Estimation of mean health care costs and incremental cost-effectiveness ratios with possibly censored data

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Abstract. In this article, we describe the hcost program for estimating mean health care costs and incremental cost-effectiveness ratios with possibly censored data. hcost estimates the mean survival time and the mean costs, as well as their variances and covariance, for a given time horizon. If the group variable is specified, hcost will report the differences between two groups, as well as the incremental cost-effectiveness ratio and its confidence interval (optional). hcost can estimate the mean costs using two methods corresponding to different types of data: the Bang and Tsiatis (2000, Biometrika 87: 329–343) estimator using only the total costs or the Zhao and Tian (2001, Biometrics 57: 1002–1008) estimator when cost-history data are available. The hcost program can also be used to specify the annual discounting rates for survival time and costs.

Keywords: st0399, hcost, mean costs, censored data, cost history, cost-effectiveness analysis

1 Introduction

Recently, the estimation of health care costs in economic evaluations of new treatments has received a lot of attention. In an environment of extremely high health care costs and limited resources, cost-effectiveness analysis to devise treatment strategies that offer a greater health benefit without imposing large economic burdens on society has become common.

In clinical trials and observational studies, survival time and accumulated health costs frequently are censored because not all patients can be observed until terminal events (for example, death) occur. Censoring poses a unique problem for cost estimation because of the “induced informative censoring” challenge, first noted by Lin et al.
As a result, traditional methods for handling censored survival data, such as the Kaplan–Meier estimator or Cox proportional hazards regression model, are no longer valid for analyzing censored cost data.

Because of the presence of censoring, the marginal distribution of cost may not be identifiable without making some parametric assumptions (Huang 2002). Thus most methods of estimating mean costs focus on restricted medical costs—that is, on costs accumulated within a time limit \( L \), where \( L \) is chosen such that a reasonable number of subjects are still available at that time. Consequently, further cost-effectiveness analysis concentrates on time-restricted costs and effectiveness.

Different methods have been proposed for estimating the time-restricted mean costs. Lin et al. (1997) proposed estimators based on survival-probability weighting using partitioned time intervals; Bang and Tsiatis (2000) proposed several consistent estimators, including the Bang and Tsiatis (BT) estimator and its Bang and Tsiatis partitioned version (BTP) using the inverse-probability weighting technique; and Zhao and Tian (2001) proposed an efficient Zhao and Tian (ZT) estimator for when cost history is available. Later, Zhao et al. (2007) described the conditions under which the BT estimator is equivalent to the Lin T estimator (when cost history is not available) and the condition under which the ZT estimator is equivalent to the BTP and the two estimators Lin A and B proposed by Lin et al. (1997) (when cost history is available). Specifically, the equality occurs when the partition boundaries coincide with those censoring times. The BTP, the Lin T estimator, and the Lin A and B estimators may change with different ways of partitioning, while the BT and ZT estimators do not depend on any type of partitioning.

In this article, we describe how to implement the BT estimator (using the total costs only) and the ZT estimator (when cost history is available) in Stata to estimate mean health care costs with possibly censored data. If the data come from two treatment groups, a cost-effectiveness analysis can also be conducted, which estimates the incremental cost-effectiveness ratio and its confidence interval based on Fieller’s method (Fieller 1954; Zhao and Tian 2001). We also discuss methods that allow discounting for costs and survival times.

2 Estimation of mean costs and mean survival time with censored data

For the \( i \)th individual in the study, \( i = 1, 2, \ldots, n \), we define \( T_i \) as the survival time from study enrollment until the occurrence of some event (for example, death or disease relapse). Because some individuals are often still living when the study ends, these are considered censored observations. The censoring time for the \( i \)th individual is denoted as \( C_i \). We can observe either the survival time or the censoring time, whichever is shorter; that is, we observe the follow-up time, \( X_i = \min(T_i, C_i) \), and the death indicator variable, \( \Delta_i = I(T_i \leq C_i) \). We define \( M_i(t) \) as the accumulated costs for patient \( i \) from time 0 to \( t \). For some applications, we observe only the total costs, \( M_i = M_i(X_i) \). However, in other studies, we may know the entire cost history, \( \{M_i(t), 0 < t < X_i\} \).
Here we briefly describe the methods for estimating the mean costs accumulated over time $L$ with censored data, where $L$ can be any time limit such that a reasonable number of subjects are still being observed at that time. The restricted survival time must be defined as $T^L_i = \min(T_i, L)$. Accordingly, the follow-up time becomes $X^L_i = \min(T^L_i, C_i)$, and the death indicator variable becomes $\Delta^L_i = I(T^L_i \leq C_i)$. We want to estimate the mean costs accumulated within time $L$, $\mu^M = E(M(T^L_i))$. For clarity, we omit the superscript $L$ in subsequent sections of this article.

Bang and Tsiatis (2000) proposed a consistent $BT$ estimator based on the inverse-probability weighting technique.

$$\hat{\mu}_{\text{BT}}^M = \frac{1}{n} \sum_{i=1}^{n} \frac{\Delta_i M_i}{\hat{K}(T_i)}$$

(1)

Here $\hat{K}(T_i)$ is the Kaplan–Meier estimator for the survival function of the censoring time $C$, $K(u) = \Pr(C > u)$. The $BT$ estimator is consistent and asymptotically normally distributed with a variance that can be estimated consistently (Bang and Tsiatis 2000) by

$$\hat{\text{Var}}(\hat{\mu}_{\text{BT}}^M) = \frac{1}{n^2} \sum_{i=1}^{n} \frac{\Delta_i (M_i - \hat{\mu}^M_{\text{BT}})^2}{\hat{K}(T_i)} + \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{\hat{K}(C_i)} \{G_i(M^2) - \hat{G}_i^2(M)\}$$

where

$$G_i(Z) = \frac{1}{nS(C_i)} \sum_{j=1}^{n} \frac{\Delta_j}{K(T_j)} Z_j I(T_j \geq C_i)$$

for any random variable $Z$ and $\hat{S}(u)$ is the Kaplan–Meier estimator for $S(u) = \Pr(T > u)$, which is the survival distribution of survival time $T$ at time $u$.

With cost history available, the $BT$ estimator is not efficient, because it does not use the cost information from censored observations. A more efficient estimator is proposed by Zhao and Tian (2001). The simplified form of the $ZT$ estimator (Pfeifer and Bang 2005) is

$$\hat{\mu}_{\text{ZT}}^M = \frac{1}{n} \sum_{i=1}^{n} \frac{\Delta_i M_i}{\hat{K}(T_i)} + \frac{1}{n} \sum_{i=1}^{n} \frac{(1 - \Delta_i)(M_i(C_i) - \overline{M}(C_i))}{\hat{K}(C_i)}$$

(2)

where $\overline{M}(C_i) = \sum_{j=1}^{n} I(X_j \geq C_i)M_j(C_i)/\sum_{j=1}^{n} I(X_j \geq C_i)$, which is the average of accumulated costs at time $C_i$ of those subjects who are living at $C_i$. 


The ZT estimator is consistent and asymptotically normally distributed with variance that can be estimated consistently (Zhao and Tian 2001) by

$$
\hat{\text{Var}}(\hat{\mu}_{ZT}) = \frac{1}{n^2} \sum_{i=1}^{n} \frac{\Delta_i (M_i - \hat{\mu}_{ZT})^2}{K(T_i)} + \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} \{G_i(M^2) - G_i^2(M)\}
$$

$$
- \frac{2}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} [G_i\{M(C_i)\} - G_i(M)G_i\{C_i\}]
$$

$$
+ \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} \left\{M(C_i)^2 - M(C_i)^2\right\}
$$

where $G_i$ and $M(C_i)$ are defined previously, and

$$
M(C_i)^2 = \frac{\sum_{j=1}^{n} I(X_j \geq C_i) M_j(C_i)^2}{\sum_{j=1}^{n} I(X_j \geq C_i)}
$$

Meanwhile, the mean survival time to time $L$ can be obtained by the area under the survival function; that is, $\hat{\mu}_T = \int_{0}^{L} \hat{S}(u) du$, where $\hat{S}(u)$ is the Kaplan–Meier estimator for $S(u) = \Pr(T > u)$. This estimator can be obtained more conveniently (Satten and Datta 2001; Zhao and Tian 2001) by

$$
\hat{\mu}_T = \frac{1}{n} \sum_{i=1}^{n} \frac{\Delta_i T_i}{K(T_i)}
$$

Following Zhao and Tian (2001), we can estimate its variance consistently by

$$
\hat{\text{Var}}(\hat{\mu}_T) = \frac{1}{n^2} \sum_{i=1}^{n} \frac{\Delta_i (T_i - \hat{\mu}_T)^2}{K(T_i)} + \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} \{G_i(T^2) - G_i^2(T)\}
$$

We can also estimate the covariance of $\hat{\mu}_{BT}$ and $\hat{\mu}_T$ (Zhao and Tian 2001) by

$$
\hat{\text{Cov}}(\hat{\mu}_{BT}, \hat{\mu}_T) = \frac{1}{n^2} \sum_{i=1}^{n} \frac{\Delta_i M_i T_i}{K(T_i)} - \frac{1}{n^3} \sum_{i=1}^{n} \frac{\Delta_i M_i}{K(T_i)} \sum_{i=1}^{n} \frac{\Delta_i T_i}{K(T_i)}
$$

$$
+ \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} \{G_i(TM) - G_i(M)G_i(T)\}
$$
We can estimate the covariance of $\hat{\mu}_{MT}^M$ and $\hat{\mu}^T$ (Zhao and Tian 2001) by

$$\text{Cov}(\hat{\mu}_{MT}^M, \hat{\mu}^T) = \frac{1}{n^2} \sum_{i=1}^{n} \frac{\Delta_i M_i T_i}{K(T_i)} - \frac{1}{n^3} \sum_{i=1}^{n} \frac{\Delta_i M_i}{K(T_i)} \sum_{i=1}^{n} \frac{\Delta_i T_i}{K(T_i)}$$

$$+ \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} \{G_i(T M) - G_i(M)G_i(T)\}$$

$$- \frac{1}{n^2} \sum_{i=1}^{n} \frac{1 - \Delta_i}{K(C_i)^2} [G_i\{T M(C_i)\} - G_i\{M(C_i)\}G_i(T)]$$

### 3 Discounting costs and survival time

In cost-effectiveness analysis, it is customary to discount the costs and survival time for a future time because a dollar and a year of life at present may be more valuable than a dollar and a year of life in the future. A 3% annual discount rate is used frequently in practice, but other discount rates may also be considered (Gold et al. 1996). Assume the annual discount rate is $\beta$. Consider a small time unit $u$, where the discount rate over the interval is $\beta u$. Then the present value for $x_u$ dollars accumulated at some future time $u$ is $x_0 = x_u (1 - \beta u)$ dollars, and the present value for $x_{2u}$ dollars at future time $2u$ is $x_0 = x_{2u} (1 - \beta u)^2$ dollars, and so on. Therefore, the present value for $x_{nu}$ dollars accumulated at future time $nu$ is $x_0 = x_{nu} (1 - \beta u)^n$. If $t = nu$, then the discounted value of $x_t$ dollars at future time $t$ is

$$x_0 = x_t (1 - \beta u)^n = x_t (1 - \beta u)^{\frac{t}{u}} \rightarrow x_t e^{-\beta t} \quad \text{as} \quad u \rightarrow 0$$

If $v(t)$ denotes costs to be incurred at future time $t$ (note that the accumulated cost $M(t)$ is $M(t) = \int_0^t v(u)du$), its discounted cost at the beginning of the study is

$$v_0 = v^d(t) = e^{-\beta t} v(t)$$

In the input data for our hcost program, each cost entry is associated with a start date and a stop date. If the option for discounting cost ($drcost()$) is chosen, the program will discount the accumulated costs using its start date. When the time interval between start and stop dates for this cost entry is shorter than the user-specified $k()$ (days), the accumulated costs are discounted using the start date. However, if the time interval is longer than $k()$, then the interval is divided into $k()$-day subintervals, and the costs at each subinterval are discounted at the start date of this subinterval. The default value for $k()$ is $k(90)$.

A similar strategy can be used to discount years of life in the future at an annual discount rate of $\beta$, as if “life” values were measured in dollars. For one year of life in the future time $t$, its present value is $e^{-\beta t}$. Therefore, the current value (discounted value) of survival time $T$ is

$$T^d = \int_0^T e^{-\beta s} ds = \frac{1}{\beta} (1 - e^{-\beta T})$$
Incremental cost-effectiveness ratio

If a new treatment has a greater health benefit but lower costs when compared with its competitor, the new strategy is undoubtedly preferred. However, if a program has higher costs but a greater health benefit than its competitor, a decision must be made about which of the two programs to adopt. The incremental cost-effectiveness ratio (ICER) is a useful measure under these circumstances.

The ICER is defined as the additional costs one must pay for saving one additional year of life. Mathematically, it can be expressed as
\[ \gamma = \frac{\mu_2^M - \mu_1^M}{\mu_2^T - \mu_1^T}. \]

It can be estimated by plugging in the ZT estimator (2) or BT estimator (1) for the mean cost \( \hat{\mu}_k^M \) as well as the estimator for mean survival time (3), \( \hat{\mu}_k^T \), for each group \( k, k = 1, 2 \).

Bootstrap methods (Efron and Tibshirani 1986, 1993; Hwang 1995; Mushlin et al. 1998; Jiang, Wu, and Williams 2000; O’Brien and Briggs 2002; Jiang and Zhou 2004) are commonly used to construct confidence intervals (CIs) for the ICER. Although many researchers believe that the bootstrap method provides better coverage, Hwang (1995) and Jiang, Wu, and Williams (2000) show that the bootstrap method is only first-order accurate, similar to the Fieller method (Fieller 1954). Therefore, the Fieller method, if used correctly, can be a reliable and computationally efficient way to obtain CIs for the ICER.

Zhao and Tian (2001) used Fieller’s Theorem to obtain CIs for the ICER. Because asymptotically \( x = \tilde{\mu}_2^M - \tilde{\mu}_1^M \) and \( y = \tilde{\mu}_2^T - \tilde{\mu}_1^T \) are bivariate normally distributed, the 100(1 - 2\( \alpha \))% confidence limits for the ICER \( \gamma \) are

\[ xy - z^2_\alpha s_{xy} \pm \frac{\{(xy - z^2_\alpha s_{xy})^2 - (x^2 - z^2_\alpha s_{xx})(y^2 - z^2_\alpha s_{yy})\}}{y^2 - z^2_\alpha s_{yy}} \frac{1}{2} \]

where \( s_{xx}, s_{yy}, \) and \( s_{xy} \) are respectively the variances of \( x \) and \( y \) and the covariance of \( x \) and \( y \), and \( z_\alpha \) is the cutoff point with tail area \( \alpha \) of the standard normal distribution. Because the two treatment groups are independent, \( s_{xx}, s_{yy}, \) and \( s_{xy} \) can be estimated by the summation of all the estimations over the two groups \( k = 1, 2 \), respectively.

Note that the cost-effectiveness analysis is normally performed when one treatment group is simultaneously more effective and more costly than the other group. The formula (4) can be used to obtain a finite CI only if there is a statistically significant difference between the mean survival times of the two treatment groups—that is, if the denominator of (4) is positive. If the denominator of (4) is negative, indicating that the difference between the effects of two treatments is not statistically significant, the CI for the ICER is exclusive and thus infinite; that is, the CI is formed by \((-\infty, \text{CL}_L) \cup (\text{CL}_U, +\infty)\), where \( \text{CL}_L \) and \( \text{CL}_U \) are the lower and upper limits obtained from (4). Although this infinite CI is correct and reflects our knowledge about the true ICER, its interpretation can be difficult (Stinnett and Mullahy 1998; Briggs, Mooney, and Wonderling 1999; Wang and Zhao 2008). Alternative methods, such as the cost-effectiveness acceptability curve, may be considered in this situation (Fenwick et al. 2006; Löthgren and Zethraeus 2000). For this reason, we do not rou-
tinely perform a cost-effectiveness analysis, but we do provide an option for calculating the ICER and its CI.

5 The hcost command

5.1 Syntax

```
hcost idvar costvar [if] [in], start(varname) stop(varname) l(#) 
[ group(varname) drsurv(#) drcost(#) icer(#) k(#) level(#) 
  method(#) ]
```

The `hcost` command estimates mean costs within a time horizon \( L \) with possibly censored data. The data must be processed by the `stset` command (see [ST] stset) before using `hcost`.

5.2 Options

`start(varname)` specifies the variable that contains the start time of accumulated costs (in days). `start()` is required unless `method()` is 0 and `drcost()` is 0.

`stop(varname)` specifies the variable that contains the end time of accumulated costs (in days). `stop()` is required unless `method()` is 0 and `drcost()` is 0.

`l(#)` specifies the time limit \( L \) (in days) for calculating mean costs and mean survival time. `l()` is required.

`group(varname)` specifies the variable that contains the treatment group information if there is more than one group. If there is only one group, this option need not be specified. There cannot be more than two groups.

`drsurv(#)` specifies the annual discount rate for survival time. The value must be between 0 and 1. The default is `drsurv(0)`, that is, survival time is already discounted in the data.

`drcost(#)` specifies the annual discount rate for costs. The value must be between 0 and 1. The default is `drcost(0)`, that is, cost is already discounted in the data. Each cost entry is discounted according to its start date. If the time interval for the cost entry is too big, it will be divided into \( k() \)-day subintervals first.

`icer(#)` specifies the option for performing cost-effectiveness analysis: 1 for calculating the ICER and its CI and 0 otherwise. The default is `icer(0)`.

`k(#)` specifies the number of days in a time interval for discounting costs. The default is `k(90)`. When the time interval between start and stop dates is smaller than \( k() \), the costs accumulated are discounted using the start date; if the time interval is bigger than \( k() \), then the interval is divided into \( k() \)-day subintervals first, and the costs at each subinterval are discounted at the start date of the subinterval.
level(#) specifies the level for CIs. The value must be between 10 and 99. The default is level(95).

method(#) specifies the method for mean costs estimation: 1 for using cost history (ZT estimator), 0 for using only total costs (BT estimator). The default is method(1). If method() is 0 and drcost() is 0, start() and stop() are not required. When start() and stop() are not provided, the costs are assumed to be collected within the time limit $L$.

The ZT estimator requires the arguments start() and stop() to be provided. On the other hand, because the BT estimator requires only total costs, the arguments start() and stop() may be omitted for the BT method. However, if the user wants to discount the costs with the BT method, start() and stop() are required for discounting purposes.

6 Example

To illustrate the hcost command, we use a small simulated dataset as an example. Among the patients in our data, 80 received the conventional therapy (group 0), and 80 received the new treatment (group 1). The first enrolled patient was followed for 69 months and the last for less than 1 month, with an average follow-up of 27 months. Cost data were simulated with start and stop dates for each entry.

Table 1 shows, as an example, several rows of the dataset. Each patient has a unique id variable; a survival time variable, surv; a death indicator variable, delta; a treatment variable, trt; and one or more cost entries representing different types of costs or costs accumulated at different time intervals defined by the start date variable, start, and the stop date variable, stop.

<table>
<thead>
<tr>
<th>id</th>
<th>start</th>
<th>stop</th>
<th>cost</th>
<th>trt</th>
<th>delta</th>
<th>surv</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>3694</td>
<td>0</td>
<td>0</td>
<td>575</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>9</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>575</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>9</td>
<td>12</td>
<td>0</td>
<td>0</td>
<td>575</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>34</td>
<td>106</td>
<td>0</td>
<td>0</td>
<td>575</td>
</tr>
<tr>
<td>2</td>
<td>5</td>
<td>11</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>1166</td>
</tr>
<tr>
<td>2</td>
<td>5</td>
<td>26</td>
<td>275</td>
<td>0</td>
<td>0</td>
<td>1166</td>
</tr>
<tr>
<td>1001</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1138</td>
</tr>
<tr>
<td>1001</td>
<td>1</td>
<td>7</td>
<td>49325</td>
<td>1</td>
<td>0</td>
<td>1138</td>
</tr>
<tr>
<td>1001</td>
<td>1</td>
<td>23</td>
<td>112</td>
<td>1</td>
<td>0</td>
<td>1138</td>
</tr>
<tr>
<td>1009</td>
<td>1</td>
<td>1</td>
<td>115</td>
<td>1</td>
<td>1</td>
<td>425</td>
</tr>
<tr>
<td>1009</td>
<td>1</td>
<td>10</td>
<td>25368</td>
<td>1</td>
<td>1</td>
<td>425</td>
</tr>
<tr>
<td>1009</td>
<td>1</td>
<td>24</td>
<td>41</td>
<td>1</td>
<td>1</td>
<td>425</td>
</tr>
</tbody>
</table>
After importing the data, we declare the data as survival data using the command `stset`. This dataset is multiple-record-per-subject survival data; that is, there may be more than one row belonging to one patient. For the BT estimator, the dataset can also be single-record-per-subject survival data as long as the total costs for each subject are provided in one record.

```
. use example
. * Declare the data as survival data by stset before using hcost
. stset surv, failure(delta)
    failure event:  delta != 0 & delta < .
    obs. time interval:  (0, surv]
    exit on or before:  failure

9882  total observations
     0  exclusions

9882  observations remaining, representing
2988  failures in single-record/single-failure data
10270570  total analysis time at risk and under observation

at risk from t = 0
earliest observed entry t = 0
last observed exit t = 2082
```

We use the command `hcost` to estimate the mean survival time and the mean costs. The time limit $L$ is chosen to be 1,461 days (that is, 4 years), at which time there are still enough subjects under observation. We estimate the mean survival time and the mean costs for the whole population using the default $ZT$ estimator without discounting either costs or survival time.

```
. hcost id cost, start(start) stop(stop) l(1461)
                  Coef.  Std. Err.      z    P>|z|     [95% Conf. Interval]
               cost    80134.84    4870.968   16.45     0.000    70587.92    89681.76
               survival   1164.878    42.04446    27.71     0.000     1082.473    1247.284

Method used:  ZT (using cost history)
Annual discounting rate for costs:  0
Annual discounting rate for survival time:  0
Time limit L (in days):  1461
```

Next, we show an example where an improperly chosen $L$ may lead to problems. We set $L$ to be 3,000 (days), which is larger than the largest observation time in the dataset (2,082 days).

```
. hcost id cost, start(start) stop(stop) l(3000)
Warning: The time limit l is too large (greater than the last observation time).
       r(198);
```
We then calculate the $ZT$ estimator with an annual discounting rate of 3% for both costs and survival time.

```
. hcost id cost, start(start) stop(stop) l(1461) drsurv(0.03) drcost(0.03)
```

|          | Coef. | Std. Err. |   z  |   P>|z| | [95% Conf. Interval] |
|----------|-------|-----------|------|--------|----------------------|
| cost     | 77578.64 | 4602.394 | 16.86 | 0.000  | 68558.12 86599.17   |
| survival | 1101.906 | 39.12913 | 28.16 | 0.000  | 1025.215 1178.598  |

Method used: $ZT$ (using cost history)
Annual discounting rate for costs: .03
Annual discounting rate for survival time: .03
Time limit L (in days): 1461

Next, we perform the estimation for the two treatment groups separately using the $BT$ estimator with an annual discounting rate of 3%. This is achieved using the `group()` argument.

```
. hcost id cost, start(start) stop(stop) l(1461) group(trt) method(0) drsurv(0.03) drcost(0.03)
```

|                      | Coef. | Std. Err. |   z  |   P>|z| | [95% Conf. Interval] |
|----------------------|-------|-----------|------|--------|----------------------|
| Estimates for Group 1 (trt=0) |       |           |      |        |                      |
| cost                 | 64927.99 | 7803.73  | 8.32 | 0.000  | 49632.96 80223.02   |
| survival             | 951.7195 | 63.09655 | 15.08| 0.000  | 828.0525 1075.386  |
| Estimates for Group 2 (trt=1) |       |           |      |        |                      |
| cost                 | 107624 | 9618.589 | 11.19| 0.000  | 88771.95 126476.1   |
| survival             | 1252.687 | 39.27494 | 31.90| 0.000  | 1175.709 1329.664  |
| Estimates for Difference Between Groups (Group 2 - Group 1) |       |           |      |        |                      |
| cost                 | 42696.05 | 12386.1  | 3.45 | 0.001  | 18419.74 66972.36   |
| survival             | 300.9673 | 74.32157 | 4.05 | 0.000  | 155.2997 446.6349  |

Method used: $BT$ (using total costs)
Annual discounting rate for costs: .03
Annual discounting rate for survival time: .03
Time limit L (in days): 1461
We can also obtain the ICER and its CI by using the argument icer(1). In this example, we use the default ZT estimator and an annual discounting rate of 3%.

```
.hcost id cost, start(start) stop(stop) l(1461) group(trt) drsurv(0.03)
> drcost(0.03) icer(1)
Estimates for Group 1 (trt=0)

| Coef. | Std. Err. | z   | P>|z|  | [95% Conf. Interval] |
|-------|-----------|-----|------|----------------------|
| cost  | 64171.55  | 6553.233 | 9.79 | 0.000 | 51327.45 - 77015.65 |
| survival | 951.7195 | 63.09655 | 15.08 | 0.000 | 828.0525 - 1075.386 |

Estimates for Group 2 (trt=1)

| Coef. | Std. Err. | z   | P>|z|  | [95% Conf. Interval] |
|-------|-----------|-----|------|----------------------|
| cost  | 92181.74  | 5535.045 | 16.65 | 0.000 | 81333.25 - 103030.2 |
| survival | 1252.687 | 39.27494 | 31.90 | 0.000 | 1175.709 - 1329.664 |

Estimates for Difference Between Groups (Group 2 - Group 1)

| Coef. | Std. Err. | z   | P>|z|  | [95% Conf. Interval] |
|-------|-----------|-----|------|----------------------|
| cost  | 28010.18  | 8577.971 | 3.27 | 0.001 | 11197.67 - 44822.7 |
| survival | 300.9673 | 74.32157 | 4.05 | 0.000 | 155.2997 - 446.6349 |
```

Method used: ZT (using cost history)
Annual discounting rate for costs: .03
Annual discounting rate for survival time: .03
Time limit L (in days): 1461
Incremental cost-effectiveness ratio (95% CI): 93.067 (38.133, 189.761)

From the two examples shown above, we can see that, as expected, the BT estimator produces larger standard errors than the ZT estimator because it does not use the cost history of censored and uncensored subjects.

7 Stored results

`hcost` stores the following in `e()` (when `group()` is not specified):

Scalars
- `e(n)` number of patients
- `e(nobs)` number of cost observations
- `e(p censor)` censoring rate

Matrices
- `e(b)` coefficient vector
- `e(V)` variance–covariance matrix of the estimators

If `group()` is specified, similar results are stored in `e(n1)`, `e(n2)`, `e(nobs1)`, `e(nobs2)`, `e(p censor1)`, `e(p censor2)`, `e(b1)`, `e(b2)`, `e(V1)`, and `e(V2)` for groups 1 and 2.
The command to list the stored results is shown below.

. ereturn list

Scalars:
- $e(n1) = 80$
- $e(nobs1) = 4869$
- $e(pcensor1) = .49$
- $e(n2) = 80$
- $e(nobs2) = 5013$
- $e(pcensor2) = .75$

Matrices:
- $e(b1) : 1 \times 2$
- $e(V1) : 2 \times 2$
- $e(b2) : 1 \times 2$
- $e(V2) : 2 \times 2$

The following commands display the estimated mean costs, mean survival time, their variances, and covariance for group 1:

. * vector of mean costs and mean survival time for Group 1
. matrix list $e(b1)$
$e(b1)[1,2]$
- cost survival
  r1 64171.554 951.7195

. * Variances and covariance between mean costs and mean survival time for Group 1
. matrix list $e(V1)$
symmetric $e(V1)[2,2]$
- cost survival
  cost 42944860
  survival 113828.12 3981.1744

8 Further comments

A program implementing Lin’s estimators for censored costs was developed by Kim and Thompson (2011). However, Lin’s estimator is appropriate only when cost data are available as grouped (for example, monthly) data. Our program can easily handle the cost data with any start and stop times using the ZT estimator and therefore is more flexible.

In addition to the survival time, quality-adjusted life-years (QALY) are also widely used to measure effectiveness of treatments. QALY are subject to the same kind of “informative censoring” challenge as costs are (Zhao and Tsiatis 1997, 1999; Willan, Lin, and Manca 2005). Hence, the analysis of QALY should not follow traditional survival techniques, similarly to the analysis of cost data. In our future work, we will extend the program such that QALY also can be used as a measure of effectiveness.

Currently, hcost estimates the mean costs and the mean survival time (limited to a time horizon $L$) without using any covariate information. However, it is often of interest to investigate how covariates affect the costs and effectiveness of different therapies. Therefore, our future work will attempt to incorporate covariate information into our program.
9 References


About the authors

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