` characteristics in the local macro `just`, with the value `lrrrrrrrr` indicating that the first column of table 3 will be left-justified and the other columns of table 3 will be right-justified.

The `listtab` command then creates a `\LaTeX` `tabular` environment and writes it to the file `texdemo1_3.tex`, where it can be copied and pasted or `\input` into a table in a `\LaTeX` document, which appears as table 3. Finally, we restore the original resultsset to the memory.

5 Gapped tables: Decode, label, reshape, insert gaps, list

Table 3 can be criticized because too much horizontal space is allocated in the first column to repetitive displays of the polymorphism groups (particularly the group “Nrf2”) so that a small font has to be used. Table 4 might be viewed as an improvement on table 3, because each polymorphism group starts with a gap row in which the polymorphism group name is displayed in bold type and left-justified, so it does not need to be repeated in the following rows. The first column therefore needs less horizontal space, allowing the other columns to have more. The gap rows make the table less dense and more clear, at the price of using more vertical space. To insert gap rows into tables, we use the `ingap` package.
Table 4. Child and maternal homozygote or heterozygote ratios (with gaps)

<table>
<thead>
<tr>
<th>Polymorphism (by group)</th>
<th>Child:</th>
<th>Maternal:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Ratio (95% CI)</td>
</tr>
<tr>
<td>AHR</td>
<td>rs2066853</td>
<td>8704</td>
</tr>
<tr>
<td>CYP1A1</td>
<td>rs1048943</td>
<td>8764</td>
</tr>
<tr>
<td></td>
<td>rs4646903</td>
<td>8587</td>
</tr>
<tr>
<td>CYP2A6</td>
<td>rs1801272</td>
<td>8666</td>
</tr>
<tr>
<td></td>
<td>rs28399433</td>
<td>8691</td>
</tr>
<tr>
<td>GCL</td>
<td>rs17883901</td>
<td>9389</td>
</tr>
<tr>
<td>GPX</td>
<td>rs713041</td>
<td>8611</td>
</tr>
<tr>
<td>GSTM1</td>
<td>GSTM1 CNV</td>
<td>8399</td>
</tr>
<tr>
<td>GSTP1</td>
<td>rs947894</td>
<td>8692</td>
</tr>
<tr>
<td>GSTT1</td>
<td>GSTT1 CNV</td>
<td>7591</td>
</tr>
<tr>
<td>IGF</td>
<td>rs35767</td>
<td>9383</td>
</tr>
<tr>
<td></td>
<td>rs2854744</td>
<td>8489</td>
</tr>
<tr>
<td>LTA</td>
<td>rs909253</td>
<td>8664</td>
</tr>
<tr>
<td>MIF</td>
<td>rs755622</td>
<td>9500</td>
</tr>
<tr>
<td>MTNR1B</td>
<td>rs10830963</td>
<td>8584</td>
</tr>
<tr>
<td>Nrf2</td>
<td>rs1806649</td>
<td>8606</td>
</tr>
<tr>
<td></td>
<td>rs1962142</td>
<td>8763</td>
</tr>
<tr>
<td></td>
<td>rs2364723</td>
<td>8725</td>
</tr>
<tr>
<td></td>
<td>rs6706649</td>
<td>8731</td>
</tr>
<tr>
<td></td>
<td>rs6721961</td>
<td>9419</td>
</tr>
<tr>
<td></td>
<td>rs6726395</td>
<td>8692</td>
</tr>
<tr>
<td></td>
<td>rs10183914</td>
<td>8671</td>
</tr>
<tr>
<td>TNF</td>
<td>rs1800629</td>
<td>8685</td>
</tr>
<tr>
<td>UGB</td>
<td>rs3741240</td>
<td>8671</td>
</tr>
</tbody>
</table>
5.1 Inserting gap rows by using ingap

The `ingap` package is downloadable from SSC and was described briefly in Newson (2003). It is a comprehensive package for inserting user-selected numbers of gap observations into user-selected positions in a dataset or by-group, with user-selected values assigned to the table row variable.

However, usually most users will probably only want to insert one gap observation at the beginning of each of a number of by-groups and to set the row variable in each of these new observations to be equal to a preexisting gap-row string variable, whose value will be constant within each by-group. The syntax for doing this is

```
by sort keyvarlist_1 gaprowvarname (keyvarlist_2): ingap, neworder(newvar)
rowlabel(varname) grexpression(gap_row_label_expression)
```

where `keyvarlist_1` is a list of key variables defining the by-groups that will each have one gap observation added at the beginning, `gaprowvarname` is the name of the preexisting gap-row string variable, `keyvarlist_2` is a list of key variables defining the order of preexisting observations within the by-group, `varname` is the name of the preexisting string row label variable, and `newvar` is the name of a new variable to be generated by `ingap` containing in each observation the sequential order of that observation within the by-group once the gap observations have been added. The dataset is usually keyed by the variables `keyvarlist_1` and `keyvarlist_2` before the execution of `ingap` and is keyed by `keyvarlist_1`, `gaprowvarname`, and `newvar` after the execution of `ingap`. Therefore, the variable `newvar` defined in `neworder()` will be 1 for the gap observation in each by-group defined by the `keyvarlist_1` and will have integer values starting from 2 in the remaining observations in each by-group.

The gap-row variable specified by `gaprowvarname` is defined, usually using `sdecode`, as a function of the variables in `keyvarlist_1` and will therefore be constant within each by-group.

For instance, in our example resultsset, once the decoding, labeling, and reshaping have been done and a gap-row variable `gaprow` has been defined as a function of the polymorphism group variable `polygp`, the command

```
. by sort polygp gaprow (poly): ingap, rowlabel(row) grexpression(gaprow)
> neworder(rowseq)
```

will insert a gap row at the beginning of each by-group defined by `polygp`, will set the value of the string variable `row` in that gap row to be equal to the value of `gaprow` in that by-group, and will create a new integer-valued variable `rowseq` equal to 1 in the gap observation and to integers starting at 2 in the other observations in the by-group in the ascending order defined by the variable `poly`. The dataset will then be keyed by `polygp`, `gaprow`, and `rowseq`. 
5.2 The code to create table 4

The code to do this in the example resultsset and its Stata log is as follows:

```
. preserve
  . keep source polygp poly N homhet min* max*
  . format homhet min* max* %3.2f
  . bmjchip homhet min* max*, esub(texsuper) prefix($) suffix($)
  . sdecode N, replace esub(texsuper) prefix($) suffix($)
  . sdecode poly, generate(row) prefix("\hfill ")
  . chardef row N homhet min* max*, char(varname) prefix("\textsl{") suffix("}
  > values("Polymorphism (by group)" N Ratio "(95\% "CI")")
  . chardef row N homhet min* max*, char(justify) values("p{1in}" r r r)
  . xrewide N homhet min* max*, i(polygp poly) j(source)
  > cjlabel(varname2) vjlabel(N) pjlabel("\textsl{") ajlabel(:)
  > lxjk(nonrowvars)

Data

<table>
<thead>
<tr>
<th></th>
<th>long →</th>
<th>wide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of obs.</td>
<td>48</td>
<td>24</td>
</tr>
<tr>
<td>Number of variables</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
<td>j variable (2 values)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xij variables:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>→ N1 N2</td>
<td></td>
</tr>
<tr>
<td>homhet</td>
<td>→ homhet1 homhet2</td>
<td></td>
</tr>
<tr>
<td>min95</td>
<td>→ min951 min952</td>
<td></td>
</tr>
<tr>
<td>max95</td>
<td>→ max951 max952</td>
<td></td>
</tr>
</tbody>
</table>
```

```
. sdecode polygp, generate(gaprow) prefix("\textbf{") suffix("}\hfill")
  . bysort polygp gaprow (poly): ingap, rowlabel(row) grexpression(gaprow)
  > neworder(rowseq)
  . listtab_vars row `nonrowvars´, rstyle(tabular) substitute(char varname) local(h1)
  . listtab_vars row `nonrowvars´, rstyle(tabular) substitute(char varname2)
  > local(h2)
  . listtab_vars row `nonrowvars´, substitute(char justify) local(just)
  . listtab row `nonrowvars´ using texdemo1_4.tex, replace rstyle(tabular)
  > headlines("\begin{tabular}{`just´}" \hline "`h2´" \hline "`h1´" \hline")
  > footlines("\hline" "\end{tabular}

. restore
```

Once again, we start by preserving the original resultsset and proceed as we did for table 3 up to `xrewide`. Here though, the row label variable `row` now depends only on `poly` and not on `polygp`. The characteristic `justfy`, for this row variable, is set to `p{1in}`, implying that this row variable will be neither left-justified nor right-justified but will have a fixed width of 1 inch in the table, with justification depending on the value of the variable `row`. In the original variable `row`, before the addition of the gap observations, the values are all prefixed with `\hfill`, which causes the values to be right-justified.

Once `xrewide` has been executed, producing a wide dataset keyed by `polygp` and `poly` as before, we use `sdecode` to produce the gap-row variable `gaprow`, containing the
decoded value of polygp, prefixed by \textbf{ and suffixed by }\hfill, which implies a bold font and left-justification. We then execute ingap, adding the gap observations. After that, we can use listtab \texttt{vars} three times as before, use listtab once as before (outputting the tabular environment to the file texdemo1.4.tex), and restore the original resultsset. The tabular environment can then be pasted or \texttt{\input} into a \LaTeX document to form table 4.

6 Creating RTF documents using rtfutil

listtab can create tables in many formats other than \LaTeX. In particular, it can be used with the SSC package rtfutil to create tables in Microsoft RTF, described in plain language by Burke (2003).

RTF tables are less simple to produce than \LaTeX tables for two reasons:

1. It is not usually a good idea to cut and paste an RTF table into an RTF document by using a text editor (although it can be done). This is because RTF documents (especially those produced using Microsoft Word) may contain a lot of incomprehensible code, making it difficult to guess where the table should be pasted. If it is pasted in the wrong place, then Microsoft Word will often fail un informatively when it tries to open the document.

2. There is no unique RTF row style, defined by \texttt{begin()}, \texttt{delimiter()}, \texttt{end()}, and \texttt{missnum()} options for listtab. This is because in RTF tables, the number of cells per row, their widths, and other formatting features are defined separately for each row as part of the \texttt{begin()} string or the \texttt{end()} string.

The first of these problems is solved using the \texttt{handle()} option of listtab, which allows listtab to output to an existing open file with a file handle as created by the \texttt{file} command of Stata (see \texttt{[P] file}). The rtfutil package has the command rtfopen, which opens an RTF file by using \texttt{file open} and outputs the beginning of an RTF document, and the command rtfclose, which outputs the end of an RTF document and closes the file by using \texttt{file close}. The listtab commands to create tables can then be inserted between the rtfopen and rtfclose commands.

The second of these problems is solved using the \texttt{rtfrstyle} command of rtfutil, which creates RTF row styles. It inputs a list of variables to be output by listtab, with their column widths and other formatting information, and it outputs to a list of local macros specified in a \texttt{local()} option, which will contain the \texttt{begin()}, \texttt{delimiter()}, \texttt{end()}, and possibly \texttt{missnum()} strings that will later be input to listtab. Column widths are specified using the \texttt{cwidths()} option of rtfrstyle and are expressed in twips (1440 twips = 1 inch), which are the units used internally by RTF documents.
From resultsets to resultstables in Stata

It is a good idea for all commands between an *rtfopen* command and an *rtfclose* command to be enclosed in a *capture noisily* block. This ensures that if any of these commands fails, then the RTF file will automatically be closed and will be available for inspection by the user.

### 6.1 The code to create an RTF version of table 4

In this example, we start with the example resultset and create an RTF document *rtfdemo1.rtf* containing an RTF version of table 4. The code is enclosed in a document-generating block starting with a *tempname* command to create a file handle name, followed by an *rtfopen* command, followed by the beginning of a *capture noisily* block, and ending with an *rtfclose* command preceded by the end of the *capture noisily* block.

```
. #delim ;
. /* Beginning of document-generating block */ ;
. tempname rtfb1;
. rtfopen `rtfb1´ using "rtfdemo1.rtf", replace
> margins(1000 1000 1000 1000) template(fnmono1);
. capture noisily {
. /* Create table */
. preserve;
. keep source polygp poly N homhet min* max*;
. format homhet min* max* %8.2f;
. bmjcip homhet min* max*, esub(rtfsuper) pref("\qr{") suff(\"\"));
. sdecode N, replace esub(rtfsuper) pref("\qr{") suff(\"\"));
. sdecode poly, generate(row) pref("\qr{") suff(\"\"));
. chardef row N homhet min* max*, char(varname) pref("\qr{\i ") suff("\i ")
> val("Polymorphism (by group)" N Ratio "(95\% "CI")");
. xrewide N homhet min* max*, i(polygp poly row) j(source)
> cjlab(varname2) vjlab(N) pjlab("\ql{\i ") sjlab(" DNA:}"
> lxjk(nonrow);
. sdecode polygp, generate(gaprow) pref("\ql{\b ") suff(\"\")
> bysort polygp gaprow (poly): ingap, row(row) grex(gaprow) neword(rouseq);
. rtfrstyle row `nonrow´, cwidths(1500 1000) local(b d e);
. listtab_vars row `nonrow´, sub(char varname) b(`b´) d(`d´) e(`e´) local(h1);
. listtab_vars row `nonrow´, sub(char varname2) b(`b´) d(`d´) e(`e´) local(h2);
. file write `rtfb1´
> "{\pard\b"
> "Geometric mean homozygote/heterozygote ratios"
> " for biallelic polymorphisms in child and maternal DNA"
> "\par}"
> _n;
```
Inside the `capture noisily` block, the commands up to `ingap` are mostly similar to the corresponding commands to create table 4 except that the prefixes and suffixes are RTF prefixes and suffixes. The prefix–suffix pair `{ql \{ and \} specifies left-justification, the prefix–suffix pair `{qr \{ and \} specifies right-justification, and both pairs can be used with or without the added `\i` or `\b` appended to the prefix to specify italic or bold font, respectively.

The `rtfstyle` command inputs the variables to be output by `listtab`, together with the option `cwidths(1500 1000)` to specify that the first column have a width of 1500 twips and that all other columns have widths of 1000 twips, and with the `local()` option to specify that the `begin()`, `delimiter()`, and `end()` strings be stored in the local macros `b`, `d`, and `e`, respectively. These local macros are used by `listtab_vars` to create the header rows, stored in the local macros `h1` and `h2`, and by `listtab` to output the table to the RTF file after the RTF table heading has been output using a `file write` command. The RTF file produced, with the name `rtfdemo1.rtf`, is part of the supplemental online material for this article and can be downloaded using Stata.

## 7 Possible extensions

The team of packages `listtab`, `sdecode`, `chardef`, `xrewide`, and `ingap` can be used with document formats other than TeX and RTF, such as HTML and possibly XML-based formats yet to be invented. The packages are good at enforcing the nested block structure commonly used in such formats, because each `prefix()` option of `sdecode` or `chardef` can have its `suffix()` option, each `pjlabel()` option of `xrewide` can have its `sjlabel()` option, each `begin()` option of `listtab` can have its `end()` option, and each `headlines()` option of `listtab` can have its `footlines()` option. Likewise, each `rtfopen` command (followed by the beginning of a `capture noisily` block) can have its `rtfclose` command (preceded by the end of the same `capture noisily` block).

It is possible to produce parallel column groups nested in larger parallel column groups by using multiple calls to `xrewide`. Similarly, it is possible to produce gap-prefixed by-groups nested in larger gap-prefixed by-groups by using multiple calls to `ingap`. The smaller by-groups might be preceded by gap-row labels in a smaller bold font, while the larger by-groups might be preceded by gap-row labels in a larger bold font.
It is also possible to produce multipage tables in a document by putting a \texttt{listtab} command with an \texttt{if} qualifier inside a program loop. Alternatively, we can put the whole set of commands from \texttt{preserve} to \texttt{restore} inside a program loop and include a \texttt{keep if} command to select the subset of observations for a page.

It should also be possible to write user-friendly front-end packages by using the low-level programming toolkit described here to produce standard documents, in particular formats containing standard-format tables. So far, I have not done this, because my colleagues and collaborators seem to prefer highly customized RTF resultstables to standard-format RTF resultstables.

Finally, a Stata program that inputs a resultset to output resultstables to \TeX, RTF, and other documents can also output resultplots of the same results to the same documents, produced using Stata graphics commands (see \cite{G-2} graph). In particular, the rtutil package includes an rtflink command to include graphical output files, such as Encapsulated PostScript files produced using graph export, as linked objects in an RTF document. The user may then convert these linked objects to embedded objects by using Microsoft Word. I find that this possibility is a major advantage of a resultset-based solution over the purely table-based solutions used by \texttt{estout}, \texttt{outreg}, \texttt{outreg2}, and \texttt{tabout}.

8 Acknowledgments

I would like to thank Ben Jann of the Eidgenössische Technische Hochschule in Zurich, Switzerland, and Adam Jacobs of Dianthus Medical Limited in London, UK, for their presentations at the 2009 UK Stata Users Group meeting, which inspired a lot of the packages and updates presented in this article. I would also like to thank Philippe Van Kerm of CEPS/INSTEAD in Luxembourg for suggesting that a \texttt{handle()} option for listtex (predecessor to listtab) would be a good idea; Ian White of the MRC Biostatistics Unit in Cambridge, UK, for suggesting that something like the listtab \texttt{vars} command of listtab would be a good idea; and Lindsey Peters of the United Kingdom Health Protection Agency for some very helpful feedback about RTF table rows output with the rtutil package. I would also like to thank Nicholas J. Cox of Durham University, UK, for coining the term “resultset” to describe a Stata dataset of results.

I would also like to thank our collaborators in the ALSPAC Study Team (Institute of Child Health, University of Bristol, Bristol, UK) for allowing the use of their data in this article. In particular, I would like to thank my ALSPAC collaborators John Holloway, John Henderson, and Seif Shaheen for helping me to define the gene labels in tables 2, 3, and 4 of this article. The whole ALSPAC Study Team comprises interviewers, computer technicians, laboratory technicians, clerical workers, research scientists, volunteers, and managers who continue to make the study possible. The ALSPAC study could not have been undertaken without the cooperation and support of the mothers and midwives who took part or without the financial support of the Medical Research Council, the Department of Health, the Department of the Environment, the Wellcome Trust, and other funders. The ALSPAC study is part of the World Health Organization–initiated
European Longitudinal Study of Pregnancy and Childhood. My own work at Imperial College London is financed by the United Kingdom Department of Health.

9 References


About the author

Roger B. Newson is a lecturer in medical statistics at the National Heart and Lung Institute, Imperial College London, UK, working principally in asthma research. He has written several Stata packages that are frequently used together and that define a dialect of the Stata language. These packages can all be installed as a group in versions compatible with Stata 9, 10, 11, or 12 by using Stata do-files (one for each Stata version), which users can download from http://www.imperial.ac.uk/nhli/r.newson/stata.htm.