Confidence intervals for rank statistics: Percentile slopes, differences, and ratios

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Abstract. I present a program, censlope, for calculating confidence intervals for generalized Theil–Sen median (and other percentile) slopes (and per-unit ratios) of \( Y \) with respect to \( X \). The confidence intervals are robust to the possibility that the conditional population distributions of \( Y \), given different values of \( X \), differ in ways other than location, such as having unequal variances. censlope uses the program somersd and is part of the somersd package. censlope can therefore estimate confounder-adjusted percentile slopes, limited to comparisons within strata defined by values of confounders, or by values of a propensity score representing multiple confounders. Iterative numerical methods have been implemented in the Mata language, enabling efficient calculation of percentile slopes and their confidence limits in large samples. I give example analyses from the \texttt{auto} dataset and from the Avon Longitudinal Study of Pregnancy and Childhood (ALSPAC).

Keywords: snp15_7, somersd, censlope, ALSPAC, robust, confidence interval, rank, nonparametric, median, percentile, slope, difference, ratio, Kendall’s \( \tau \), Somers’ \( D \), Theil–Sen, Hodges–Lehmann, confounder adjusted, propensity score

1 Introduction

The Theil–Sen median slope is a rank-based parameter, defined in terms of Kendall’s \( \tau \), but expressed in \( y \)-axis units per \( x \)-axis unit and interpreted as a “typical” difference in \( Y \) associated with a unit difference in \( X \). Knowing this slope is therefore useful if we want to use rank methods to make monetary or other practical decisions. It was introduced by Theil (1950) and developed by Sen (1968), who derived a confidence interval formula. If the \( X \) variable is binary, then the Theil–Sen median slope is known as the Hodges–Lehmann median difference and is expressed in \( y \)-axis units. This median difference was introduced by Hodges and Lehmann (1963) and developed by Lehmann (1963), who derived a confidence interval formula that is a special case of the one in Sen (1968).

The median difference was made popular by Conover (1980), Campbell and Gardner (1988), and Altman et al. (2000) and implemented in Stata by Wang (1999) and in Patrick Royston’s \texttt{ssc} package \texttt{cid}. Sprent and Smeeton (2001) gives a good general introduction to confidence interval formulas for median slopes and differences.

Most existing confidence interval formulas for median slopes and differences assume that, if \( \beta \) is the median slope, \( Y - \beta X \) is statistically independent of \( X \). This independence in turn implies that the conditional distributions of \( Y \), given different values of \( X \), differ only in location and not in other ways such as unequal variance. These
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problems are discussed in Wilcox (1998), which describes a possible solution using the percentile bootstrap. Also the median differences and slopes are usually defined as crude differences and slopes, assumed to apply to the whole population, and not as adjusted differences and slopes, assumed to apply within subpopulations with similar values of stratification or confounding variables. These assumptions may limit the usefulness of these confidence interval formulas.

In a previous paper (Newson 2002), I argued that median differences and slopes belong to a unified family of rank parameters, with a unified system of confidence interval formulas. In this family of parameters, median differences and slopes are naturally defined in terms of Somers’ $D$, which in turn is naturally defined in terms of Kendall’s $\tau_a$. This paper introduced the _somersd_ package, downloadable from SSC, as a way of calculating some of these confidence intervals. The _somersd_ package then contained two modules, namely, _somersd_, described in Newson (2000a), for calculating confidence intervals for Somers’ $D$ and Kendall’s $\tau_a$, and _cendif_, described in Newson (2000b), for calculating confidence intervals for Hodges–Lehmann median differences. In 2005, Stata 9 introduced the Mata programming language, which made it possible to update _somersd_ to estimate many extended versions of Somers’ $D$ and Kendall’s $\tau_a$ and to do so more quickly. This update was reported in Newson (2006b), which contains the syntax, formulas, and methods, and in Newson (2006a), which describes the Mata algorithm used.

This article describes a third module, _censlope_, which has been added to the _somersd_ package in a recent update and which estimates generalized Theil–Sen median (and other percentile) slopes, differences, and ratios. In particular, these slopes, differences, and ratios may be adjusted for confounding variables, allowing the user to use rank methods to answer many questions that could previously be answered only by using regression methods. In section 2, I describe the current version of the program _censlope_. In section 3, I present, for reference, the methods and formulas used by _censlope_. In section 4, I demonstrate a range of examples.

2 The program _censlope_

2.1 Syntax

```
censlope yvarname xvarname [if] [in] [weight] [, centile(numlist) eform 
   _ystargenerate(newvarlist) estaddr somersd_options iteration_options]
```

where _yvarname_ and _xvarname_ are variable names.

_fweights_, _iweights_, and _pweights_ are allowed; see [U] 11.1.6 weight. They are interpreted as for _somersd_.

_bootstrap_, _by_, _jackknife_, and _statsby_ are allowed; see [U] 11.1.10 Prefix commands.
2.2 Description

censlope calculates confidence intervals for generalized Theil–Sen median slopes and other percentile slopes of a Y variable specified by yvarname with respect to an X variable specified by xvarname. These confidence intervals are robust to the possibility that the population distributions of Y, conditional on different values of X, are different in ways other than location. This difference might happen if, for example, the conditional distributions had different variances. For positive-valued Y variables, censlope can be used to calculate confidence intervals for median per-unit ratios or other percentile per-unit ratios associated with a unit increment in X. If X is binary with values 0 and 1, then the generalized Theil–Sen percentile slopes are the generalized Hodges–Lehmann percentile differences between the group of observations whose X value is 1 and the group of observations whose X value is 0. censlope is part of the somersd package and requires the somersd program to work. It executes the somersd command,

\texttt{somersd xvarname yvarname [if] [in] [weight] [, somersd_options]}

and then estimates the percentile slopes. The estimates and confidence limits for the percentile slopes are evaluated using an iterative numerical method, which the user may change from the default, using iteration_options.

2.3 Options

centile\texttt{(numlist)} specifies a list of percentile slopes to be reported and defaults to centile\texttt{(50)} (median only) if not specified. Specifying centile\texttt{(25 50 75)} will produce the 25th, 50th, and 75th percentile differences.

eform specifies that exponentiated percentile slopes be given. This option is used if yvarname specifies the log of a positive-valued variable. Then confidence intervals are calculated for percentile ratios or per-unit ratios between values of the original positive variable instead of for percentile differences or per-unit differences.

ystargenerate\texttt{(newvarlist)} specifies a list of variables to be generated, corresponding to the percentile slopes, containing the differences \( Y^*(\beta) = Y - \beta X \), where \( \beta \) is the percentile slope. The variable names in the newvarlist are matched to the list of percentiles specified by the centiles\texttt{()} option, sorted in ascending order of percentage. If the two lists have different lengths, then censlope generates a number nmin of new variables equal to the minimum length of the two lists, matching the first nmin percentiles with the first nmin new variable names. Usually, there is only one percentile slope (the median slope) and one new ystargenerate\texttt{()} variable, whose median can be used as the intercept when drawing a line through the data points on a scatterplot.

estaddr specifies that the results saved in \texttt{r()} also be saved in \texttt{e()} (see section 2.5). This option makes it easier to use censlope with parmby, to create an output dataset (or resultsset) with 1 observation per by-group and data on confidence intervals for
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Somers’ D and median slopes. parmby is part of the package parmest, downloadable from SSC. The online help for censlope gives an example using the estaddr option with parmby.

somersd_options are any of the options used by somersd.

2.4 Iteration options

Table 1: Iteration options used by censlope

<table>
<thead>
<tr>
<th>iteration_options</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>fromabs(#)</td>
<td>initial estimate for absolute magnitude of slopes</td>
</tr>
<tr>
<td>brackets(#)</td>
<td>maximum number of rows for the bracket matrix</td>
</tr>
<tr>
<td>technique(algorithm_spec)</td>
<td>iterative numerical solution technique</td>
</tr>
<tr>
<td>iterate(#)</td>
<td>perform maximum of # iterations; default is iterate(16000)</td>
</tr>
<tr>
<td>tolerance(#)</td>
<td>tolerance for the percentile slopes</td>
</tr>
<tr>
<td>log</td>
<td>display an iteration log of the brackets during bracket</td>
</tr>
<tr>
<td></td>
<td>convergence</td>
</tr>
</tbody>
</table>

where algorithm_spec is

\[algorithm \ [ \ # \ [ \ algorithm \ [ \ # \ ] \ ] \ ...]\n
and algorithm is \{bisect|regula|ridders\}

The censlope command calculates estimates and confidence limits for a median or other percentile slope \( \beta \) by solving numerically a scalar equation in \( \beta \) with an iterative method. The options controlling the exact iterative method will probably not be used often, because censlope is intended to have sensible defaults. Nontechnical readers may therefore skip this section. However, users who wish to change the default method may do so using a set of options similar to the maximization options used by Stata’s maximum likelihood estimation commands (see [R] maximize). These options are listed in table 1 and are described as follows:

fromabs(#) specifies an initial estimate of the typical absolute magnitude of a percentile slope. If fromabs() is not specified, it defaults to the aspect ratio \( (ymax – ymin)/(xmax – xmin) \) (where xmax and xmin are the maximum and minimum X values and ymax and ymin are the maximum and minimum Y values) if that ratio is defined and nonzero and to 1 otherwise. This magnitude is used in constructing the bracket matrix. Candidate bracket \( \beta \) values will have values of zero or of \( \pm \text{fromabs} \times 2^K \), where \( K \) is a nonnegative integer. The bracket matrix is a matrix with two columns and three or more rows, each row containing a candidate \( \beta \) value.
in column 1 and the corresponding $\zeta^*$ value in column 2. The bracket matrix is used to find an initial pair of $\beta$ values for input into the iterative numerical solution method, which attempts to find a solution in $\beta$ between the two initial $\beta$ values. The bracket matrix is initialized to have $\beta$ values $-\text{fromabs}$, 0, and $+\text{fromabs}$, as well as $\zeta^*$ values corresponding to these $\beta$ values. If a target $\zeta$ value is outside the range of the $\zeta^*$ values of the bracket matrix, then the bracket matrix is extended by adding new rows before the first row by successively doubling the $\beta$ value in the first row or by adding new rows after the last row by successively doubling the $\beta$ value in the last row, until there is a $\zeta^*$ value in the second column on either side of the target $\zeta$ value. For an explanation of this terminology, see section 3.

The bracket matrix is initialized to have $\beta$ values $-\text{fromabs}$, 0, and $+\text{fromabs}$, as well as $\zeta^*$ values corresponding to these $\beta$ values. If a target $\zeta$ value is outside the range of the $\zeta^*$ values of the bracket matrix, then the bracket matrix is extended by adding new rows before the first row by successively doubling the $\beta$ value in the first row or by adding new rows after the last row by successively doubling the $\beta$ value in the last row, until there is a $\zeta^*$ value in the second column on either side of the target $\zeta$ value. For an explanation of this terminology, see section 3.

$\textit{brackets(\#)}$ specifies a maximum number of rows for the bracket matrix. The minimum is $\text{brackets(3)}$. The default is $\text{brackets(1000)}$.

$\textit{technique(\text{algorithm_spec})}$ specifies an iterative solution method for finding a solution in $\beta$ to the equation to be solved. The following algorithms are currently implemented in $\text{censlope}$:

- $\textit{technique(bisect)}$ specifies an adapted version of the bisection method for step functions.
- $\textit{technique(regula)}$ specifies an adapted version of the regula falsi (or false position) method for step functions.
- $\textit{technique(ridders)}$ specifies an adapted version of the method of Ridders (1979) for step functions.

The default is $\textit{technique(ridders 5 bisect iterate)}$, where $\text{iterate}$ is the value of the $\text{iterate()}$ option. The bisection method is guaranteed to converge in a number of iterations similar to the binary logarithm of the $\text{tolerance()}$ option. The regula falsi and Ridders methods are usually faster if the $\zeta^*$ function is nearly continuous but may sometimes be slower if the $\zeta^*$ function is a discrete-step function. All methods are modified versions for step functions of the methods of the same names described in Press et al. (1992).

You can switch between algorithms by specifying more than one in the $\text{technique()}$ option. By default, $\text{censlope}$ will use an algorithm for five iterations before switching to the next algorithm. To specify a different number of iterations, include the number after the technique in the option. For example, specifying the option $\text{technique(ridders 10 bisect 1000)}$ requests that $\text{censlope}$ perform 10 iterations with the Ridders algorithm, perform 1,000 iterations with the bisection algorithm, and then switch back to Ridders for 10 iterations, and so on. The process continues until convergence or until the maximum number of iterations is reached.

$\textit{iterate(\#)}$ specifies the maximum number of iterations. When the number of iterations equals $\text{iterate()}$, the iterative solution program stops and records failure to converge. If convergence is declared before this threshold is reached, it will stop
when convergence is declared. The default value of \texttt{iterate(\#)} is the current value of \texttt{set maxiter}, which is \texttt{iterate(16000)} by default.

\texttt{tolerance(\#)} specifies the tolerance for the percentile differences. When the relative difference between the current $\beta$ brackets is less than or equal to \texttt{tolerance()}, the \texttt{tolerance()} convergence criterion is satisfied. \texttt{tolerance(1e-6)} is the default.

\texttt{log} specifies that an iteration log showing the progress of the numerical solution method be displayed. If an iteration log is displayed, there will be four separate iteration sequences per percentile, estimating the left estimate, the right estimate, the lower confidence limit, and the upper confidence limit. For this reason, the default is not to produce an iteration log. However, if \texttt{censlope} is expected to be slow (as for large datasets), an iteration log can be specified to reassure the user that progress is being made.

### 2.5 Saved results

\texttt{censlope} saves the following results in \texttt{r(\)}:

Scalars
- \texttt{r(level):} confidence level
- \texttt{r(fromabs):} value of the \texttt{fromabs()} option
- \texttt{r(tolerance):} value of the \texttt{tolerance()} option

Macros
- \texttt{r(yvar):} name of the \texttt{Y} variable
- \texttt{r(xvar):} name of the \texttt{X} variable
- \texttt{r(eform):} \texttt{eform} if specified
- \texttt{r(centiles):} list of percentages for the percentiles
- \texttt{r(technique):} list of techniques from the \texttt{technique()} option
- \texttt{r(techn_steps):} list of step numbers for the techniques

Matrices
- \texttt{r(cimat):} confidence intervals for percentile differences or ratios
- \texttt{r(rcmat):} return codes for entries of \texttt{r(cimat)}
- \texttt{r(bracketmat):} bracket matrix
- \texttt{r(techstepmat):} column vector of step numbers for the techniques

The matrix \texttt{r(cimat)} has one row per percentile and columns containing the percentages, percentile estimates, lower confidence limits, and upper confidence limits, labeled \texttt{Percent}, \texttt{Pctl Slope}, \texttt{Minimum}, and \texttt{Maximum} if \texttt{eform} is not specified, or \texttt{Percent}, \texttt{Pctl Ratio}, \texttt{Minimum}, and \texttt{Maximum} if \texttt{eform} is specified. The matrix \texttt{r(rcmat)} has the same numbers of rows and columns as \texttt{r(cimat)}, with the same labels, and the first column contains the percentages, but the other entries contain return codes for the estimation of the corresponding entries of \texttt{r(cimat)}. These return codes are equal to 0 if the $\beta$ value was estimated successfully, 1 if the corresponding $\zeta^*$ value could not be calculated, 2 if the corresponding $\zeta^*$ value could not be bracketed, 3 if the $\beta$ brackets failed to converge, and 4 if the $\beta$ value could not be calculated from the converged $\beta$ brackets. The matrix \texttt{r(bracketmat)} is the final version of the bracket matrix described in the help for the \texttt{fromabs()} and \texttt{brackets()} options of \texttt{censlope} and has one row per $\beta$ bracket and two columns labeled \texttt{Beta} and \texttt{Zetastar} containing the $\beta$ brackets and the corresponding $\zeta^*$ values. The matrix \texttt{r(techstepmat)} is a column vector, with
one row for each of the techniques listed in the \texttt{technique()} option, with a row label
equal to the name of the technique and a value equal to the number of steps for that
technique. The \texttt{fromabs()}, \texttt{brackets()}, \texttt{tolerance()}, and \texttt{technique()} options are
described in section 2.4 above.

censlope also saves in \texttt{e()} a full set of estimation results for the \texttt{somersd} command
as described in section 2.2 above. If \texttt{estaddr} is specified, this set of estimation results
is expanded by adding a set of \texttt{e()} results with the same names and contents as the
\texttt{r()} results. This expansion allows the user to pass a censlope command to \texttt{parmby},
producing an output dataset (or resultsset) with 1 observation per by-group and data
on confidence intervals for Somers’ $D$ and for the median slope.

3 Methods and formulas

This section is intended mainly as a reference for the extensive family of methods and
formulas used by the censlope program. Less technically minded readers may skip or
skim through this section and progress to the Examples.

The Theil–Sen median slope was introduced by Theil (1950) and developed further
by Sen (1968). If $X$ is binary with values 0 and 1, the Theil–Sen slope is the Hodges–
Lehmann median difference of Hodges and Lehmann (1963) and Lehmann (1963). The
methods used by censlope are a generalization of the methods of Theil and Sen. They
include, as a special case, the methods used by cendif (Newson 2000b), which calculates
confidence intervals for generalized Hodges–Lehmann median differences and is part of
the somersd package. However, cendif (like ttest) estimates the median difference
between $Y$ values associated with the smaller $X$ value and $Y$ values associated with
the larger $X$ value, whereas censlope (like regress), given a binary $X$ variable with
values 0 and 1, estimates the median difference between $Y$ values associated with the
larger $X$ value and $Y$ values associated with the smaller $X$ value.

Percentile slopes are defined in terms of the parameters Somers’ $D$ (Somers 1962)
and Kendall’s $\tau_a$ (Kendall and Gibbons 1990). A discussion of the connections between
these parameters appears in Newson (2002). For censlope, we will define Somers $D$
and Kendall’s $\tau_a$ in the general sense used in Newson (2006b). Given two random
variables $U$ and $V$, we denote by $\tau(U,V)$ the Kendall’s $\tau_a$ of $U$ and $V$ and denote by
$D(U|V)$ the Somers’ $D$ of $U$ with respect to $V$. In brief, if two $(U,V)$ pairs, $(U_i, V_i)$
and $(U_j, V_j)$, are sampled from some population of such pairs by using some sampling
scheme, $\tau(U,V)$ is the difference between the probability that the two $(U,V)$ pairs are
concordant (meaning that the larger $U$ value is paired with the larger $V$ value) and the
probability that the two $(U,V)$ pairs are discordant (meaning that the larger $U$ value
is paired with the smaller $V$ value). I define $D(U|V)$ as the difference between the
corresponding conditional probabilities, given that the two $V$ values are strictly ordered
(meaning that one $V$ value is known to be larger than the other $V$ value). Both $\tau(U,V)$
and $D(U|V)$ are differences between probabilities, and therefore both may have values
ranging from $-1$ (for a “perfect negative association”) to $+1$ (for a “perfect positive
association”), but $\tau(U,V)$ is always symmetric in $U$ and $V$, whereas $D(U|V)$ is not. I
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will use the notation $\theta(U,V)$ to stand for the value of either $\tau(U,V)$ or $D(U|V)$ in the population and denote the corresponding sample value as $\hat{\theta}(U,V)$. The somersd package allows us to choose between Somers’ $D$ and Kendall’s $\tau_a$ by using the taua option and provides other options to specify a version of either parameter corresponding to a specific sampling scheme.

For an outcome variable $Y$, a predictor variable $X$, and a proportion $q$ such that $0 \leq q \leq 1$, a 100$q$th percentile slope of $Y$ with respect to $X$ is defined as a value $\beta$ satisfying

$$\theta(Y - \beta X, X) = 1 - 2q$$

If $q = 0.5$, then $1 - 2q = 0$, and a solution in $\beta$ to (1) is known as a Theil–Sen median slope, as defined in Theil (1950) and Sen (1968). There is not always a unique solution to (1) in $\beta$. If the joint population distribution of $Y$ and $X$ is discrete (as are all population distributions sampled by applied statisticians in the real world), $\theta(Y - \beta X, X)$ will be a monotonically nonincreasing step function of $\beta$, and there may be no exact solution or an interval of exact solutions. However, the confidence intervals derived here will contain any solution with the specified confidence level, if a solution exists.

If $\theta(\cdot, \cdot)$ stands for Somers’ $D$ rather than Kendall’s $\tau_a$, the value of the parameter $\theta(Y - \beta X, X)$ depends only on the conditional distribution of pairs of bivariate observations $(X_1, Y_1)$ and $(X_2, Y_2)$ satisfying $X_1 < X_2$. For such pairs of observations, the pairwise slope $(Y_2 - Y_1)/(X_2 - X_1)$ is always defined. If neither $X$ nor $Y$ is subject to left or right censorship, the equality (1) becomes

$$1 - 2q = D(Y - \beta X | X) = \Pr(Y_1 - \beta X_1 < Y_2 - \beta X_2) - \Pr(Y_1 - \beta X_1 > Y_2 - \beta X_2) \quad (2)$$

Therefore, a 0.5th percentile (or median) slope has the expected property that a pairwise slope is equally likely to be above or below it. If in addition the distributions of $X$ and $Y$ are limited to finite sets of discrete values, the distribution of pairwise slopes will be bounded, and a 0th percentile slope will be any number below all possible pairwise slopes, and a 100th percentile slope will be any number above all possible pairwise slopes.

We aim to include a value $\beta$ in a confidence interval for a 100$q$th percentile slope if and only if the sample $\hat{\theta}(Y - \beta X, X)$ is compatible with a population $\theta(Y - \beta X, X)$ equal to $1 - 2q$. The methods of Newson (2006b), used by the program somersd, typically use a monotonically increasing transformation $\zeta(\cdot)$, which may be normalizing and/or variance stabilizing when applied to $\hat{\theta}(Y - \beta X, X)$. We define

$$\zeta^*(\beta) = \zeta(\hat{\theta}(Y - \beta X, X)) \quad (3)$$

$\zeta^*(\beta)$ is a randomly variable function of $\beta$, with a population standard error $\text{SE}\{\zeta^*(\beta)\}$, estimated consistently by a corresponding sample standard error $\hat{\text{SE}}\{\zeta^*(\beta)\}$, whose formula is one of those described in Newson (2006b). We will assume that, if $\theta(Y - X\beta, X) = 1 - 2q$, the pivotal quantity

$$\{\zeta^*(\beta) - \zeta(1 - 2q)\} / \text{SE}\{\zeta^*(\beta)\} \quad (4)$$
has a standard normal distribution. In general, the sample $\zeta^*(\beta)$ is a monotonically nonincreasing step function of $\beta$, bounded above by $\zeta(-1)$ and below by $\zeta(1)$, either of which may be infinite, depending on the choice of transformation $\zeta(\cdot)$.

Figure 1 illustrates an example of a function $\zeta^*(\beta)$ from the auto data. Here the observations are car models, the $Y$ variable is trunk (trunk space in cubic feet), the $X$ variable is foreign (a binary variable indicating non-U.S. origin), the transformation is the hyperbolic arctangent or Fisher’s $z$ (as recommended by Edwardes [1995]), and a slope $\beta$ is a difference (expressed in cubic feet) between cars made by non-U.S. and U.S. companies. The function $\zeta^*(\beta)$ is plotted against the differences $\beta$ over the range of differences for which the absolute value of $\zeta^*(\beta)$ is finite. (As there are no differences between non-U.S. and U.S. cars more than 9 ft$^3$ or less than $-18$ ft$^3$, the value of $\zeta^*(\beta)$ is $-\infty$ for $\beta > 9$ and $+\infty$ for $\beta < -18$. ) This plot was made using the program cendif, which is restricted to binary $X$ variables, and calculates the full set of differences in the $Y$ variable between observations in the two groups. The square data points give values of $\zeta^*(\beta)$ for differences $\beta$ actually observed in the auto data, and the solid line gives values of $\zeta^*(\beta)$ for values of $\beta$ between these observed values. The sample $\zeta^*(\beta)$ is a monotonically nonincreasing step function of $\beta$, which is discontinuous at the observed differences and constant within the open intervals between consecutive observed differences. This outcome implies that a unique exact solution for (1) does not usually exist, as there is usually either no exact solution or an interval of exact solutions between two consecutive observed differences. In a finite sample, this will be true for observed slopes in general, whether or not the $X$ variable is binary.

(Continued on next page)
If we knew the value of \( \text{SE}\{\hat{\zeta}^*(\beta)\} \), then a 100\((1 - \alpha)\)% confidence interval for a 100\(q\)th percentile difference might be the interval of values \( \beta \) for which

\[
\zeta(1 - 2q) - z_\alpha \text{SE}\{\hat{\zeta}^*(\beta)\} \leq \zeta^*(\beta) \leq \zeta(1 - 2q) + z_\alpha \text{SE}\{\hat{\zeta}^*(\beta)\}
\]

where \( z_\alpha \) is the 100\((1 - (1/2)\alpha)\)th percentile of the standard normal distribution. To construct such a confidence interval, we proceed as follows. Given a value \( \zeta \) in the range of \( \zeta(\cdot) \), we define

\[
B_L(\zeta) = \sup \{ \beta : \zeta^*(\beta) > \zeta \}
\]

\[
B_R(\zeta) = \inf \{ \beta : \zeta^*(\beta) < \zeta \}
\]

\[
B_C(\zeta) = \begin{cases} 
\text{Undefined,} & \text{if } B_L(\zeta) = -\infty \text{ and } B_R(\zeta) = \infty, \\
B_L(\zeta), & \text{if } B_L(\zeta) > -\infty \text{ and } B_R(\zeta) = \infty, \\
B_R(\zeta), & \text{if } B_R(\zeta) < +\infty \text{ and } B_L(\zeta) = -\infty, \\
\{ B_L(\zeta) + B_R(\zeta) \}/2, & \text{otherwise.}
\end{cases}
\]

By convention, the supremum (or infimum) of a set unbounded to the right (or left) are defined as \(+\infty\) (or \( -\infty \)), respectively, and the supremum and infimum for an empty set are \(-\infty\) and \(+\infty\), respectively. Clearly, \( B_L(\zeta) \leq B_C(\zeta) \leq B_R(\zeta) \), and the values of \( B_L(\zeta) \) and \( B_R(\zeta) \) (if finite), can be either the same observed slope or two successive observed slopes. The confidence interval for a 100\(q\)th percentile slope is centered on the sample 100\(q\)th percentile slope, defined as

\[
\hat{\xi}_q = B_C \{ \zeta(1 - 2q) \}
\]
The lower and upper confidence limits for a qth percentile slope are, respectively,
\[ \hat{\xi}_{q}^{(\text{min})} = B_{L} \left[ \xi (1 - 2q) - z_{\alpha} \frac{\text{SE}}{\xi} \{ \hat{\xi}_{q}^{*} \} \right], \quad \hat{\xi}_{q}^{(\text{max})} = B_{R} \left[ \xi (1 - 2q) + z_{\alpha} \frac{\text{SE}}{\xi} \{ \hat{\xi}_{q}^{*} \} \right] \] (8)

If `tdist` is specified, `censlope` uses the t distribution with \( \nu = N - 1 \) degrees of freedom if there are \( N \) unclustered observations or with \( \nu = N_{\text{clust}} - 1 \) degrees of freedom if there are \( N_{\text{clust}} \) clusters, instead of the normal distribution, and therefore \( t_{\nu, \alpha} \) replaces \( z_{\alpha} \) in (8). The upper and lower confidence limits may occasionally be infinite for extreme percentiles and/or very small sample numbers. `censlope` codes these infinite limits as plus or minus the Stata `creturn` value `c(maxdouble)`, which is the system maximum double-precision value (see [P] `creturn`).

Figure 1 illustrates these formulas for the Y variable `trunk` and the X variable `foreign` in the `auto` data. The median difference in trunk capacity \( \hat{\xi}_{0.5} \) and its lower and upper 95% confidence limits are shown as reference lines on the horizontal axis. The estimated median difference in trunk space between non-U.S. and U.S. cars is \( -3 \) ft\(^3\), with 95% confidence limits from \(-5\) to \(-1\) ft\(^3\). The reference lines on the vertical axis are the optimum, minimum, and maximum values of \( \zeta^{*}(\beta) \) required for \( \beta \) to be in the confidence interval.

`censlope` inherits all the options of `somersd`, so \( \theta(X, Y - \beta X) \) in (1) can stand for any of the generalized versions of Somers’ D and Kendall’s \( \tau_{a} \) described in Newson (2006b). We can therefore estimate generalized percentile slopes or differences, defined in terms of generalized Somers’ D or Kendall’s \( \tau_{a} \) parameters. For instance, we can use the `wstrata()` option to estimate median slopes and differences restricted to comparisons within strata defined by a confounding variable, or we might use the option `funtype(wcluster)` to estimate within-cluster median differences and slopes. In the terminology of Serfling (1980), the Theil–Sen percentile slope is an \( M \) estimate if `funtype(wcluster)` is specified, a hybrid between an \( M \) estimate and a \( U \) statistic if `funtype(bcluster)` is specified, and a hybrid between an \( M \) estimate and a \( V \) statistic if `funtype(vonmises)` is specified.

### 3.1 Numerical evaluation of \( B_{L}(\zeta) \) and \( B_{R}(\zeta) \)

We can see by (6), (7), and (8) that the key to calculating confidence intervals for percentile slopes is calculating \( B_{L}(\zeta) \) and \( B_{R}(\zeta) \) for a given \( \zeta \). Traditionally, this task has been done by calculating every possible pairwise slope \((Y_i - Y_j)/(X_i - X_j)\) for each pair of observations in the sample to make a dataset of all pairwise slopes and by using this dataset to find the median and other percentile slopes. This approach requires an amount of computational time and data storage space proportional to \( N^2 \), where \( N \) is the number of observations. For this reason, confidence intervals for median slopes have traditionally been calculated only for small samples, as have confidence intervals for other rank statistics, such as Somers’ D and Kendall’s \( \tau_{a} \), which are also commonly calculated by comparing all \((X, Y)\) pairs. See Sprent and Smeeton (2001) for some examples using traditional methods.
Comparing all \((X, Y)\) pairs is not necessary. \texttt{somersd} uses the algorithm of Newson (2006a), which calculates Somers’ \(D\), Kendall’s \(\tau_a\), and their jackknife variances in a time asymptotically proportional to \(N \log N\), using a search tree to avoid having to compare every \((X, Y)\) pair. We can therefore use \texttt{somersd} to calculate \(\zeta^*(\beta)\) for any \(\beta\) in a time proportional to \(N \log N\). \texttt{censlope} uses versions of some of the iterative numerical methods of chapter 9 of Press et al. (1992), modified for step functions, to evaluate \(B_L(\zeta)\) and \(B_R(\zeta)\), for a given \(\zeta\). This method is done by defining the object function \(\omega(\beta) = \zeta^*(\beta) - \zeta\) and attempting to find a solution in \(\beta\) to the equation

\[ 0 = \omega(\beta) = \zeta^*(\beta) - \zeta \]  

(9)

using \texttt{somersd} to calculate \(\omega(\beta)\). This calculation requires a computational time of order \(N_{\text{eval}} N \log N\), where \(N_{\text{eval}}\) is the number of evaluations of the object function in the iteration sequence. For very large datasets \((N > 1,000)\), this approach will typically take less time than a quadratic algorithm that compares all \((X, Y)\) pairs. However, in small datasets, such as the \texttt{auto} data, \texttt{cendif} typically takes much less time to calculate a Hodges–Lehmann median difference by using its quadratic algorithm than \texttt{censlope} takes by using one of its iterative algorithms to do the same. This finding is not surprising. The performance study of Newson (2006a) seems to imply that, if there are fewer than 100 observations, then the execution time of \texttt{somersd} is dominated by “constant” terms not dependent on sample size, whether \texttt{somersd} is using a quadratic algorithm or a search tree algorithm. Therefore, we would expect the computational time for an iteration sequence, involving \(N_{\text{eval}}\) calls to \texttt{somersd}, to have a component proportional to \(N_{\text{eval}}\), which will dominate execution time if the sample size is small and the number of iterations is large.

The algorithms used by \texttt{censlope} are implemented in the Mata language and use versions of standard bracket convergence methods for finding roots, modified for step functions. To solve an equation of the form (9), we would normally start with two \(\beta\) values, \(\beta_0\) and \(\beta_1\), whose corresponding respective \(\omega\) values, \(\omega_0\) and \(\omega_1\), bracket zero, meaning that \(\omega_0 \omega_1 < 0\) (because the two \(\omega\) values have opposite signs). If \(\omega(\cdot)\) is continuous, then by the intermediate-value theorem, there will be a solution to (9) between \(\beta_0\) and \(\beta_1\), and this solution will be unique if \(\omega(\cdot)\) is strictly monotonic. However, \(\omega(\cdot)\) is not continuous but is a nonincreasing step function similar to figure 1. Therefore, instead of expecting to find a unique solution to (9), we try to find a supremum (or infimum) of the set of \(\beta\) values with positive (or negative) values of the object function. In this case, the two \(\omega\) values are said to bracket zero if and only if

\[ \text{sign}(\omega_1) \neq 0 \quad \text{and} \quad \text{sign}(\omega_1) \neq \text{sign}(\omega_0) \]  

(10)

\(\omega_1\) is a strict bracket, which must not be zero, whereas \(\omega_0\) is a partial bracket, which may either be zero or have the opposite sign of \(\omega_1\). During each iteration, we compute a new \(\beta\) value \(\beta_{\text{new}}\), between \(\beta_0\) and \(\beta_1\), with a corresponding \(\omega\) value \(\omega_{\text{new}} = \omega(\beta_{\text{new}})\). In the next iteration, the pair \((\beta_{\text{new}}, \omega_{\text{new}})\) will replace \((\beta_1, \omega_1)\) if \(\text{sign}(\omega_{\text{new}}) = \text{sign}(\omega_1)\) and will replace \((\beta_0, \omega_0)\) otherwise. Iterations proceed until \(\beta_0\) and \(\beta_1\) have a relative difference no more than the value of the \texttt{tolerance()} option. Once this threshold has been reached, we can use either of the \(\beta\) values to estimate \(B_L(\zeta)\) or \(B_R(\zeta)\) (depending on whether we initialized \(\beta_1 < \beta_0\) or \(\beta_0 < \beta_1\)).
The numerical methods specified by the `technique()` option differ in how they calculate \( \beta_{new} \). \texttt{bisection} does this using the simple bisection formula, \( \beta_{new} = (\beta_0 + \beta_1)/2 \). \texttt{regula} uses simple bisection if \( \omega_0 = 0 \) and uses the regula falsi (or false position) method otherwise. \texttt{ridders} uses simple bisection if \( \omega_0 = 0 \) and uses the method of Ridders (1979) otherwise. The simple bisection method is guaranteed to converge slowly, whereas the modified regula falsi and Ridders methods will be faster if the object function \( \omega(\cdot) \) is nearly continuous but may be much slower if \( \omega(\cdot) \) is very discrete. The user may specify a combination of methods, such as starting with the regula falsi or Ridders method for earlier iterations (when the object function is nearly continuous over a long interval) and moving to the bisection method later (when the object function is highly discrete over a short interval).

For each percentage \( 100q \), \texttt{censlope} attempts to evaluate \( B_L\{\zeta(1-2q)\} \) and \( B_R\{\zeta(1-2q)\} \), attempts to evaluate the percentile estimate \( \hat{\xi}_q \), and then (if this evaluation is successful) evaluates the two confidence limits. This pathway implies four sequences of iterations to evaluate: the “left estimate”, the “right estimate”, and the two confidence limits, respectively. Typically, using the default tolerance of \( \text{1e-6} \) and the “slow but sure” bisection method implies four sets of around 20 iterations. Together with the initialization of the brackets, this method implies many (80–100) calls to \texttt{somersd}. However, that number is usually fewer than 100 evaluations per percentile, implying less work than (say) bootstrapping Somers’ \( D \), which would typically involve at least 1,000 evaluations. On the other hand, if the sample size is large, this method would probably be unthinkable for practical statisticians without the algorithm of Newson (2006a).

### 3.2 Comparisons with existing methods

Sen (1968) developed a confidence interval formula for \( \hat{\xi}_q \) in the special case where \( q = 0.5 \), \( \theta(Y, X) = \tau(Y, X) \), and \( \zeta(\theta) = \theta \), using methods similar to those in this article. In this special case, (1) becomes simply \( \tau(Y - \beta X, X) = 0 \). The main difference from the present method was in how the distribution of \( \zeta^*(\beta) \) was calculated. Sen assumed that the variables \( X \) and \( Y - \beta X \) were not only “Kendall uncorrelated” but also statistically independent. For small sample sizes \( (N \leq 10) \), the confidence interval was calculated using tables of the exact distribution of the sample Kendall’s \( \tau_a \), based on that assumption. For larger sample sizes, the population standard error \( \text{SE}[\zeta^*(\beta)] \) was calculated from the marginal sample distribution of \( X \), using the same assumption. (See Kendall and Gibbons [1990] for tables of the exact distribution for small sample sizes and for a demonstration that the Central Limit Theorem works well at sample sizes as small as 8 for the sample Kendall’s \( \tau_a \) under the null hypothesis of independence.) The assumption of independence between the predictor variable \( X \) and the “residuals” \( Y - \beta X \) implies that the conditional population distributions of \( Y \), given each value of \( X \), are different only in location and may not differ in the conditional variance, or indeed in any other conditional moment about the mean. The original Sen method therefore does not use the assumption of normality but does use the assumption of homoskedasticity, which typically causes more problems when it is wrong.
Lehmann (1963) derived a confidence interval for the Hodges–Lehmann median difference, which is the Theil–Sen slope for binary X variables, based on the same assumption of independence. This method was made popular by Conover (1980), Campbell and Gardner (1988), and Altman et al. (2000) and is available in unofficial Stata, using Duolao Wang’s npshift routine (Wang 1999) or Patrick Royston’s cid routine, downloadable from SSC. The method is essentially a special case of the Sen (1968) method and is presumably subject to the same cautions.

The method used by censlope and cendif, by contrast, can estimate percentile differences other than the median difference. Even for a median difference, the predictor variable X and the “residuals” Y – βX are assumed to be only “Kendall uncorrelated” and not necessarily independent. The population standard error $\hat{SE}[\zeta^*(\beta)]$ is estimated using the sample standard error $\hat{SE}[\zeta^*(\beta)]$, which is calculated using an infinitesimal jackknife method described in Newson (2006b). This method is robust to heteroskedasticity, probably at the price of being less robust to extremely small sample sizes than the traditional methods. Therefore, the method of censlope can be compared with the original Sen method as Huber confidence intervals can be compared with maximum likelihood or quasilikelihood confidence intervals, and the method of cendif can be compared with the Lehmann method as the unequal-variance t test can be compared with the equal-variance t test. Lehmann’s method, like the equal-variance t test, assumes that you can use data from the larger of two samples to estimate the population variability of the smaller sample. The censlope method, like the unequal-variance t test, assumes that you can use data from the smaller of the two samples to estimate the population variability of the smaller sample. At present, if the tdist option is specified for censlope or cendif, the number of degrees of freedom is set to one less than the sum of the two sample numbers. This method is in contrast to the unequal-variance t test, which typically uses a more complicated formula (Satterthwaite 1946) and is usually less generous with degrees of freedom if the smaller sample size is very small.

The issue of heteroskedasticity, as it affects the t test, is discussed in Moser, Stevens, and Watts (1989) and in Moser and Stevens (1992), which explored the issue, using exact analytical formulas to compare the equal-variance t test with the Satterthwaite unequal-variance t test. Their conclusion (as I understand it) appears to be that we should view the equal-variance t test as a special method for use only when we “know” that the subpopulation variances are equal, rather than following the more traditional practice of viewing the unequal-variance t test as a special method for use only when we “know” that the subpopulation variances are unequal. I have carried out some unpublished simulations, comparing cendif to the Lehmann method and to the two t tests. These simulations, some of which are briefly described in Newson (2000b) and in Newson (2002), seem to point to a similar recommendation regarding the two types of rank-based methods for median differences. However, more work is probably required on this issue.

Another method of defining heteroskedasticity-consistent confidence intervals for the Theil–Sen median slope is the percentile bootstrap, recommended by Wilcox (1998). Bootstrapping censlope or cendif may be an option, at least for small samples, where the computational cost of evaluating one sample median slope or difference, using a
quadratic or iterative method, is low enough to allow us to evaluate many subsample median slopes or differences. \texttt{censlope} adds the options of estimating clustered and/or stratified median slopes and differences, as well as the option of nonbootstrap confidence intervals for large samples. The infinitesimal jackknife method, used by \texttt{somersd}, is usually considered to be an inferior substitute for the bootstrap method applied to the same parameter. However, in this case, the infinitesimal jackknife standard error calculated by \texttt{censlope} is not for the median slope itself but for another parameter (Somers’ $D$ or Kendall’s $\tau_a$), for which the central limit theorem works fast, especially under the null hypothesis (Kendall and Gibbons 1990). This setup might limit the advantage of the bootstrap over the infinitesimal jackknife. On the other hand, a possible future compromise might be to modify \texttt{censlope} to allow it to bootstrap Somers’ $D$ or Kendall’s $\tau_a$ and thereby to substitute bootstrap-based formulas for formulas (5) and (8) when calculating confidence intervals for the percentile slope itself. Whether we use the bootstrap or the infinitesimal jackknife, it is probably a good idea, if the sample size is large, to calculate the Theil–Sen median slope by using a nonquadratic algorithm, which does not require calculation of all the individual pairwise slopes.

4 Examples

These examples introduce some of the capabilities of \texttt{censlope}. There are more examples in the online help for \texttt{censlope} and in \texttt{censlope.pdf}, which is distributed with the \texttt{somersd} package.

4.1 Weight per inch in the auto data

In the \texttt{auto} data, we can use \texttt{censlope} to estimate the median slope of \texttt{weight} (in U.S. pounds) with respect to \texttt{length} (in U.S. inches) as follows:

\begin{verbatim}
  (1978 Automobile Data)
. censlope weight length, tdist
Outcome variable: weight
  Somers' D with variable: length
  Transformation: Untransformed
  Valid observations: 74
  Degrees of freedom: 73
  Symmetric 95% CI

       Jackknife                     [95% Conf. Interval]
   length          Coef.    Std. Err.    t    P>|t|         Lower          Upper
weight    .8286359     .0275321   30.10    0.000    .7737644    .8835073

95% CI(s) for percentile slope(s)
Percent Pctl_Slope Minimum Maximum
  50    32.745114    30.508468    35.185195
\end{verbatim}
Confidence intervals for rank statistics

The untransformed Somers’ $D$ of weight with respect to length is 0.83, with a confidence interval from 0.77 to 0.88, indicating that, in the population from which these cars were sampled, a longer car is 77%–88% more likely to be heavier than a shorter car than to be lighter than a shorter car. Each additional inch of length typically adds 30.51–35.19 pounds of weight.

If we use the $z$ transform for Somers’ $D$, the results are as follows:

```
. censlope weight length, tdist transf(z)
Outcome variable: weight
Somers’ D with variable: length
Transformation: Fisher’s z
Valid observations: 74
Degrees of freedom: 73
Symmetric 95% CI for transformed Somers’ D

| length | Coef. | Std. Err. | t     | P>|t| | [95% Conf. Interval] |
|--------|-------|-----------|-------|------|---------------------|
| weight | 1.183767 | .0878602 | 13.47 | 0.000 | 1.008662 1.358873 |

Asymmetric 95% CI for untransformed Somers’ D

<table>
<thead>
<tr>
<th>Somers_D</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>weight</td>
<td>.82863585</td>
<td>.76520811</td>
</tr>
</tbody>
</table>

95% CI(s) for percentile slope(s)

<table>
<thead>
<tr>
<th>Percent Pctl_Slope</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>32.745093</td>
<td>30.571414</td>
</tr>
</tbody>
</table>
```

This time, Somers’ $D$ is 0.77–0.88, implying (again) that longer cars are 77%–88% more likely to be heavier than shorter cars than to be lighter than shorter cars. The typical increase in weight per additional inch of length is 30.57–35.12 pounds per inch, which is similar to the previous confidence interval.

Transformations such as Fisher’s $z$ are more likely to be important in estimating percentile slopes other than the median. We can ask for the 25th and 75th percentiles as well, using the `centile()` option:

```
. censlope weight length, tdist transf(z) centile(25(25)75)
Outcome variable: weight
Somers’ D with variable: length
Transformation: Fisher’s z
Valid observations: 74
Degrees of freedom: 73
Symmetric 95% CI for transformed Somers’ D

| length | Coef. | Std. Err. | t     | P>|t| | [95% Conf. Interval] |
|--------|-------|-----------|-------|------|---------------------|
| weight | 1.183767 | .0878602 | 13.47 | 0.000 | 1.008662 1.358873 |
```
Asymmetric 95% CI for untransformed Somers’ D

<table>
<thead>
<tr>
<th>weight</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.76520811</td>
<td>0.87613131</td>
<td></td>
</tr>
</tbody>
</table>

95% CI(s) for percentile slope(s)

<table>
<thead>
<tr>
<th>Percent</th>
<th>Pctl_Slope</th>
<th>Minimum</th>
<th>Maximum</th>
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</thead>
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<tr>
<td>25</td>
<td>24.102562</td>
<td>19.999992</td>
<td>27.06897</td>
</tr>
<tr>
<td>50</td>
<td>32.745093</td>
<td>30.571414</td>
<td>35.121969</td>
</tr>
<tr>
<td>75</td>
<td>41.818174</td>
<td>38.620683</td>
<td>46.200022</td>
</tr>
</tbody>
</table>

We see that the 25th percentile slope is 20.00–27.07 pounds per inch and that the 75th percentile slope is 38.62–46.20 pounds per inch.

We can also produce plots of observed and fitted values, using the `ystargenerate()` option of `censlope`. After executing `censlope`, we use `egen` to calculate the median of the variable `resid`, generated by the `ystargenerate()` option, which stores the “residuals” \( Y - \beta X \), where \( Y \) is weight, \( X \) is length, and \( \beta \) is the median slope. This median is stored in a new variable, named `intercept`. Then we generate the fitted values of `weight` in a new variable `wthat`, calculated by subtracting \( Y - \beta X \) from \( Y \) to obtain \( \beta X \) and then adding `intercept`. These fitted values are plotted as a line against `length`, and the observed weight values are superimposed to create the graph of figure 2.
4.2 Prenatal paracetamol and immunoglobulin E

The Avon Longitudinal Study of Pregnancy and Childhood (ALSPAC) is a birth cohort study based at the University of Bristol, Bristol, UK. For more information, see the study web site at http://www.alspac.bris.ac.uk. As part of the study, the mothers of 12,127 children were asked whether they ever used paracetamol (acetaminophen) in weeks 20–32 of pregnancy. At 7 years of age, total immunoglobulin E (IgE) was measured in the blood of 4,848 of these children. IgE level is viewed as a measure of allergic tendency and is raised in individuals suffering from allergic diseases such as asthma. Shaheen et al. (2005) reported that in ALSPAC the children of paracetamol users typically had higher IgE levels than those of children of paracetamol nonusers, based on estimates of geometric mean ratios.

The distribution of total IgE, expressed in kilounits per liter (kU/L), in the 4,848 children with data on IgE and on maternal paracetamol use in late pregnancy, is given in figure 3. The distribution is nonnormal and has a long tail of extremely high values. A total of 2,051 of these children had mothers who reported using paracetamol in late pregnancy, and the remaining 2,797 had mothers who reported not using paracetamol.
Figure 3: Distribution of blood IgE in 4,848 children in ALSPAC

We used \texttt{censlope} to compare the IgE levels in children exposed and unexposed to maternal paracetamol use in late pregnancy, using Somers’ \(D\) and the Hodges–Lehmann median IgE ratio. Given a randomly sampled paracetamol-exposed child and a randomly sampled unexposed child, Somers’ \(D\) is the difference between the probability that the exposed child has the higher IgE level and the probability that the unexposed child has the higher IgE level. The Hodges–Lehmann median ratio is the median ratio between IgE levels in two such randomly sampled children and is defined as the exponential of the Hodges–Lehmann median difference between the logged IgE values in the two groups and estimated using the \texttt{eform} option of \texttt{censlope}. The results are as follows:

```
censlope lnigetot para32g, transf(z) eform
Outcome variable: lnigetot
Somers’ \(D\) with variable: para32g
Transformation: Fisher’s \(z\)
Valid observations: 4848
Symmetric 95% CI for transformed Somers’ \(D\)

| para32g | Coef. | Std. Err. | z    | P>|z| | [95% Conf. Interval] |
|---------|-------|-----------|------|------|---------------------|
| lnigetot | .0533954 | .0168421 | 3.17 | 0.002 | [.0203856 .0864053] |
```

Asymmetric 95% CI for untransformed Somers’ \(D\)

```
Somers_D | Minimum | Maximum
lnigetot | .05334475 | .02038276 | .0861909
```

95% CI(s) for percentile ratio(s)

```
Percent Pctl_Ratio | Minimum | Maximum
50 | 1.172549 | 1.0616111 | 1.2944986
```
Confidence intervals for rank statistics

Here lnigetot is the natural log of total IgE, and para32g is a binary variable, indicating paracetamol exposure during weeks 20–32 of gestation. From the asymmetric confidence interval for the untransformed Somers’ D, we see that, if we choose an exposed child and an unexposed child at random, the exposed child is 2.0%–8.6% more likely than the unexposed child to have the higher IgE level. From the confidence interval for the 50th percentile (or median) ratio, we can see that the median ratio is 1.06–1.29, implying that the exposed child typically has 6%–29% more IgE than the unexposed child.

However, these are only crude, unadjusted estimates, and the effects that they represent could be due to potential confounding variables. To produce confounder-adjusted estimates, we used a propensity score, as defined in Rosenbaum and Rubin (1983) and Rosenbaum (2003). We defined this score by fitting a logistic regression model with para32g as the outcome to data from the 12,127 children with paracetamol data. The predictors in this model were the following confounders: gender, maternal age, prenatal tobacco exposure, mother’s education, housing tenure, parity, maternal anxiety, maternal ethnic origin, multiple pregnancy, birth weight, gestational age at birth, head circumference, use of antibiotics in pregnancy, alcohol intake in pregnancy, maternal disease and infection history, younger siblings, presence of pets, breast feeding, day care, dampness problems, passive smoking exposure after birth, and obesity index at 7 years. (Not all these confounders could have had a causal effect on prenatal paracetamol exposure, but they could all be indirect indicators of prenatal proneness to paracetamol exposure.) The propensity score was defined as the predicted log odds of paracetamol exposure from this regression model. Using the xtile command (see [D] pctile), we defined 32 paracetamol propensity groups, with approximately equal numbers.

somersd, and therefore censlope, has a wstrata() option, allowing stratified versions of Somers’ D and median slopes, restricted to comparisons between pairs of observations in the same stratum. We measured the confounder-adjusted paracetamol effect by using censlope with the option wstrata(pg_para32g), where pg_para32g is a discrete variable indicating which of the 32 paracetamol-propensity groups a child belongs to, based on that child’s confounder values. The results were as follows:

```
censlope lnigetot para32g, transf(z) eform wstrata(pg_para32g)
Outcome variable: lnigetot
Somers’ D with variable: para32g
Transformation: Fisher’s z
Within strata defined by: pg_para32g
Valid observations: 4848

Symmetric 95% CI for transformed Somers’ D

| para32g | Coef. | Std. Err. | z    | P>|z| | [95% Conf. Interval] |
|---------|-------|-----------|------|-----|---------------------|
| lnigetot| .0416191 | .018089 | 2.30 | .021 | .0061653 to .0770729 |
```
Asymmetric 95% CI for untransformed Somers’ D

<table>
<thead>
<tr>
<th>Somers_D</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>lnigetot</td>
<td>.04159508</td>
<td>.00616518</td>
</tr>
<tr>
<td></td>
<td>.07692067</td>
<td></td>
</tr>
</tbody>
</table>

95% CI(s) for percentile ratio(s)

<table>
<thead>
<tr>
<th>Percent</th>
<th>Pctl_Ratio</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>1.1256541</td>
<td>1.0165742</td>
<td>1.2556066</td>
</tr>
</tbody>
</table>

This time, the adjusted Somers’ D is between 0.006 and 0.077, and the adjusted Hodges–Lehmann median ratio is between 1.017 and 1.256. Therefore, if we sample a random-exposed child and a random-unexposed child from the same propensity stratum, it is 0.6%–7.7% more likely that the exposed child will have the higher IgE level than that the unexposed child will have the higher IgE level, and the exposed child will typically have 1.7%–25.6% more IgE than the unexposed child. Therefore, sampling similarly paracetamol-prone children does not seem to alter the relative exposed–unexposed IgE difference much. These conclusions are (reassuringly) similar to those of Shaheen et al. (2005).

The sample size of 4,848 is much larger than those of most samples conventionally analyzed using rank methods and is in the range at which the computational methods used by censlope begin to have an advantage. The unadjusted analysis presented above typically takes 2 minutes with censlope and 4 minutes with cendif on my system, which is based on a 2.79-GHz Intel Pentium 4 CPU with 0.99 GB of RAM running Windows XP. As sample size increases further, so will the ratio of time and space requirements between cendif (which uses a quadratic-time and quadratic-space algorithm) and censlope.

5 Summary

The censlope module is a major extension to the somersd package, enabling the estimation of generalized Theil–Sen percentile slopes and Hodges–Lehmann percentile differences, corresponding to the generalized Somers’ D and Kendall’s $\tau_a$ parameters introduced in Newson (2006b). All these generalized parameters are estimated with confidence intervals and may be restricted to comparisons within or between clusters and/or strata defined by a confounder or by a propensity score summarizing multiple confounders. The somersd package therefore allows users to do more with rank methods than they were probably accustomed to do, although we may still need regression methods to define a propensity score.

Rank parameters of the Somers’ D family have the advantage of being robust to distributional assumptions. Somers’ D and Kendall’s $\tau_a$ have “democratic” influence functions, based on a principle of “one comparison, one vote”, causing the central limit theorem (usually) to work faster than it would for comparable regression parameters. (See Hampel [1974] and Hampel, Ronchetti, Rousseeuw, and Stahel [1986] for more about influence functions.) This robustness must, to an extent, be purchased at the price of being less robust to small sample numbers. The argument of Fisher (1935) implies that, if we know the distributional family a priori, an estimate for a median slope or dif-
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ference based on maximum likelihood estimators will have a lower asymptotic variance than the corresponding Theil–Sen or Hodges–Lehmann statistics. The contrast in power may be spectacular at tiny sample sizes, when using a $t$ test may reduce the minimum detectable difference from infinity to a finite difference (which is why censlope and cendif can produce infinite confidence limits). At larger sample sizes, there is typically a more modest contrast in power, such as a 5% reduction in the minimum detectable difference, and even this may be conditional on guessing the distributional family right in advance. However, more work is needed (and is in progress) to find out more about the tradeoffs involved.

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7 References


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About the author

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