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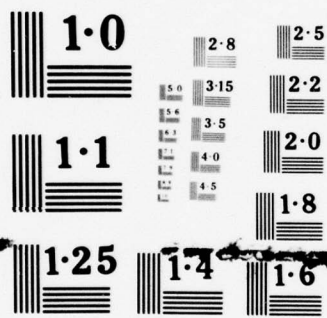
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THESIS

NONPARAMETRIC ESTIMATION
FROM CENSORED DATA

by

Lee Won Hyung

March 1978

Thesis Advisor:

Donald P. Gaver

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This paper proposes alternative estimators and compares them to the product limit method. A computer simulation is used to generate the times of death and truncation from a variety of assumed distributions. No single estimator gives the best fit to the "true" distribution of death under all situations. However, other estimators are shown to be better than the product limit estimator under all of the assumed situations. ↗

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Nonparametric Estimation

from Censored Data

by

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Major, Korean Army
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Submitted in partial fulfillment of the
requirements for the degree of

MASTER OF SCIENCE IN OPERATIONS RESEARCH

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ABSTRACT

For nearly two decades we have witnessed an intensive development of a statistical methodology for assessing length of life and reliability of performance from empirical data. The initial stimulus for research on statistical problems in life testing and reliability came from the need to answer pressing practical questions which could not be treated by the existing statistical techniques. Because life and performance tests are so time consuming and expensive to run, it is a practical necessity to terminate them as soon as possible.

For the statistician this means developing estimation and decision procedure for data, which are severely curtailed in one way or another long before all items on test have actually failed. The estimation is more complicated when the data are truncated, i.e. when the observer loses track of some individuals before death occur. The product limit method of Kaplan and Meier is one way of estimating $p(t)$ when the mechanism causing truncation is independent of the mechanism causing death.

This paper proposes alternative estimators and compares them to the product limit method. A computer simulation is used to generate the times of death and truncation from a variety of assumed distributions. No single estimator gives the best fit to the "true" distribution of death under all situations. However, other estimators are shown to be better than the product limit estimator under all of the assumed situations.

TABLE OF CONTENTS

I.	INTRODUCTION - - - - -	6
II.	THEORY - - - - -	8
III.	THE ESTIMATORS - - - - -	16
	A. STEP-FUNCTION ESTIMATORS - - - - -	17
	B. POINT ESTIMATORS - - - - -	22
	C. BAYESIAN ESTIMATORS - - - - -	27
	D. JACKKNIFE ESTIMATORS OF LOGISTIC TRANSFORMATION - - - - -	30
	E. A PARAMETER ESTIMATOR - - - - -	36
IV.	INSTRUCTIONS FOR USING PROGRAM - - - - -	38
V.	RESULTS OF THE SIMULATION - - - - -	43
	APPENDIX A. ESTIMATORS FOR GROUPED DATA - - - - -	60
	APPENDIX B. LISTING OF COMPUTER PROGRAM - - - - -	63
	LIST OF REFERENCES - - - - -	74
	INITIAL DISTRIBUTION LIST - - - - -	75

I. INTRODUCTION

Let the random variable T denote the time that elapses until an event occurs; the event may for example be an equipment failure, an individual's death, or the detection of a target. Denote by $p(t)$ the probability of survival to time t ,

$$p\{T > t\} = p(t)$$

Picturesquely, T is called a lifetime, and $p(t)$ is a survival probability;

$F(t) = 1 - p(t)$ is the distribution function of T .

In the medical field, one might wish to estimate the probability, $p(t)$ that a patient survives t after a certain surgical procedure for cancer. In electronics, one wishes to estimate the probability of continuous failure-free operation of an equipment for time t . In the military, one might be interested in the probability of conducting a certain mission, under specified environmental conditions, without detection by the enemy. The event of interest may be a human death, equipment malfunctions, or sonar detection. Following Kaplan and Meier, Reference (1), this paper will refer to the event of interest as a "death". The test element in the sample may be a human, a radio, or a submarine. This paper will refer to the test elements as "individuals". Suppose that observed values of T are $t_1, t_2, t_3, \dots, t_N$, so that N lifetimes are observed. In this case an appropriate (unbiased) estimates of survival to time t is

$$\tilde{p}(t) = \frac{\text{number of } t_i \text{'s} > t}{N}$$

Under many circumstances complete lifetimes are not observed; censoring occurs at certain, x_i , beyond which the life of an individual is not known. In such cases construction of an appropriate estimate of the survival probability is more difficult. In this paper various estimates of survival probability are studied when lifetimes are randomly censored. This means that censoring times are assumed to be realizations random variables independent of the actual lifetimes.

The product-limit estimator of Kaplan and Meier, Reference (1), is an accepted method of dealing with the problem of censored data. This paper presents thirteen non-parametric estimators, including the product limit function. Censored data sets are simulated. The thirteen estimators are compared by examining their performance on the simulated data bases.

II. THEORY

There are two approaches to the empirical estimation of the survival probability, $p(t)$:

- (1) one may use the observed fraction of survivors at arbitrarily selected times (step function estimator), or
- (2) one may focus attention on the times of the observed deaths (point estimator).

The initial discussion is based on the assumption that all observations are complete, i.e., it is assumed that all individuals remain under observation until their time of death. This initial assumption is for the purpose of simplifying the discussion. Then, later in this paper, the discussion is broadened to include incomplete data with observations of both death and censoring events.

Survival Probabilities; No Censoring

Let $0 = t_0 < t_1 < t_2 \dots < t_i < t_{i+1} < \dots$ be a sequence of fixed times. Then if T is a lifetime

$$p(t_i) = p\{T > t_i\}$$

and denote the conditional probability of survival to time t_i , given survival to t_{i-1} by

$$\begin{aligned} p(t_i | t_{i-1}) &= p\{T > t_i | T > t_{i-1}\} \\ &= \frac{p\{T > t_i\}}{p\{T > t_{i-1}\}} = \frac{p(t_i)}{p(t_{i-1})} \end{aligned} \quad (1)$$

If $p(t_{i-1}) = 0$, define $p(t_i | t_{i-1}) = 0$.

Then

$$\begin{aligned} p(t_i) &= p(t_i | t_{i-1}) p(t_{i-1}) = p(t_i | t_{i-1}) \cdot p(t_{i-1} | t_{i-2}) p(t_{i-2}) \\ &= \prod_{j=1}^i p(t_j | t_{j-1}) \end{aligned} \quad (2)$$

where $p(t_i | t_0) = p(t_i)$; $p(0) = 1$.

Observations on Uncensored Data at Fixed Times

Let a sample of N individuals come under observation. They are all observed from birth (or the appropriate event defining time zero) until death. With the first approach, preselects a series of times, $0 < t_1 < t_2 < \dots$ before examining the observed time of death. In the medical follow-up example, one might select the times corresponding to exactly 1, 2, 3, ... years after a surgical procedure for cancer. An estimate of the conditional probability of survival to t_i , given survival to t_{i-1} is

$$\tilde{p}(t_i | t_{i-1}) = \frac{N_i - r_i}{N_i} \quad (3)$$

With N_i elements were present at the beginning of the interval, i.e., at time t_{i-1} , and r_i elements failed during the interval.

For a set of data which is not censored, $N_i = N_{i-1} - r_{i-1}$. Now replace probabilities by their estimates in (2):

$$\tilde{p}(t_i) = \prod_{j=1}^i \tilde{p}(t_j | t_{j-1}) = \prod_{j=1}^i \left(\frac{N_j - r_j}{N_j} \right)$$

$$\begin{aligned}
&= \left(\frac{N-r_1}{N}\right) \left(\frac{N-r_1-r_2}{N-r_1}\right) \cdots \left(\frac{N-r_1-\dots-r_{i-2}}{N-r_1-\dots-r_{i-2}}\right) \left(\frac{N-r_1-\dots-r_i}{N-r_1-\dots-r_{i-1}}\right) \\
&= 1 - \frac{\sum_{j=1}^i r_j}{N}
\end{aligned}$$

Now the estimate $p(t_i)$ is of the form

$$\tilde{p}(t_i) = \frac{N-(r_1 + r_2 + \dots + r_i)}{N},$$

and this is the same as

$$\tilde{p}(t_i) = \frac{S_i}{N}$$

where S_i is the number of the original group, of size N , that survive to t_i . If it is assumed that the N individuals each have the survival probability $p(t)$, and that they die independently, then S_i , the random number that survive to time t_i is binomially distributed, with S_i being a realized value of S_i . Then, considering the estimate as a random variable,

$$\tilde{p}(t_i) = \frac{S_i}{N}$$

and

$$E[\tilde{p}(t_i)] = \frac{N p(t_i)}{N} = p(t_i)$$

and

$$\text{Var}[\tilde{p}(t_i)] = \frac{p(t_i)(1-p(t_i))}{N}$$

Consequently $\tilde{p}(t_i)$ is an unbiased and consistent estimate of $p(t_i)$. This is true for every t_i , and can be shown to be true for all t_i , $i=1, 2, \dots, I$, as will.

All of this indicates that the estimate suggested is likely to be a good one if the sample size, N , is large.

Clearly $\tilde{p}(t_i) \leq \tilde{p}(t_{i-1})$. The survival probability, $p(t)$, is thus estimated at a fixed sequence of times. At each time point, t_i being a typical one, there are r_i fewer survivors than at t_{i-1} , where $r_i = 0, 1, 2, \dots, N$. Consequently a plot of $\tilde{p}(t_i)$ shows a non-decreasing step function, with downward steps of varying sizes at t_1, t_2, \dots .

If the above times are close together, and if the time of death T , has a density function, then one can anticipate seeing values of r_i that are either zero or unity.

The so-called second approach is really a limiting case of the first, as the time of intervals of measurement decrease indefinitely. Thus when a death (or loss) occurs it is only a single event.

When no losses take place, the case now considered, the time t_i of the i th death is a really a realization of a random variable, denoted by \underline{t}_i ; this means that $p(\underline{t}_i)$ the probability of surviving \underline{t}_i , is a random variable. It can be shown that the expected value of $p(\underline{t}_i)$ is

$$E[p(\underline{t}_i)] = \frac{N-i+1}{N+1}, \quad i=1, 2, \dots, N$$

where $\underline{t}_1 < \underline{t}_2 < \dots < \underline{t}_N$.

The derivation involves integrating

$$\begin{aligned} E[p(\underline{t}_i)] &= \int_0^{\infty} p(t) \cdot \frac{N!}{(i-1)!(N-i+1)!} [1-p(t)]^{i-1} \left(-\frac{dp(t)}{dt}\right) [p(t)]^{N-i} dt \\ &= \frac{N-i+1}{N+1} \end{aligned}$$

by transformation from $p(t)$ to x ; see Cramér, Mathematical Methods of Statistics, H. Cramér, Princeton University Press, 1946.

Thus one is led to use

$$\tilde{p}(t_i) = \frac{N-i+1}{N+1} \quad (4)$$

as an estimate of the value of $p(t_i)$, t_i being the i th time of death. Expression (4) provides estimator of the survival function at times of observed deaths when there are no losses because of censoring. The estimator at the points $t_i: t_1 < t_2 < \dots < t_N$, can be connected by straight lines, or a step function with step sizes $1/(N+1)$ may be used.

The estimators of equation (4) give intuitively acceptable results. For example, if the sample consists of only a single individual ($N=1$), then death is equally likely to occur before or after the time at which the true (but unknown) survival function equals one half. Thus, the result of equation (4) is reasonable:

$$E[\tilde{p}(t_1)] = \frac{1}{2}$$

The point estimates of the second approach always occur at the times of discontinuity forestimates from the first approach. For example, consider a data base ($N=4$) with deaths observed at times 1, 3, 4 and 7. The first approach gives the following step function estimate of the survival function:

$$\tilde{p}(t) = \begin{cases} 1.0 & 0 \leq t < 1 \\ 0.75 & 1 \leq t < 3 \\ 0.5 & 3 \leq t < 4 \\ 0.25 & 4 \leq t < 7 \\ 0.0 & t < 7 \end{cases}$$

The second approach gives the following point estimates

$$p(0) = 1.0$$

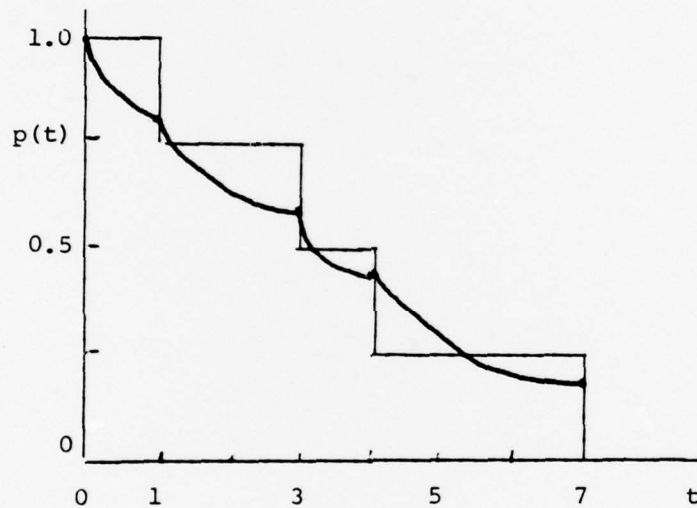
$$p(1) = 0.8$$

$$p(3) = 0.6$$

$$p(4) = 0.4$$

$$p(7) = 0.2$$

A graphic comparison of the results of the two approaches is given below:



It is difficult to decide how to smooth out the step functions that result from the first approach. By connecting the tops of the "stairsteps," one places an upper bound on reasonable estimates. By connecting the bottom corners of the stairsteps, one places a lower bound on reasonable estimates. One might draw a smooth, decreasing curve that passes through all (or almost all) of the vertical faces of the step-function estimate. The second approach suggests method of selecting a unique point on each of these vertical segments.

Incomplete observations

When some of the observations are incomplete, equation (4) requires modification. The expected value of the survival function at the time of the first observed death may be written:

$$E[\tilde{p}(t_1)] = \frac{N_1}{N_1 + 1} \quad (5)$$

Here N_1 is the effective size of the sample during the interval terminated by the time observed for the first death $(0, t_1)$. In the special case of no censoring events, the value of N_1 is unambiguous. It is equal to the initial sample size ($N_1 = N$). In this case equation (5) reduces to equation (4).

Subsequent point estimates for t_2, t_3, \dots may be calculated iteratively:

$$E[\tilde{p}(t_i)] = \frac{N_i}{N_i + 1} \cdot E[p(t_{i-1})]$$

where $t_0 = 0$ and N_i is the effective sample size over the time interval (t_{i-1}, t_i) . Thus,

$$E[\tilde{p}(t_i)] = \prod_{j=1}^i \left(\frac{N_j}{N_j + 1} \right) \quad (6)$$

Variance of the estimators

Kaplan and Meier, reference (1), give an expression for the exact calculation of the variance of step functions. They also discuss "Greenwood's formula," a large sample approximation that ignores terms of order $1/N_i^2$.

Herd, reference (2), presents without derivation an expression for the variance of estimates using the second approach (point estimators):

$$V(t_i) = \text{Var} \{E[\tilde{p}(t_i)]\} = \prod_{j=1}^i \left(\frac{N_j}{N_j+2}\right) - \prod_{j=1}^i \left(\frac{N_j}{N_j+1}\right)^2$$

The notation here follows that for the estimating equation (6).

III. THE ESTIMATORS

This section describes the nine non-parametric estimators and four jackknife estimators of the survival probability. It also describes the parametric estimator for an exponential decay function. Exponential life distributions are the starting point for much of reliability theory and practice. The estimator derived from the exponential is regarded as "par" when the simulated data is based on an underlying exponential decay distribution for deaths. Thus, when deaths are exponentially distributed, the non-parametric estimators may be compared relative to each other, and they may be compared with the parametric estimator as a standard.

A hypothetical data base, consisting of five individuals, is used to illustrate each of the estimators. This sample data base is as follows:

<u>Individual</u>	<u>Time of Death</u>	<u>Time of Truncation</u>
A	1	-
B	Unknown (>2)	2
C	3	-
D	Unknown (>6)	6
E	7	-

The data have been arranged in time sequence of the death and truncation events. In the medical example, the data might indicate that patients A, C and E were observed to die exactly 1, 3 and 7 years, respectively, after their surgery. However, B and D moved away or otherwise became unavailable to the observer at these times. Further, the

cause of the unobservability is unrelated to the patient's health and life expectancy.

A. STEP-FUNCTION ESTIMATORS

1. The First Estimator, " $\tilde{p}_1(t)$ "

$\tilde{p}_1(t)$ is a naive estimator; it is expected to perform poorly relative to the other estimators. \tilde{p}_1 only depends on the data from individuals whose deaths are observed. It ignores any information from the partial lifetimes noted for the censored observations. $\tilde{p}_1(t)$ is simply the fraction of individuals surviving to at least time t among those individuals whose time of death is known. It is a step function:

<u>t</u>	<u>$\tilde{p}_1(t)$</u>
0-1	1.0
1-3	0.667
3-7	0.333
7- ∞	0.00

The naive estimator, $\tilde{p}_1(t)$, takes no account of the successful survival intervals observed for the censored individuals. Therefore it is biased in a downward (pessimistic) direction.

2. The Second Estimator, " $\tilde{p}_2(t)$ "

$\tilde{p}_2(t)$ is the product-limit estimate. Kaplan and Meier, reference (1), have shown that this is the maximum likelihood estimator. The observed events, both deaths and truncations, are arranged in increasing order of occurrence: t_1, t_2, \dots, t_N ; where N is the number of individuals in the sample.

Let $p(t_i)$ denote the cumulative probability of survival of an individual from time zero to time t_i . Let $p(t|t_i)$ denote the conditional probability of surviving to time $t(> t_i)$, given that the individual has

already survived to time t_i . Then,

$$\tilde{p}_2(t_i) = p_2(t_{i-1}) \cdot p_2(t_i | t_{i-1}) \quad (E-1)$$

If we define $t_0 = 0$ and $p(0) = 1$, then

$$\tilde{p}_2(t_i) = \prod_{j=1}^i p_2(t_j | t_{j-1}) \quad (E-2)$$

The product limit estimator is in the form of equation (E-2) with

$$\tilde{p}_2(t_j | t_{j-1}) = \begin{cases} \frac{N_j}{N_j} = 1 & \text{If the event at } t_j \text{ is truncation} \\ \frac{N_j - 1}{N_j} & \text{If the event at } t_j \text{ is a death} \end{cases} \quad (E-3)$$

Here n_j is the number of individuals observed surviving in the interval $t_{j-1} < t < t_j$. This formulation causes the product limit estimator to be insensitive to the exact time of the censoring events.

The estimator is unity from time zero to the time of the first event, t_1 , reflecting the fact that all individuals in our example are observed to live until at least time t_1 .

- If the event at time t_1 is a truncation, then the estimator remains at unity until at least time t_2 . Again, no deaths are observed in the sample before t_2 .
- If the event at time t_1 is a death, then the estimator drops to $(N-1)/N$. This drop reflects the observed death of $1/N$ of the survival sample just prior to t_1 .

Values of the estimator \tilde{p}_2 are calculated iteratively at successive values of t_i ($i=1,2,\dots,N$).

The size of the survival sample declines as truncations and deaths remove individuals from observation. For the hypothetical data base listed above, one obtains:

<u>t</u>	<u>$\tilde{p}_2(t)$</u>
0-1	5/5 = 1.0
1-2	4/5 = 0.8
2-3	(4/5) x (3/3) = 0.8
3-6	(4/5) x (2/3) = 0.533
6-7	(8/15) x (1/1) = 0.533
7- ∞	(8/15) x (0/1) = 0.0

The product-limit estimator explicitly accounts for the survival of these individuals (up to the time of the last death before each censoring event). Thus, $\tilde{p}_2(t)$ is a step function with a value that is not less than $\tilde{p}_1(t)$ for any value of t . If the sample contains no censoring, then $\tilde{p}_1(t)$ and $\tilde{p}_2(t)$ are identical.

If the last event in the sample is a truncation rather than a death, then the modified data give the following estimate, i.e., individual E had disappeared from the observer at time 6.5 (so that the fact of E's death at time 7 is unknown).

<u>t</u>	<u>$\tilde{p}_2(t)$ - Modified data</u>
0-1	1.0
1-3	0.8
3-6.5	0.533

Since the time of the death for individual E is now unknown, one can only estimate that:

$$0 \leq \tilde{p}_2(t) \leq 0.533 \text{ for } t > 6.5$$

If the analyst is willing to assume a functional form for the survival function, then he may calculate the manner in which the estimator $\tilde{p}_2(t)$ decreases to zero. However, the data alone are insufficient when a strictly non-parametric estimator is used.

The product-limit estimator is a useful and intuitively appealing method of dealing with incomplete observations. It has been widely used and studied. However, the product-limit has one disturbing characteristic:

Most of the biological, physical or other causes of deaths produce a survival probability that continuously decreases in time. It is, therefore, one may be a little uncomfortable estimating the survival probability with a step function. One is tempted to smooth the estimator to make it a monotonic decreasing function of t .

3. The Third Estimator, " $\tilde{p}_3(t)$ "

$\tilde{p}_3(t)$ is a modification of $\tilde{p}_2(t)$. Like $\tilde{p}_2(t)$, it is a step function with discrete drops at those times corresponding to the observed deaths in the sample population. It may also be expressed as a product of conditional probabilities:

$$\tilde{p}_3(t_i) = \prod_{j=1}^i \tilde{p}_3(t_k | t_{k-1}) \quad (E-4)$$

where the t_k are the times of observed deaths and t_0 is zero. The conditional probabilities on the right-hand side of Equation (E-4) differ somewhat from those in Equation (E-2):

$$\tilde{p}_3(t_k | t_{k-1}) = \frac{N_k - 1}{N_k} \quad (E-5)$$

Equation (E-5) differs from Equation (E-3) in the interpretation of the numbers of individuals at risk. Here, the value of N_k is taken to be the average number of individuals observed surviving in the interval between the $(k-1)$ st observed death and the k th observed death. The number of observed survivors decrease at intermediate times if events are censored, and hence the N_k are not necessary integers.

The value of N_k is regarded as the effective sample size for the interval from t_{k-1} to t_k . In the sample data base shown above, individual B is known to have survived from time 1 to time 2, or half of the interval between the first death at $t=1$ and the second death at $t=3$. Therefore, the estimator p_3 treats individual B as half a participant in the interval between the death of individuals A and C.

The effective sample size for this interval is then 3.5 ($n = 3 + \frac{(2-1)}{(3-1)} = 3.5$) (full contributions from individuals C, D and E, plus a half contribution from B). For our hypothetical data base, the following values are calculated for \tilde{p}_3 :

<u>t</u>	<u>$\tilde{p}_3(t)$</u>
0-1	5/5 = 1.0
1-3	4.5 x 1.0 = 0.8
3-7	(2.5/3.5) x 0.8 = 0.571
(7)	(1.75/2.75) x 0.571 = 0.364

The value of $\tilde{p}_3(t)$ can never be less than the corresponding value of $\tilde{p}_2(t)$. In the special case with no censoring events the estimators $\tilde{p}_1(t)$, $\tilde{p}_2(t)$ and $\tilde{p}_3(t)$ are identical.

One might perturb the data by shifting the time of B's truncation event down to $1+\epsilon$ or up to $3-\epsilon$, ϵ arbitrarily small. The dependence of the estimator \tilde{p}_3 upon the exact time of the censoring events may now be demonstrated.

For purposes of illustration, the time of the censoring event for individual $B(t_2)$ is decreased from 2 to 1.1, then increased to 2.9.

<u>t</u>	<u>$\tilde{p}_3(t), t_2 = 2$</u>	<u>$\tilde{p}_3(t), t_2 = 1.1$</u>	<u>$\tilde{p}_3(t), t_2 = 2.9$</u>
0-1	1.0	1.0	1.0
1-3	0.80	0.80	0.80
3-7	0.571	0.538	0.597
(7)	0.364	0.342	0.380

This example demonstrates an intuitively appealing characteristic of the estimator, \tilde{p}_3 . As the total observed survival time increases for the individuals in our sample (with deaths held constant), the value of the estimating function increases over at least a portion of its range.

We may safely assume that the true survival function eventually tends to zero with time, since no physical or biological system lives forever. However, there are no observations on the survival of individuals beyond time 7. The data only indicate that our step-function estimator drops to a value of .364 at $t=7$, but the nonparametric estimator gives no information about the survival function's subsequent decline from .364 to zero. However, the data alone are insufficient when a strictly nonparametric estimator is used.

B. POINT ESTIMATOR

As mentioned above, the estimators \tilde{p}_1 , \tilde{p}_2 and \tilde{p}_3 are somewhat undesirable because they give step-function estimates for a continuous survival function. The next three estimators \tilde{p}_4 , \tilde{p}_5 and \tilde{p}_6 are modification of the first three. Again they provide estimates of the survival function only at those points in time that correspond to observed deaths.

These estimators are specified by Equations (E-2) and (E-4), except for a substitution of the term $(N+1)$ in place of (N) .

Since the point estimators have rigorous definitions at only discrete points in time, it is necessary to offer an interpolation rule. That is, we need a method of "connecting the dots." The method proposed here is to assume that the survival function declines in a piece-wise exponential decay between the discrete points in time. This procedure is equivalent to assuming that the hazard function is essentially constant between a consecutive pair of the discrete times, but that the hazard varies from one time period to the next. Such an assumption is intuitively acceptable unless one suspects violent fluctuations in the hazard function.

1. The Estimator, " $\tilde{p}_4(t)$ "

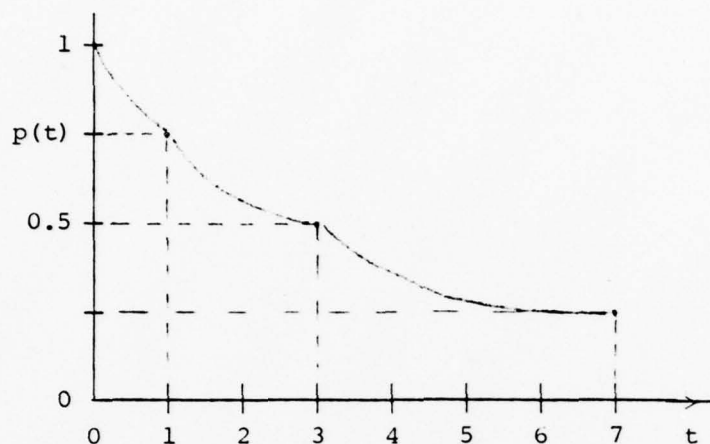
$\tilde{p}_4(t)$ is analogous to $\tilde{p}_1(t)$ in that only those individuals observed to die are included in the sample. These two estimators are naive because they suppress all data from the survival times of individuals terminated from observation by censoring.

These estimates, i.e., $\tilde{p}_1(t)$ and $\tilde{p}_4(t)$, tend to ignore information from the more long-lived individuals in the sample, and they may be expected to give biased estimates of the survival function.

The point estimator $\tilde{p}_4(t)$ gives the following values with sample data base presented earlier in this section.

t	$\tilde{p}_4(t)$	Interpolation	
		t	$\tilde{p}_4(t)$
0	1.0	0-1	$e^{t \cdot \ln(3/4)}$
1	$3/4 = 0.75$		
3	$(\frac{2}{3}) \times 0.75 = 0.5$	1-3	$(\frac{3}{4}) e^{\frac{t-1}{2} \cdot \ln(2/3)}$
7	$(\frac{1}{2}) \times 0.5 = 0.25$	3-7	$(\frac{1}{2}) e^{\frac{t-3}{4} \cdot \ln(1/2)}$

The interpolation for connecting the dots are as follows:



t	$0 \leq t \leq 1$	$1 \leq t \leq 3$	$3 \leq t \leq 7$
p(t)	$e^{-t/\tau}$	$p(t-1) e^{-\frac{t-1}{\tau}}$	$p(t-1) e^{-\frac{t-3}{\tau}}$
interpolation	$t = 1$ $\frac{3}{4} = e^{-\frac{1}{\tau}}$ $\frac{1}{\tau} = -\ln(\frac{3}{4})$	$t = 3$ $\frac{1}{2} = \frac{3}{4} e^{-\frac{2}{\tau}}$ $\frac{1}{\tau} = \frac{-\ln(\frac{2}{3})}{2}$	$t = 7$ $\frac{1}{4} = \frac{1}{2} e^{-\frac{4}{\tau}}$ $\frac{1}{\tau} = \frac{-\ln(\frac{1}{2})}{4}$
p(t)	$e^{t \cdot \ln(\frac{3}{4})}$	$(\frac{3}{4}) e^{\frac{t-1}{2} \cdot \ln(\frac{2}{3})}$	$(\frac{1}{2}) e^{\frac{t-3}{4} \cdot \ln(\frac{1}{2})}$

2. The Estimator, " $\tilde{p}_5(t)$ "

The estimator $\tilde{p}_5(t)$ similarly corresponds to the product-limit estimator $\tilde{p}_2(t)$. These two estimators use information from the individuals on whom there are censored observations. \tilde{p}_5 , like \tilde{p}_2 , does not exploit information about that portion of the censored observation after the death event (of some other individual) preceding the censoring event.

For our hypothetical data base the following values are calculated for $\tilde{p}_5(t)$:

t	$\tilde{p}_5(t)$	Interpolation	
		t	$\tilde{p}_5(t)$
0	1.0	0-1	$e^{t \cdot \ln(t/6)}$
1	$5/6 = 0.833$		
3	$(\frac{3}{4}) \times 0.833 = 0.625$	1-3	$(\frac{5}{6}) e^{\frac{t-1}{2} \ln(\frac{3}{4})}$
7	$(\frac{1}{2}) \times 0.625 = 0.312$	3-7	$(\frac{5}{8}) e^{\frac{t-3}{4} \ln(\frac{1}{2})}$

Whenever censored observations are present, the estimator $\tilde{p}_4(t)$ never exceeds $\tilde{p}_5(t)$.

For $\tilde{p}_5(t)$, the value of N_i is taken to be the number of surviving individuals in the sample just before the observation of the ith death. This value is smaller than the number of surviving individuals just after the (i-1)st death if any truncation events occur in the interval. In fact, N_i is the smallest number of surviving individuals observed at any time during the interval (t_{i-1}, t_i) . Thus \tilde{p}_5 might be expected to introduce a bias by using values of N_i that are, on the average, too small. However, this bias would be much less severe than the bias anticipated for the estimator $\tilde{p}_4(t)$.

The estimators \tilde{p}_4 and \tilde{p}_5 are insensitive to the precise times of the censoring events. A change in the time of the censoring event for individual B to $1+\epsilon$ to $3-\epsilon$, ϵ arbitrarily small, does not alter the estimates from \tilde{p}_4 and \tilde{p}_5 given in the preceding paragraph.

3. The Estimator, " $\tilde{p}_6(t)$ "

The estimator $\tilde{p}_6(t)$ corresponds to $\tilde{p}_3(t)$ by accounting for all of the survival time for the truncated observations. For our hypothetical data base, the following values are calculated for $p_6(t)$:

t	$\tilde{p}_6(t)$	Interpolation	
		t	$\tilde{p}_6(t)$
0	1.0	0-1	$e^{t \cdot \ln(5/6)}$
1	$5/6 = 0.833$		
3	$(\frac{3.5}{4.5}) \times 0.833 = 0.648$	1-3	$(\frac{5}{6}) e^{\frac{t-1}{2} \ln(\frac{3.5}{4.5})}$
7	$(\frac{1.75}{2.75}) \times 0.648 = 0.412$	3-7	$(0.648) e^{\frac{t-3}{4} \ln(\frac{1.75}{2.75})}$

The estimator $\tilde{p}_6(t)$ is based on the average number of surviving individuals noted in the various time intervals. These estimators give part credit for individuals whose lifetime is censored in mid-interval. The value of N_i for $\tilde{p}_6(t)$ is an unweighted time average. If the observation of an individual is truncated after 23% of the interval has elapsed, then that individual contributes a value of 0.23 to N_i . Individuals who are observed to survive the entire interval, and the individual whose death terminates the interval each contribute a value of 1.0 to N_i . This interpretation of the effective sample size is approximate if the hazard is approximately constant over the interval. If the hazard function changes markedly within a time interval containing censored events, then this interpretation of the effective sample size is biased. Therefore, the procedure of determining the value of N_i for the estimator $\tilde{p}_6(t)$ is based on the implicit assumption that the survival function is locally

exponential. If the hazard function may be assumed to vary slowly over each of the time intervals (t_{i-1}, t_i) then \tilde{p}_6 would appear to be biased on an acceptable approximation.

The estimator \tilde{p}_6 , like \tilde{p}_3 , depends on the precise times of all deaths and censoring events.

<u>t</u>	<u>$\tilde{p}_6(t), t_2 = 2$</u>	<u>$\tilde{p}_6(t), t_2 = 1.1$</u>	<u>$\tilde{p}_6(t), t_2 = 2.9$</u>
0	1.0	1.0	1.0
1	5/6 = 0.833	5/6 = 0.833	5/6 = 0.833
3	$(\frac{3.5}{4.5}) \times 0.833 = 0.648.$	$(\frac{3.05}{4.05}) \times 0.833 = 0.628.$	$(\frac{3.95}{4.95}) \times 0.833 = 0.665$
7	$(\frac{1.75}{2.75}) \times 0.648 = 0.412.$	$(\frac{1.75}{2.75}) \times 0.628 = 0.399.$	$(\frac{1.75}{2.75}) \times 0.665 = 0.423$

This illustrates that an increase (or decrease) in the total observed survival time causes an increase (or decrease) in the estimate \tilde{p}_6 over at least some of its time range.

If the last event is a censored, and not an observed, death, these estimators also require definition for the time period starting with the time of the last death and ending with the time of the final censoring event.

The method proposed here for $\tilde{p}_4(t)$ and $\tilde{p}_5(t)$ is to continue the exponential function used in the interval terminated by the time of the last death. This procedure can be illustrated with the modified data base used above in the discussion of \tilde{p}_2 and \tilde{p}_3 .

C. THE BAYESIAN ESTIMATORS

Consideration is next given to quasi-Bayesian estimators based on a uniform prior distribution on the unit interval. Let X_1, \dots, X_N be the true survival times of N individuals which are censored on the right by N follow-up times Y_1, \dots, Y_N . It is assumed that the X_i are independent,

identically distributed random variables with common distribution $p(t)$ and we wish to estimate the survival function

$$p(t) = \Pr(X > t)$$

However, we only have available the data,

$$Z_i = \min \{X_i, Y_i\}$$

$$\delta_i = \begin{cases} 1 & \text{if } X_i \leq Y_i \\ 0 & \text{if } X_i > Y_i, i=1, \dots, n \end{cases}$$

If $\delta_i = 0$, then Z_i is called "a loss", and if

$\delta_i = 1$, then Z_i is called "a death".

Then $p_r[\delta_i = 1] = p_r[X_i > t] = p(t)$, $i=1, \dots, N$.

The maximum likelihood estimator for $p(t)$ is

$$\hat{p}(t) = \frac{s}{N} \quad \text{where} \quad s = \sum_{i=1}^N \delta_i$$

is the number of successful tests, s has the binomial distribution.

$$P(S|p) = \binom{N}{s} p^s (1-p)^{N-s}, \quad s=0, 1, \dots, N, \quad 0 < p < 1$$

$$f_p(p) = 1, \quad 0 < p < 1$$

The joint density of s and p is

$$f_{s,p}(s,p) = \binom{N}{s} p^s (1-p)^{N-s}, \quad 0 < p < 1, \quad s=0, 1, \dots, N.$$

The marginal for s is

$$p_s(s) = \int_0^1 \binom{N}{s} p^s (1-p)^{N-s} dp = \binom{N}{s} \cdot \frac{s! (N-s)!}{(N+1)!} = \frac{1}{N+1}$$

for $s=0,1,\dots,N$. Thus, averaging over the values of p , all of which are assumed to be equally likely, the values of s are equally likely to occur. The posterior for p then is

$$f_{p|S}(p|S) = \frac{\Gamma(N+2)}{\Gamma(s+1)\Gamma(N-s+1)} p^s (1-p)^{N-s}, \quad 0 < p < 1,$$

a beta density with parameters $s+1$ and $N-s+1$. The mean of the posterior is $(s+1)/(N+2)$ and the modal (maximum value) of the posterior is s/N ; thus the Bayes estimate of p (given s survivors occur in the sample of N) is

$$p^* = \frac{s+1}{N+2} \quad (C-1)$$

Then, equation (C-1) yields a step function and also has shown that the uniform prior has the effect of adding two individuals to the population at risk with one dying at time zero and the other essentially immortal.

The Bayesian estimators based on a uniform prior distribution on the unit interval are denoted $\tilde{p}_{11}(t)$, $\tilde{p}_{12}(t)$ and $\tilde{p}_{13}(t)$, that correspond, respectively, to the estimators $\tilde{p}_1(t)$, $\tilde{p}_2(t)$ and $\tilde{p}_3(t)$. The sample data base thus gives the following estimates of the survival function:

<u>t</u>	<u>$\tilde{p}_{11}(t)$</u>	<u>$\tilde{p}_{12}(t)$</u>	<u>$\tilde{p}_{13}(t)$</u>
0-1	4/5 = 0.8	6/7 = 0.857	6/7 = 0.857
1-3	3/5 = 0.6	$(\frac{5}{6}) \times 0.857 = 0.714$	$(\frac{5}{6}) \times 0.857 = 0.714$
3-7	2/5 = 0.4	$(\frac{3}{4}) \times 0.714 = 0.536$	$(\frac{3.5}{4.5}) \times 0.714 = 0.556$
(7)	1/5 = 0.2	$(\frac{1}{2}) \times 0.536 = 0.268$	$(\frac{1.75}{2.75}) \times 0.556 = 0.354$

At the time of the final event (whether a death or a truncation), these step-function estimators drop to some positive value. Again, we have no data to indicate how the survival function proceeds to zero at subsequent times.

D. THE JACKKNIFE ESTIMATOR

We will assume that we observed, or have generated in a simulation, a survival probability $p(t_j)$, $j=1, \dots, n$, from various sample sizes. Furthermore we have some parameter or characteristic $p(t_j)$ of the sample size which we wish to estimate with an estimator $\hat{p}(t_j)$. The jackknife estimator $\tilde{p}(t, n)$ described below is an approximately unbiased estimator of $p(t_j)$. A modification of it has other useful properties.

$\tilde{p}_{-i}(t, n-1)$ is the estimator from the sample of n of the X_i 's with the i th value deleted from the sample.

$$\tilde{p}_i(t, n) = n \tilde{p}(t, n) - (n-1) \tilde{p}_{-i}(t, n-1) \quad i=1, \dots, n$$

$$\tilde{p}(t, n) = \frac{1}{n} \sum_{i=1}^n \tilde{p}_i(t, n) = n \tilde{p}(t, n) - \frac{n-1}{n} \sum_{i=1}^n \tilde{p}_{-i}(t, n-1)$$

the $\tilde{p}_i(t, n)$, called the PSEUDO-values.

The PSEUDO-values can be used to obtain variance estimates of $\tilde{p}(t, n)$ and to set approximate confidence limits, using Student's t .

The idea is that the PSEUDO-values will be approximately independently and normally distributed. The jackknife estimator $\tilde{p}(t, n)$ is a sample average so we form an estimate $S_{\tilde{p}(t, n)}^2$ of its variance given by the following relationship (Miller, 1974):

$$S^2 = \frac{\sum \tilde{p}_i^2(t, n) - \frac{1}{n} (\sum \tilde{p}_i(t, n))^2}{n-1}$$

$$S_{\tilde{p}(t, n)}^2 = \frac{S^2}{n}$$

This procedure is particularly useful if the number of data points is small, but it must be used with care. Note, that the estimator $\tilde{p}(t, n)$ is designed to eliminate a $\frac{1}{n}$ bias term in the estimator $\tilde{p}(t, n)$. Of course the computational aspects of the complete jackknife can be quite onerous, especially if $\tilde{p}(n)$ were, say, a complicated maximum likelihood estimator. Miller, reference (4) has shown that the product limit estimator is its own jackknife.

Logistic Transformation

Although one can legitimately jackknife the Kaplan-Meier estimate directly, there is some reason to believe that a preliminary transformation will give improved results. Consequently, consider the transformation

$$\ell = \ln\left(\frac{\tilde{p}(t)}{1-\tilde{p}(t)}\right)$$

and notice that where the range of $\tilde{p}(t)$ is from zero to unity, the above transformation makes the range of ℓ run from $-\infty$ to ∞ . The procedure utilized will be as follows.

- (A) Compute the overall estimate at a time point t , using all N data points, and using a "continuity" correction that has the effect of removing the effect of a zero in the logarithm (see D.R. Cox, Analysis of Binary Data, Methuen Monograph):

$$\ell_N = \ln\left(\frac{\tilde{p}_N(t) + \frac{1}{2N}}{1 - \tilde{p}_N(t) + \frac{1}{2N}}\right)$$

- (B) Compute the ℓ -values by leaving out each data point in turn when computing $\tilde{p}(t)$: for $i=1, 2, \dots, N$.

$$\ell_{N-1,i} = \ln\left(\frac{\tilde{p}_{N-1,-i}(t) + \frac{1}{2(N-1)}}{1 - \tilde{p}_{N-1,-i}(t) + \frac{1}{2(N-1)}}\right)$$

(C) Form the pseudo-values

$$z_i = N\ell_N - (N-1)\ell_{N-1,-i}$$

(D) Compute \bar{z} , S_z^2

(E) Put approximate confidence $(1-\alpha) \cdot 100\%$ limits on $E[\ell]$ as follows

$$L \leq E[\ell] \leq H$$

$$\text{where } H(L) = \bar{z} + (-) t_{1-\alpha} (N-1) \sqrt{\frac{S_z^2}{N}}$$

(F) Transform back to obtain

$$\frac{e^L}{1+e^L}, \text{ and } \frac{e^H}{1+e^H}$$

The true value, $p(t)$, should be enclosed between these levels for roughly $(1-\alpha) \cdot 100\%$ of all samples. The coverage properties of this procedure will now be checked by simulation: successive samples of size N will be selected, the jackknife limits H and L will be computed for each, and a check will be made as to whether $\frac{e^L}{1+e^L} \leq p(t) \leq \frac{e^H}{1+e^H}$ or not. Tables illustrating performance are given subsequently.

Let

$\ell_{N-1,-i}$ is the logistic transformation estimator from the sample n of the X_i 's with the i th value deleted from the sample.

$$\ell_{N-1,-i} = \ln \left(\frac{\tilde{p}_{N-1,-i}(t) + \frac{1}{2(N-1)}}{1 - \tilde{p}_{N-1,-i}(t) + \frac{1}{2(N-1)}} \right)$$

$\ell_{N-1,-i}$

i	1	2	3	4	5
t					
t_1	3.04	0.98	0.98	0.98	0.98
t_2	3.04	0.98	0.98	0.98	0.98
t_3	0.63	0	0.98	-0.46	-0.46
t_4	0.63	0	0.98	-0.46	-0.46
t_5		-3.04	-3.04	-3.04	-1.89

$$z_i = N\ell_N - (N-1)\ell_{N-1,-i}$$

$$= N \ln \left(\frac{\tilde{p}_N(t) + \frac{1}{2N}}{1 - \tilde{p}_N(t) + \frac{1}{2N}} \right) - (N-1) \ln \left(\frac{\tilde{p}_{N-1,-i}(t) + \frac{1}{2(N-1)}}{1 - \tilde{p}_{N-1,-i}(t) + \frac{1}{2(N-1)}} \right)$$

$z_i(N)$ are called PSEUDO-values of logistic transformation, the following values are calculated:

z_i :

i	1	2	3	4	5
t					
t_1	-6.05	2.198	2.198	2.198	2.198
t_2	-6.05	2.198	2.198	2.198	2.198
t_3	-1.9	0.606	-3.314	2.446	2.446
t_4	-1.9	0.606	-3.314	2.446	2.446
t_5	-3.0626	-3.0626	-3.0626	-3.0626	-7.162

Average of the pseudo-values

$$\bar{z} = \frac{1}{N} \sum_{i=1}^N z_i$$

Invert to find jackknife estimator of logistic transformation

$$\bar{z} = \ln \left(\frac{\tilde{p}(t) + \frac{1}{2N}}{1 - \tilde{p}(t) + \frac{1}{2N}} \right)$$

$$\tilde{p}(t) = \frac{(1 + \frac{1}{2N}) e^{\bar{z}} - \frac{1}{2N}}{1 + e^{\bar{z}}}$$

called the jackknife estimator
of logistic transformation

Variance of the z_i

$$s_z^2 = \text{Var}(z) = \frac{1}{n-1} \sum_{i=1}^n z_i - \bar{z}$$

The following values are calculated:

t	\bar{z}	$\tilde{p}(t)$	Var J
t_1	0.5484	0.646	13.6
t_2	0.5484	0.646	13.6
t_3	0.0568	0.516	6.727
t_4	0.0568	0.516	6.727
t_5	-3.882	0	3.361

The jackknife estimator for estimating variability and giving confidence interval.

Tukey, reference (3) has suggested that in the jackknife procedure we consider the pseudo values $z_i(N)$ as approximately independent and identically distributed and consequently, since \bar{z} is an average of

the $Z_i(N)$, proceed as if

$$\frac{N^{\frac{1}{2}} \bar{z} - \ell_N}{\left\{ \frac{1}{N-1} \sum_{i=1}^n (z_i - \bar{z})^2 \right\}^{\frac{1}{2}}}$$

has t-distribution with $N-1$ d.F.

If the z_i are approximately normal variates (Miller has shown) confidence bands for the unknown $\tilde{p}(t)$ are given, as for the mean of any normal variate when estimated from sample size n .

$$\bar{z} \pm \frac{s_z}{\sqrt{n}} t_{1-\alpha/2}^{(N-1)} \quad (D-1)$$

i.e.

$$\bar{z} - \frac{s_z}{\sqrt{n}} t_{1-\alpha/2}^{(N-1)} \leq \ell_n \left(\frac{\tilde{p}(t) + \frac{1}{2N}}{1 - \tilde{p}(t) + \frac{1}{2N}} \right) \leq \bar{z} + \frac{s_z}{\sqrt{n}} t_{1-\alpha/2}^{(N-1)}$$

$$\bar{L}(n) = \bar{z} - \frac{s_z}{\sqrt{N}} t_{1-\alpha/2}$$

$$\bar{L}(n) = \bar{z} + \frac{s_z}{\sqrt{N}} t_{1-\alpha/2}$$

$$\frac{(1 + \frac{1}{2N})e^{\bar{L}(N)} - \frac{1}{2N}}{1 + e^{\bar{L}(N)}} \leq \tilde{p}(t) \leq \frac{(1 + \frac{1}{2N})e^{\bar{L}(N)} - \frac{1}{2N}}{1 + e^{\bar{L}(N)}}$$

The following values are calculated:

t	4	$t_{1-\alpha/2} = 2.776$
	Lower Int.	Upper Int.
t_1	0	1.0
t_2	0	1.0
t_3	0	1.0
t_4	0	1.0
t_5	0	0.14

The basis for this leap of the imagination seems to be that if $\bar{z} = \bar{x} = \bar{x}_n$ then the procedure for obtaining confidence intervals using equation (D-1) and pseudo-values is the same as the procedure using jackknife. Then if $\bar{x}_N = \bar{z}$ and

$$\bar{z} = \frac{1}{n} \sum_{i=1}^n z_i \quad \text{we have}$$

$$\begin{aligned} z_i &= N \ell_N - (n-1) \ell_{N-1, -i} \\ &= N \bar{x}_N - (N-1) \frac{\left\{ \sum_{j=1}^N x_j \right\} - x_i}{N-1} \\ &= \sum_{j=1}^N x_j - \left[\sum_{j=1}^N x_j \right] + x_i = x_i \end{aligned}$$

Thus the pseudo value

$$z_i = x_i \quad \text{and} \quad \bar{z} = \frac{1}{n} \sum_{i=1}^n x_i = \bar{x}_n$$

The pseudo values are independent if $\bar{z} = \bar{x}_n$ and they are normal if x_i is normal.

E. PARAMETRIC ESTIMATOR, " $\tilde{p}_7(t)$ "

This paper considers one additional estimator, denoted $\tilde{p}_7(t)$. It is a parametric estimator. Therefore, it is not really a competitor to the thirteen non-parametric estimators considered here. In general, a parametric estimator would not be used if the functional form were regarded as unknown. Similarly, a non-parametric estimator would not

normally be used if the survival function were strongly suspected to have a specified form.

$\tilde{p}_7(t)$ is the well known maximum likelihood estimator for the exponential distribution:

$$\tilde{p}_7(t) = e^{-t/\tau}$$

where
$$\tau = \frac{\sum t_i}{\text{number of observed death}}$$

In our sample data base, the total observed survival time is 19, and three deaths are observed. Thus,

$$\sum t_i = 1 + 2 + 3 + 6 + 7 = 19$$

$$\tau = \frac{19}{3}$$

and
$$\tilde{p}_7(t) = e^{-3t/19}$$

Calculations for selected times of interest yield the following estimates:

$$p_7(0) = 1.0$$

$$p_7(1) = 0.854$$

$$p_7(3) = 0.623$$

$$p_7(7) = 0.331$$

The thirteen non-parametric estimators are compared for a variety of generating distributions for both the death mechanism and censoring mechanism.

IV. INSTRUCTIONS FOR USING PROGRAM

INPUT

Each input card bears nine variables. The distribution of time of death is entered in the first set of (five) columns, the censoring distribution is entered in the second set of (ten) columns, a parameter of the censoring distribution is entered in the third set of (ten) columns, the number of replication is entered in the fourth set of (five) columns, the number of the event is entered in the fifth set of (five) columns. For the purpose of all print output used code "0" and "1" in the sixth set of (five) columns, the seed number is entered in the seventh set of (five) columns, after the card giving the time of the last event of a data set, a card with "0" or "1" in the column 50 is inserted, i.e., the "0" indicating more data sets to follow and "1" indicating the last data sets and t value is entered in the ninth set of (eight) columns.

The distribution of time of death and of censoring time used code as follows:

<u>Code</u>	<u>Type of Distribution</u>
1	Uniform
2	Exponential
3	Delta function

OUTPUT

The output lists:

- 1) the time of each observed failure
- 2) estimated survival probability at that time
- 3) the variance of that estimator
- 4) result of goodness fit

- a) mean error
- b) mean absolute error (ABS)
- c) root-mean-square error (RMS)
- 5) total number of observed death
- 6) confidence interval at particular time

Definition of Fortran Variables

NDIE : the distribution of time of death

NTRUNC : the distribution of censoring time

XTRUNC : the parameter of the distribution of censoring time

NREPL : number of replication

NEVENT : number of event

NWRITE : write all output or partial output of simulation

NEND : indicate more data sets or last data set

TN : t statistic value

P_1 : the estimator, $\tilde{p}_1(t)$

P_2 : the estimator, $\tilde{p}_2(t)$

P_3 : the estimator, $\tilde{p}_3(t)$

P_4 : the estimator, $\tilde{p}_4(t)$

P_5 : the estimator, $\tilde{p}_5(t)$

P_6 : the estimator, $\tilde{p}_6(t)$

P_7 : parametric estimator, $\tilde{p}_7(t)$

P_8 : jackknife estimator of logistic transformation of $\tilde{p}_4(t)$

P_9 : jackknife estimator of logistic transformation of $\tilde{p}_5(t)$

P_{10} : jackknife estimator of logistic transformation of $\tilde{p}_6(t)$

P_{11} : Bayesian estimator of $\tilde{p}_1(t)$

P_{12} : Bayesian estimator of $\tilde{p}_2(t)$

P_{13} : Bayesian estimator of $\tilde{p}_3(t)$

p_{14} : jackknife estimator of logistic transformation of $\tilde{p}_2(t)$
 $SL(I,J)$: PSEUDO-value
 $SBAF$: average of pseudo-value
 Var : variance of estimator, $\tilde{p}(t)$
 Var_J : variance of jackknife estimator
 $u(I,J)$: mean of goodness fit
 $w(I,J)$: absolute mean of goodness fit
 $s(I,J)$: root mean square error
 C_1 : upper confidence interval of $p_{14}(t)$
 C_2 : lower confidence interval of $p_{14}(t)$
 C_3 : upper confidence interval of $p_8(t)$
 C_4 : lower confidence interval of $p_8(t)$
 C_5 : upper confidence interval of $p_9(t)$
 C_6 : lower confidence interval of $p_9(t)$
 C_7 : upper confidence interval of $p_{10}(t)$
 C_8 : lower confidence interval of $p_{10}(t)$

To compare RMS with product limit ($p_2(t)$) and jackknife estimator of logistic transformation ($p_{14}(t)$)

2	1	4.0000	1000	20	1	505	: Input				
1	1	C.01275	C.95000	C.93726	: Output						
1	1	C.01581	0.0	C.00000							
1	1	C.03085	C.85722	C.88376							
1	1	C.05450	C.84444	0.83159							
1	1	C.11825	C.75167	C.78256							
1	1	C.12671	C.73885	C.73445							
1	0	C.17874	0.0	0.05254							
1	1	C.18582	0.0	C.00000							
1	1	C.19034	C.67731	0.68724							
1	1	C.19818	C.61574	C.63201							
1	1	C.27670	0.55417	0.57755							
1	1	C.28000	C.45255	C.52356							
1	0	C.31870	0.0	0.00000							
1	1	C.48075	0.42222	0.47045							
1	1	C.65596	C.35185	0.40828							
1	1	C.70112	0.28148	0.34624							
1	1	C.77885	C.21111	0.28413							
1	1	C.18657	C.14074	0.22201							
1	0	C.25712	0.0	0.05254							
1	1	C.47370	0.0	0.05254							
		C.100	C.200	0.300	0.400	0.500	C.600	C.700	0.800	C.900	
P1	-0.002	0.000	0.002	-0.000	-0.001	-0.001	0.007	0.028	0.066	MEAN	
P2	-0.012	-0.007	0.004	0.015	0.029	0.044	0.065	0.109	0.166	MEAN	
P1	0.052	C.071	0.084	0.090	0.097	0.093	0.091	0.077	0.078	ABS	
P2	0.057	C.068	0.075	0.080	0.087	0.087	0.094	0.116	0.166	ABS	
P1	0.065	C.091	0.105	0.113	0.119	0.115	C.113	0.100	0.102	RMS	
P2	0.071	0.085	0.094	0.101	0.108	0.108	0.120	0.140	0.185	RMS	
C1	C.985	C.951	C.501	0.841	0.781	0.722	0.681	0.678	0.751	CCNF	
C2	C.787	C.551	0.459	0.355	0.267	0.185	0.116	0.057	0.035	CCNF	
	57.726	91.254	97.085	97.959	93.542	99.125	97.959	98.542	97.668	PER	
	1000	1000	1000	1000	998	990	951	796	343		

2	2	4.0000	1000	10	1	1509	: Input				
1	1	C.03888	0.0	C.00000	: Output						
1	1	C.14510	C.88885	C.90756							
1	0	C.22306	0.0	0.00000							
1	1	C.22401	C.76190	0.78995							
1	0	C.30447	0.0	C.00000							
1	1	C.48355	0.60552	0.67485							
1	1	C.71228	0.45714	C.54479							
1	1	C.79300	0.30476	0.42580							
1	1	C.14695	0.15238	C.36025							
1	1	C.27255	C.0	0.06065							
		C.100	C.200	0.300	0.400	0.500	0.600	0.700	0.800	C.900	
P1	-0.002	-0.002	-0.005	-0.001	0.006	0.018	0.031	0.062	0.111	MEAN	
P2	0.005	C.013	0.020	0.041	0.067	0.100	0.142	0.200	0.274	MEAN	
P1	0.073	C.100	0.116	0.129	0.131	0.128	0.115	0.107	C.111	ABS	
P2	0.074	C.096	0.105	0.114	0.119	0.128	0.148	0.200	0.274	ABS	
P1	0.058	C.130	0.149	0.160	0.165	0.160	0.146	0.142	0.148	RMS	
P2	0.053	C.121	0.133	0.142	0.150	0.162	0.182	0.227	0.291	RMS	
C1	0.975	C.953	0.522	0.387	0.259	0.144	0.054	0.096	-0.360	CCNF	
C2	0.840	C.640	0.455	0.336	0.242	0.163	0.092	0.033	-0.015	CCNF	
	52.667	70.933	78.933	91.722	95.733	97.600	97.067	97.067	96.800	PER	
	1000	1000	1000	996	984	944	853	656	275		

Computer output of the fourteen estimators

	2	1	4.0000	1000	20	0	505						
	C.100	C.200	0.300	0.400	0.500	C.600	0.700	0.800	C.900				
P1	-0.0033	-0.0558	-0.0759	-0.1000	-0.1114	-0.1211	-0.1114	-0.0992	-0.0449	MEAN			
P2	-0.0000	0.0000	0.0000	-0.0000	-0.0001	-0.0003	0.0003	0.0118	0.0366	MEAN			
P3	-0.0001	0.0001	0.0001	0.0002	0.0002	0.0002	0.0005	0.0026	0.0067	MEAN			
P4	-0.0055	-0.0073	-0.0086	-0.0099	-0.1006	-0.1006	-0.0054	-0.0070	-0.0209	MEAN			
P5	-0.0021	-0.0015	-0.0010	-0.0007	-0.0003	-0.0001	0.0007	0.0017	0.0040	MEAN			
P6	-0.0004	-0.0014	-0.0008	-0.0004	0.0001	0.0006	0.0018	0.0037	0.0077	MEAN			
P7	-0.0000	-0.0007	-0.0008	-0.0008	-0.0007	-0.0005	0.0000	0.0010	0.0033	MEAN			
P8	-0.0045	-0.0065	-0.0085	-0.0099	-0.1007	-0.1009	-0.1000	-0.0800	-0.0339	MEAN			
P9	-0.0011	-0.0013	-0.0009	-0.0005	0.0001	0.0006	0.0015	0.0025	0.0031	MEAN			
P10	-0.0077	-0.0087	-0.0093	-0.0004	0.0002	0.0007	0.0018	0.0032	0.0060	MEAN			
P11	-0.0038	-0.0027	-0.0016	-0.0008	0.0001	0.0011	0.0028	0.0054	0.0103	MEAN			
P12	-0.0038	-0.0026	-0.0015	-0.0006	0.0004	0.0015	0.0033	0.0061	0.0111	MEAN			
P13	-0.0038	-0.0026	-0.0015	-0.0006	0.0004	0.0015	0.0033	0.0061	0.0111	MEAN			
P14	0.0011	0.0014	0.0020	0.0023	0.0029	0.0039	0.0064	0.0117	0.0222	MEAN			
P1	C.074	0.100	0.118	0.133	0.142	C.141	0.131	0.104	C.054	ABS			
P2	C.053	0.071	0.084	0.090	0.097	0.093	0.092	0.077	C.070	ABS			
P3	C.053	0.071	0.084	0.090	0.097	0.093	0.092	0.080	C.079	ABS			
P4	C.053	0.066	0.078	0.083	0.085	0.087	0.085	0.076	C.066	ABS			
P5	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P6	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P7	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P8	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P9	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P10	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P11	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P12	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P13	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P14	C.053	0.066	0.078	0.083	0.085	0.086	0.083	0.074	C.066	ABS			
P1	C.096	0.129	C.150	0.165	0.173	0.168	0.153	0.119	0.068	RMS			
P2	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P3	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P4	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P5	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P6	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P7	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P8	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P9	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P10	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P11	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P12	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P13	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
P14	C.065	0.091	0.105	0.113	0.119	0.116	0.114	0.099	C.074	RMS			
C1	C.568	C.932	0.833	0.826	0.768	0.717	0.657	0.741	0.656	CCNF			
C2	C.761	C.570	0.452	0.357	0.273	0.196	0.129	0.071	0.024	CCNF			
C3	C.561	C.513	0.350	0.276	0.206	0.141	0.079	0.020	0.018	CCNF			
C4	C.707	C.486	0.360	0.267	0.191	0.127	0.079	0.045	C.326	CCNF			
C5	C.567	C.525	0.375	0.280	0.206	0.141	0.079	0.045	C.326	CCNF			
C6	C.760	C.570	0.452	0.357	0.273	0.196	0.129	0.071	0.024	CCNF			
C7	C.561	C.513	0.350	0.276	0.206	0.141	0.079	0.020	0.018	CCNF			
C8	C.760	C.569	0.453	0.361	0.282	C.209	0.146	0.096	0.055	CCNF			
	1000	1000	1000	1000	1000	1000	994	933	556				

V. RESULTS OF THE SIMULATION

This paragraph presents graphical comparisons of the 13 estimators based on simulated data. Each comparison is based on 1000 replications of a simulated data base. The bias and RMS error (square-root of mean-squared error) of each estimator depends on the parameters that control the simulated data base. No single estimator dominates all others under all conditions.

The bias and RMS errors of the estimators depend on several factors:

(A) The sample size (NEVENT) of individuals under observation at time zero affects the accuracy of the estimators. In general, a larger sample size leads to a better estimate than a smaller sample. Values of NEVENT selected for simulation are 5, 10, 25, and 50 (plus one simulation with NEVENT = 100).

(B) The distribution of times at which the observations are censored (unless the individual dies earlier) affects the performance of the various estimators. This distribution is particularly important in conjunction with the distribution of lifetimes (do most individuals die before censoring is likely?, are deaths and censoring events about equally likely at all times?, are most observations censored before death?). Three types of distributions are assumed to underlie the censoring mechanism:

- (1) Some of the samples are generated on the assumption that no censoring occurs.
- (2) Some samples are generated from a uniform distribution of times of censoring.

(3) Other data bases are generated from an exponential distribution of censoring times.

(C) The distribution of lifetimes (ignoring the possibility of censoring) also affects the performance of the various estimators. Two types of distributions are assumed to underlie the death mechanism:

(1) Some of the samples are generated from a uniform distribution of lifetimes.

(2) Other data bases are generated from an exponential distribution of lifetimes.

If a uniform distribution of lifetimes is selected, its range is always over the interval from time 0 to time 1. If an exponential distribution is selected, it always has a mean lifetime of 1. The distributions of truncation times (uniform or exponential) have parameters .25, .5, .667, .75, 1, 1.333, 1.5, 2 and 4. A wide variety of samples may be simulated by mixing various pairs of distributions (for censoring times and deaths). Since the time units are arbitrary, the restriction on mean lifetimes is irrelevant.

The true value of the survival function is, $p(t)$, and the form of this function affects the relative performance of the 13 nonparametric estimators. For example, the Bayesian estimator $p_{12}(t)$ tends to be better as measured by square-root of mean-squared error than its counterpart (the product-limit estimator, $p_2(t)$) for the time frame in which

$$.3 < p(t) < .9$$

However, the product limit estimator tends to be better for those times when $p(t)$ is close to zero or unity.

The point estimators, $p_5(t)$ and $p_6(t)$ tend to be better than the product-limit estimator ($p_2(t)$) for all time periods. The jackknife

estimators of logistic transformation ($p_8(t)$, $p_9(t)$, $p_{10}(t)$) of point estimators tends as same as its counterpart point estimators ($p_4(t)$, $p_5(t)$, $p_6(t)$) for all time periods. And also the estimator formed by jackknifing the logistic transformation ($p_{14}(t)$) of the product limit estimator tends to be better than its counterpart product limit ($p_2(t)$) for the time frame in which

$$.1 < p(t) < .7$$

However, the product limit estimator tends to be better for those times when $p(t)$ is close to unity. Point estimators, $p_5(t)$ and $p_6(t)$ tend to be same for the time frame in which

$$0.1 < p_T(t) < 0.9$$

However, the $p_5(t)$ tends to be better for those times when $p_T(t)$ is close to unity. The jackknife procedure may be validated, in an empirical sense, by sampling experiments or computer simulation in the following manner. First, times of censoring and death are obtained by drawing random numbers from postulated distributions. Second, the jackknifed estimator of the logistic-transformed product-limit estimation is found, and confidence limits are computed by the method of Tukey, reference (3). Since the true value of survival function, $p(t)$, is known, so is the theoretical value of A . The jackknife confidence intervals can be checked for coverage: if $L_\alpha \leq A \leq H_\alpha$ then the particular interval covers, while otherwise (if $A < L_\alpha$ or $H_\alpha < A$) it does not cover. Finally, the above procedure can be repeated many times (say 1000) and the fraction of repetitions which contains the true value of A is recorded. This fraction of the coverage should desirably be close to $(1-\alpha)$, the nominal confidence level.

The jackknife confidence limit procedure can be said to be robust of validity, ref (7), if the actual coverage is close to the nominal coverage, $1-\alpha$, for a various distributions. Such seems to be true for large n ($n \geq 50$). However, the jackknife confidence limits do not cover accurately when the true value of $p(t)$ is close to unity.

The following tables illustrate confidence limits of jackknife method of product limit ($p_2(t)$). Many computer generated graphics are presented on the following pages to complete this section.

Table 1

Simulation Experiments Validating Table
95% Confidence Limits (t value = 2.776)

95% Confidence Limits (t value = 2.776)											
True Value of Death		Distinguishing Trunc. Parameter of Censoring	0.9	0.8	0.7	0.6	0.5	0.4	0.3	0.2	0.1
			Upper Int.	Lower Int.	Coverage	Upper Int.	Lower Int.	Coverage	Upper Int.	Lower Int.	Coverage
Exponential	Exponential	2.0	0.962	0.935	0.917	0.912	0.922	0.945	0.971	0.995	1.0
			0.709	0.515	0.366	0.259	0.173	0.115	0.068	0.034	0.009
			56.735	66.122	75.408	85.714	90.816	96.939	98.980	95.918	96.939
Uniform	Exponential	1.3333	0.962	0.935	0.918	0.914	0.923	0.942	0.969	0.992	1.0
			0.741	0.559	0.422	0.313	0.238	0.166	0.10	0.059	0.068
			57.209	65.116	72.093	79.070	86.047	93.029	95.023	95.023	86.047
Uniform	Exponential	4.0	0.955	0.910	0.871	0.846	0.825	0.819	0.827	0.837	0.855
			0.744	0.552	0.407	0.301	0.217	0.163	0.113	0.078	0.047
			50.792	68.358	74.194	85.630	92.669	96.774	92.150	95.308	88.886
Exponential	Uniform	0.6667	0.957	0.923	0.901	0.893	0.898	0.904	0.911	0.915	0.932
			0.775	0.618	0.494	0.393	0.324	0.276	0.202	0.166	0.025
			54.545	65.545	73.636	82.727	92.273	94.091	86.364	87.27	72.787
Exponential	Uniform	2.0	0.964	0.931	0.911	0.901	0.906	0.917	0.940	0.963	**
			0.760	0.558	0.419	0.305	0.231	0.171	0.125	0.122	**
			55.887	56.259	72.340	82.496	87.943	91.489	85.668	87.943	**
Uniform	Exponential	1.3333	0.957	0.924	0.902	0.891	0.897	0.914	0.979	**	**
			0.757	0.580	0.456	0.380	0.305	0.256	0.247	**	**
			45.887	54.255	72.340	85.496	88.15	93.52	96.688	**	**
Uniform	Uniform	4.0	0.948	0.906	0.862	0.836	0.821	0.816	0.821	0.837	0.863
			0.718	0.533	0.385	0.291	0.216	0.155	0.112	0.080	0.049
			53.117	61.039	68.571	87.987	93.506	96.429	91.558	95.130	89.844
Uniform	Uniform	1.0	0.953	0.917	0.893	0.882	0.883	0.891	0.907	0.924	0.938
			0.751	0.60	0.496	0.399	0.318	0.276	0.251	0.131	0.049
			58.0	70.0	78.333	78.667	78.667	80.0	70.0	73.333	66.667

Table 2

Simulation Experiments Validating Table
95% Confidence Limits (t value = 2.262)

95% Confidence Limits (t value = 2.262)												
n=10	True Value		0.9	0.8	0.7	0.6	0.5	0.4	0.3	0.2	0.1	
	Dist. of Death	of Censoring										
Expo- nential	Expo- nential	4.0	Upper Int.	0.975	0.953	0.922	0.887	0.859	0.844	0.854	0.896	0.966
			Lower Int.	0.840	0.640	0.455	0.336	0.242	0.163	0.092	0.033	0.0
			Coverage	52.667	70.933	78.933	91.733	95.733	97.60	97.067	97.067	96.8
Uni- form	Expo- nential	1.0	Upper Int.	0.966	0.930	0.891	0.861	0.851	0.858	0.892	0.938	0.976
			Lower Int.	0.582	0.359	0.241	0.176	0.125	0.084	0.058	0.032	0.015
			Coverage	62.0	87.50	94.643	98.214	100.0	100.0	100.0	98.214	98.214
Uni- form	Expo- nential	1.0	Upper Int.	0.954	0.899	0.843	0.792	0.753	0.730	0.731	0.745	0.760
			Lower Int.	0.612	0.410	0.305	0.250	0.204	0.160	0.120	0.083	0.060
			Coverage	57.798	77.982	89.297	96.330	99.083	98.165	99.083	96.024	76.453
Expo- nential	Uni- form	2.0	Upper Int.	0.956	0.897	0.834	0.772	0.716	0.673	0.646	0.637	0.649
			Lower Int.	0.615	0.391	0.286	0.227	0.138	0.151	0.120	0.089	0.058
			Coverage	53.542	78.125	84.167	94.167	98.333	98.542	99.167	97.917	82.083
Expo- nential	Uni- form	4.0	Upper Int.	0.975	0.948	0.913	0.874	0.840	0.821	0.828	0.865	0.937
			Lower Int.	0.546	0.295	0.165	0.100	0.068	0.048	0.030	0.012	0.003
			Coverage	60.0	86.154	96.923	98.174	100.0	100.0	100.0	100.0	100.0
Uni- form	Uni- form	2.0	Upper Int.	0.940	0.896	0.855	0.814	0.787	0.785	0.806	0.851	**
			Lower Int.	0.860	0.709	0.578	0.468	0.379	0.308	0.251	0.168	**
			Coverage	57.6	51.2	78.933	91.733	96.0	98.133	97.333	96.8	**
Uni- form	Uni- form	1.0	Upper Int.	0.949	0.896	0.837	0.794	0.767	0.761	0.779	0.800	0.840
			Lower Int.	0.596	0.423	0.324	0.265	0.225	0.182	0.150	0.130	0.097
			Coverage	59.091	79.545	93.182	93.182	100.0	97.727	97.727	72.727	56.818
Uni- form	Uni- form	4.0	Upper Int.	0.957	0.902	0.836	0.767	0.703	0.647	0.603	0.583	0.589
			Lower Int.	0.602	0.383	0.272	0.220	0.184	0.153	0.121	0.089	0.055
			Coverage	57.093	79.159	87.566	94.921	99.124	100.0	100.0	98.949	86.165

Table 3

Simulation Experiments Validating Table
95% Confidence Limits (t value = 2.093)

Dist. of Die		True Value		0.9		0.8		0.7		0.6		0.5		0.4		0.3		0.2		0.1	
Die		of Die		Trunc. Parameter		distinguish		Upper Int.		Lower Int.		Coverage		Upper Int.		Lower Int.		Coverage		Upper Int.	
Exponential	Exponential	2.0	0.986	0.787	0.603	0.477	0.893	0.833	0.774	0.717	0.682	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805
Exponential	Exponential	1.333	0.941	0.897	0.848	0.778	0.893	0.833	0.774	0.717	0.682	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805
Exponential	Exponential	0.667	0.967	0.910	0.851	0.794	0.893	0.833	0.774	0.717	0.682	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805
Exponential	Exponential	4.0	0.948	0.911	0.864	0.809	0.778	0.730	0.675	0.643	0.715	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805
Exponential	Exponential	1.333	0.922	0.860	0.792	0.730	0.675	0.643	0.715	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805	0.798	0.805
Exponential	Exponential	2.00	0.951	0.878	0.799	0.714	0.627	0.541	0.455	0.372	0.309	0.250	0.221	0.186	0.146	0.100	0.054	0.005	0.000	0.000	0.000
Exponential	Exponential	1.00	0.950	0.877	0.796	0.717	0.638	0.559	0.490	0.428	0.372	0.309	0.250	0.221	0.186	0.146	0.100	0.054	0.005	0.000	0.000
Exponential	Exponential	2.00	0.952	0.879	0.797	0.715	0.624	0.541	0.455	0.372	0.309	0.250	0.221	0.186	0.146	0.100	0.054	0.005	0.000	0.000	0.000
Exponential	Exponential	1.00	0.952	0.879	0.797	0.715	0.624	0.541	0.455	0.372	0.309	0.250	0.221	0.186	0.146	0.100	0.054	0.005	0.000	0.000	0.000
Exponential	Exponential	2.00	0.952	0.879	0.797	0.715	0.624	0.541	0.455	0.372	0.309	0.250	0.221	0.186	0.146	0.100	0.054	0.005	0.000	0.000	0.000

Table 4

Simulation Experiments Validating Table
95% Confidence Limits (t value = 2.010)

n = 50

Dist. of Death	Uni- form	True Value	0.9	0.8	0.7	0.6	0.5	0.4	0.3	0.2	0.1
Expo- nential	Uni- form	1.0	Upper Int.	0.938	0.856	0.772	0.692	0.626	0.636	**	**
			Lower Int.	0.300	0.302	0.290	0.268	0.238	0.215	**	**
			Converge	94.650	94.870	95.140	95.520	95.910	96.720	**	**
		1.5	Upper Int.	0.950	0.876	0.799	0.717	0.636	0.566	**	**
			Lower Int.	0.238	0.245	0.244	0.228	0.202	0.168	**	**
			Converge	94.860	95.120	95.210	95.830	95.960	96.120	**	**
	Expo- nential	3.0	Upper Int.	0.968	0.908	0.839	0.766	0.689	0.612	0.541	0.503
			Lower Int.	0.121	0.142	0.153	0.149	0.134	0.111	0.082	0.049
			Converge	94.275	95.237	95.237	95.618	95.820	95.820	96.012	96.947
		0.6667	Upper Int.	0.959	0.892	0.822	0.753	0.689	0.648	0.670	0.756
			Lower Int.	0.201	0.199	0.198	0.182	0.155	0.121	0.083	0.046
			Converge	92.381	94.325	94.875	95.312	95.312	96.012	96.463	96.463
Uni- form	Expo- nential	1.3333	Upper Int.	0.970	0.913	0.848	0.780	0.711	0.646	0.602	0.635
			Lower Int.	0.128	0.136	0.144	0.137	0.121	0.097	0.066	0.034
			Converge	94.568	94.761	95.327	95.327	96.032	96.032	95.882	95.882
		2.0	Upper Int.	0.974	0.920	0.858	0.791	0.721	0.653	0.594	0.585
			Lower Int.	0.103	0.114	0.124	0.120	0.107	0.087	0.061	0.030
			Converge	94.523	94.876	95.038	95.523	95.823	96.720	96.720	96.720
	Uni- form	1.0	Upper Int.	0.937	0.855	0.769	0.681	0.598	0.520	0.465	0.472
			Lower Int.	0.318	0.312	0.301	0.277	0.246	0.209	0.165	0.122
			Converge	94.667	95.222	95.667	95.778	95.778	96.033	96.112	94.778
		4.0	Upper Int.	0.936	0.853	0.766	0.677	0.585	0.490	0.393	0.295
			Lower Int.	0.296	0.303	0.295	0.275	0.247	0.211	0.168	0.118
			Converge	94.826	95.362	96.012	96.723	96.723	97.023	97.023	94.234

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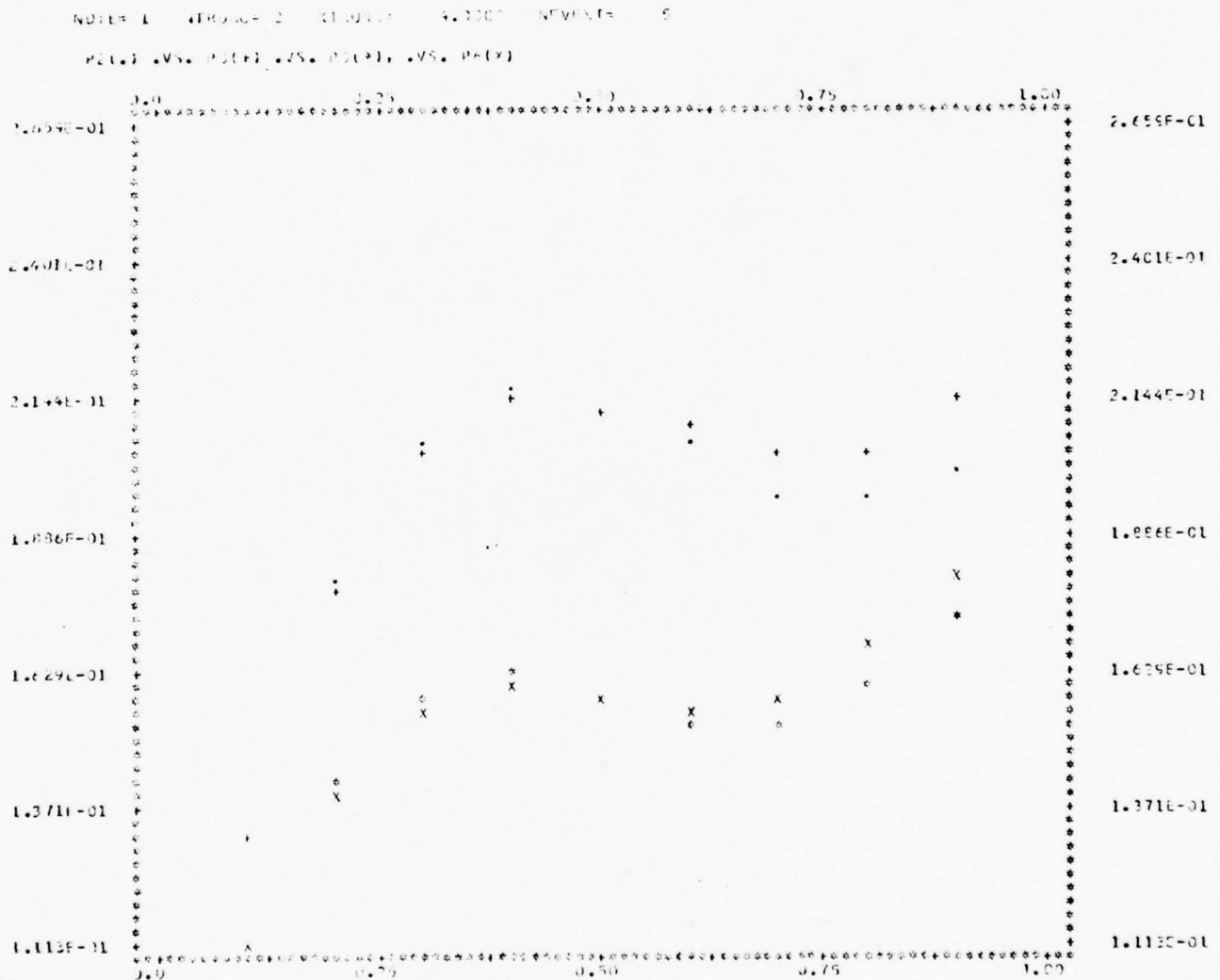


Fig. 1: Comparison of RMS derived from sample size 5.
(Step function estimators vs. point estimators)

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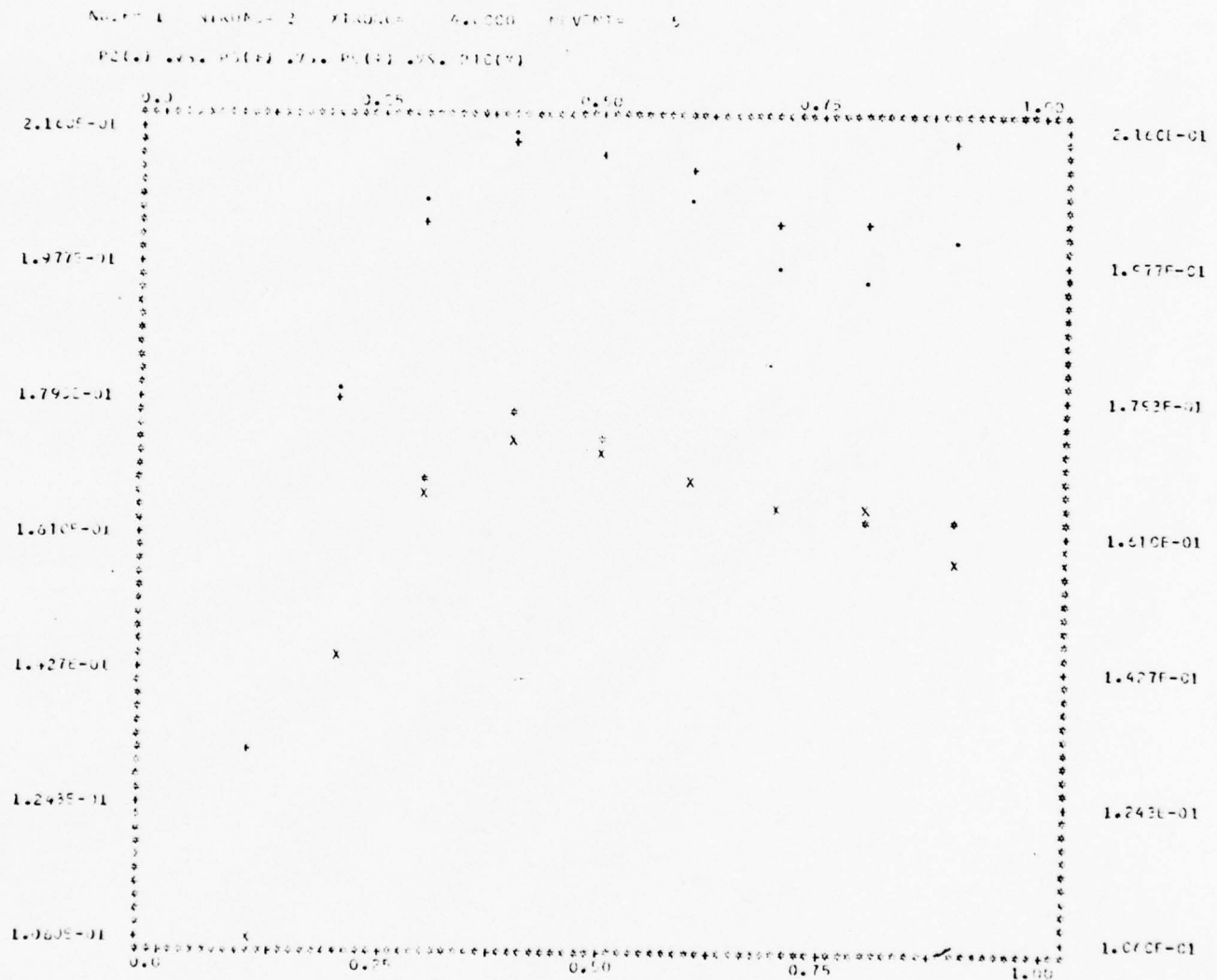


Fig. 2: Comparison of root mean square derived from sample size 5.
(Step function estimators vs. jackknife estimators of
logistic transformation)

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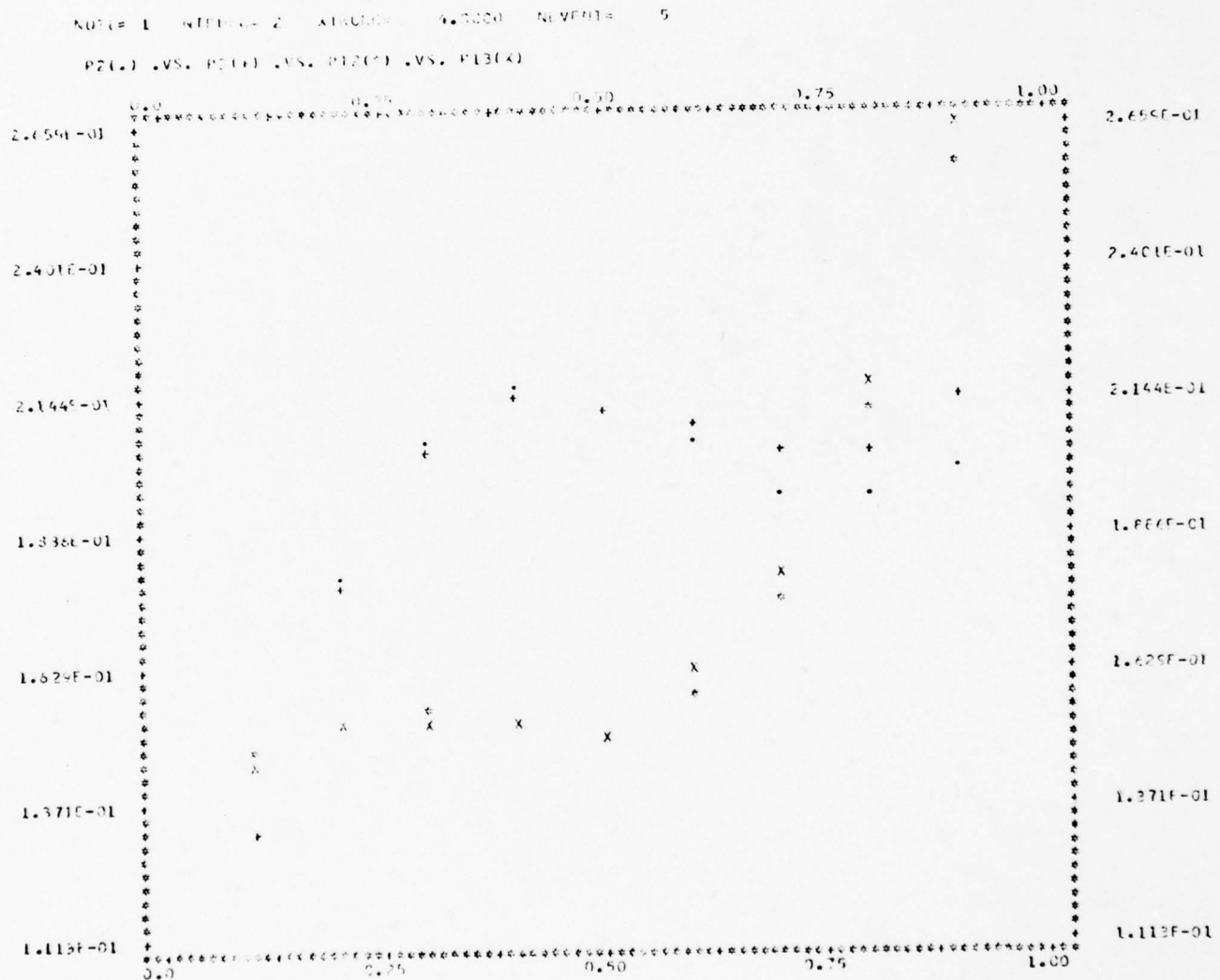


Fig. 3: Comparison of root mean square derived from sample sizes.
(Step function estimators vs. Bayesian estimators)

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NOTE: 1. NUMBER OF X-VALUES 4,000 NUMBER OF Y-VALUES 10

P2(X) VS. P2(X) VS. P2(X) VS. P2(X)

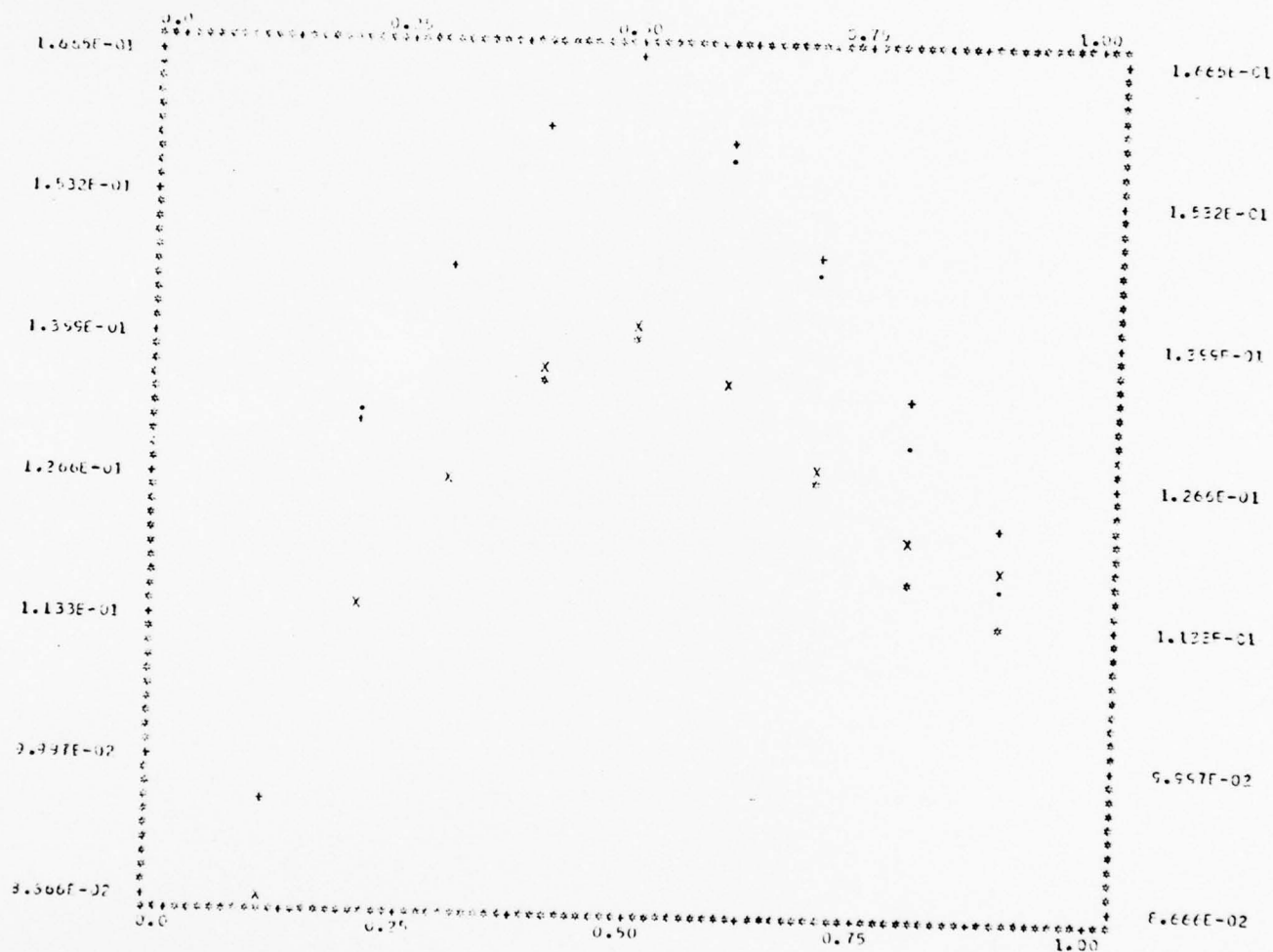


Fig. 4: Comparison of root mean square derived from sample size 10.
(Step function estimators vs. point estimators)

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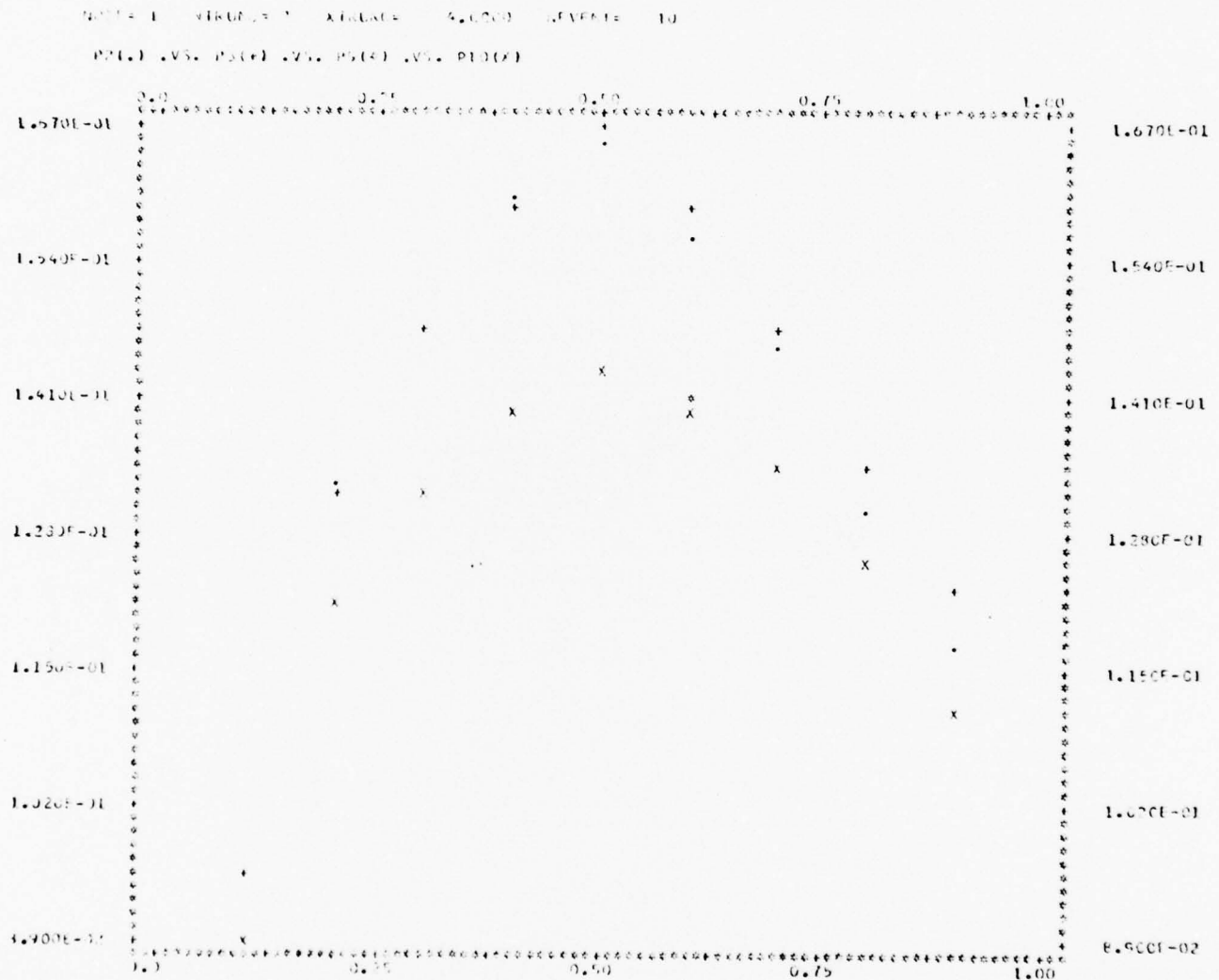


Fig. 5: Comparison of root mean square derived from sample size 10.
(Step function estimators vs. jackknife estimators of
logistic transformation)

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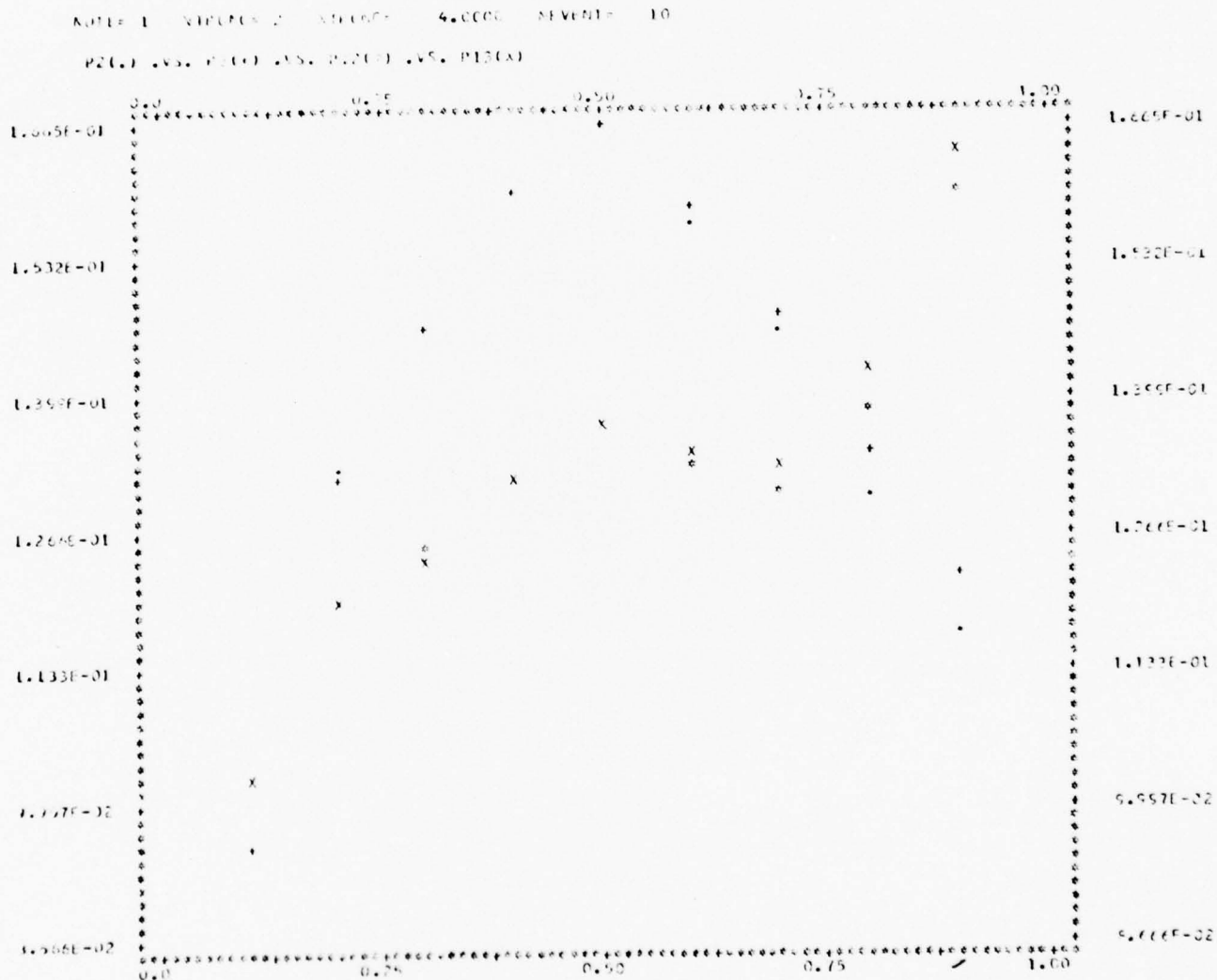


Fig. 6: Comparison of RMS derived from sample size 10.
(Step function estimators vs. Bayesian estimators)

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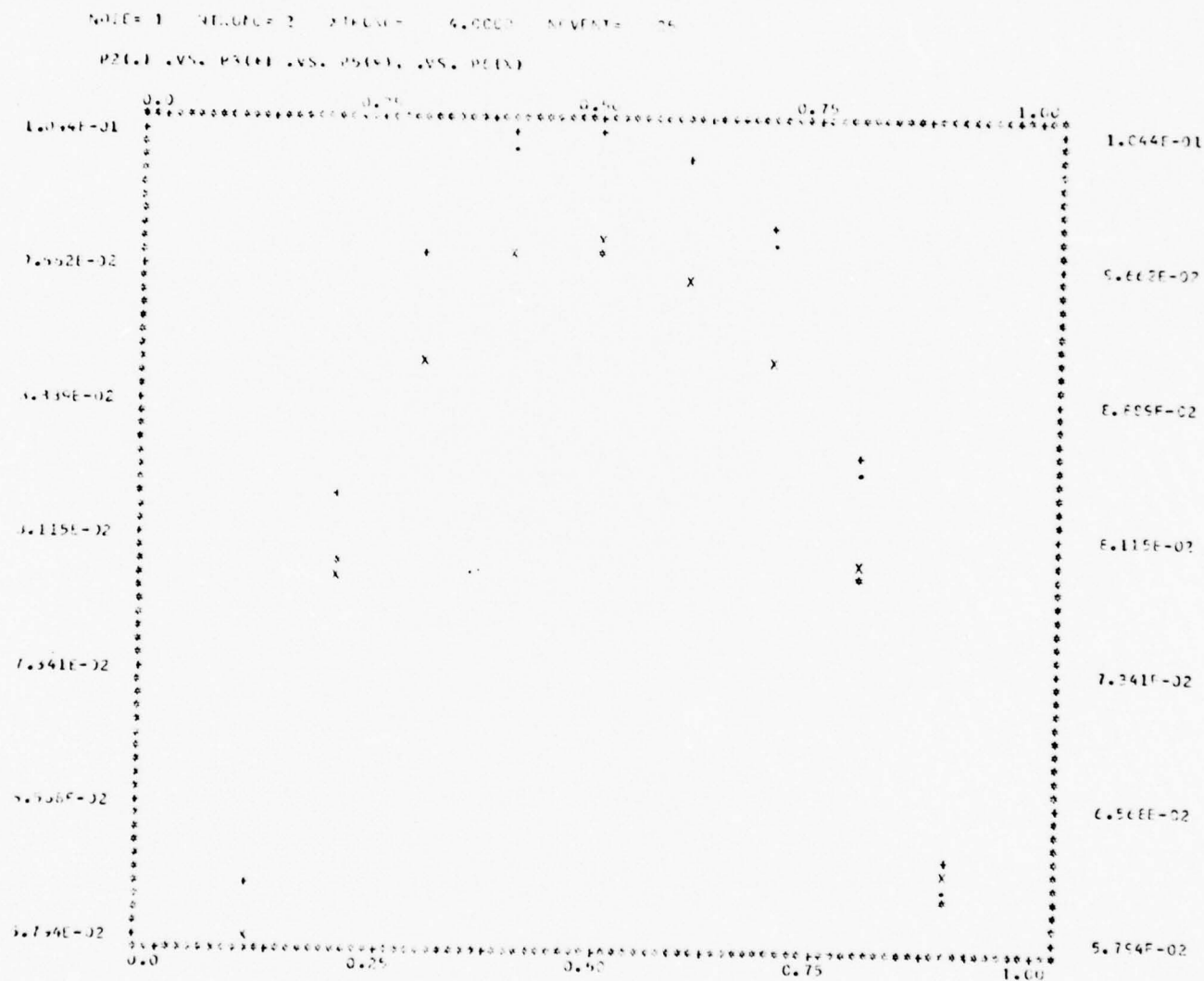


Fig. 7: Comparison of RMS derived from sample size 25.
(Step function estimators vs. point estimators)

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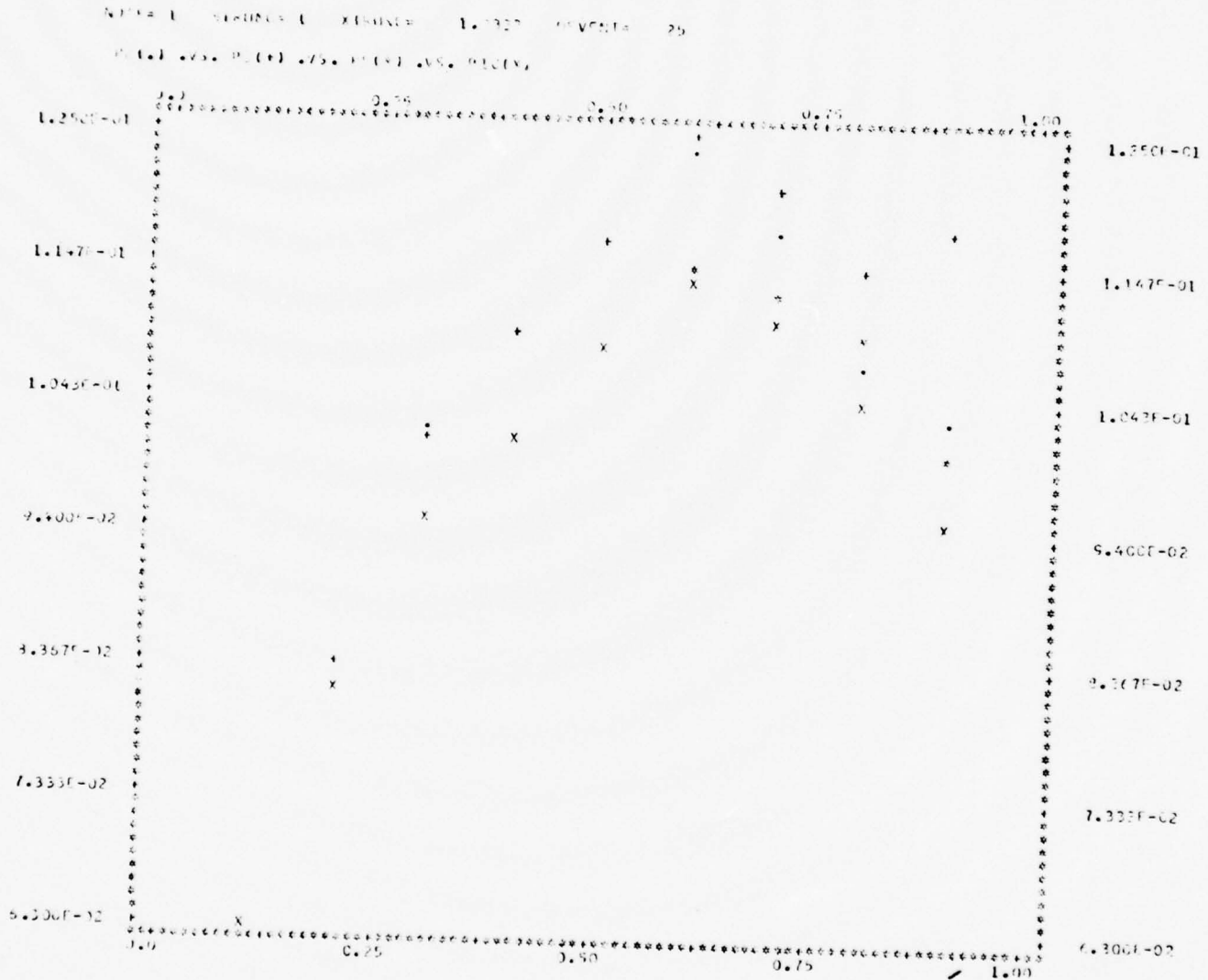


Fig. 8: Comparison of RMS derived from sample size 25.
(Step function estimators vs. jackknife estimators of
logistic transformation)

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NOTE: 1. FIGURE 2. XI. DATES 4.10.11. NINE. 15.

P21(1) .VS. P3(1) .VS. P12(1) .VS. P13(1)

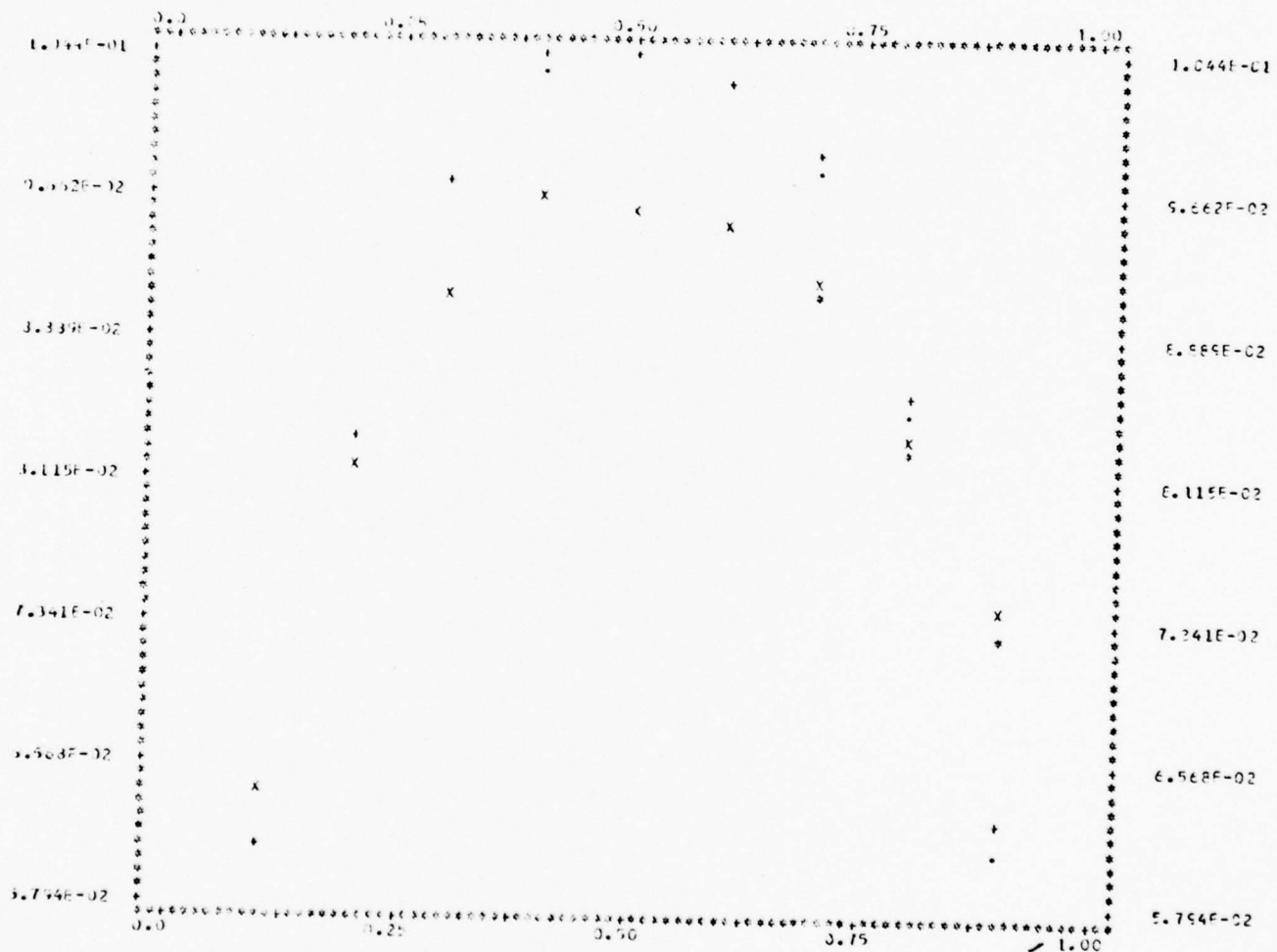


Fig. 9: Comparison of RMS derived from sample size 25.
(Step function estimators vs. Bayesian estimators)

APPENDIX A

ESTIMATORS FOR GROUPED DATA

With grouped censored data the definition of $p(t_i/t_{i-1})$ given by equation (5) does not hold unless the assumption is made that all truncations occur at the end of the time interval. If, on the other hand, it is assumed that all truncations occur at the beginning of Δt_i the equivalent form of equation (5) is

$$p(t_i/t_{i-1}) = \frac{N_i - a_i - r_i}{N_i - a_i} \quad (G-1)$$

With N_i elements were presents at beginning of interval, i.e., at time t_{i-1} , r_i elements failed during the interval, and a_i elements truncated from the sample during the interval but prior to failing. As a hypothesis, assume that all aborts occur simultaneously somewhere within the time interval, so that r' failures occur prior to the truncations and time remaining $r_i - r'$ after the truncations. Then

$$p(t_i/t_{i-1}) = \frac{N_i - r'}{N_i} \cdot \frac{N_i - a_i - r_i}{N_i - r_i - a_i} \quad (G-2)$$

Thus, the value of $p(t_i/t_{i-1})$ depends on when the truncations occur. It is assumed that this is not known for the grouped data case. Nevertheless, it is possible to place limits on the value of $p(t_i/t_{i-1})$ since equation (G-2) always gives values between those of equation (5) and (G-1). Thus

$$\frac{N_i - a_i - r_i}{N_i - a_i} \leq p(t_i/t_{i-1}) \leq \frac{N_i - r_i}{N_i} \quad (G-3)$$

For average sample size approximation, a simpler expression from the point of view of computational ease may be derived by substituting $a/2$ for a in equation (G-1) giving

$$p(t_i) = \frac{N - \frac{a}{2} - r}{N - \frac{a}{2}} \quad (G-4)$$

The equation (G-4) may be thought of as the result of assuming that the average number of elements in the time interval is the number at the beginning decreased by half the number of truncations.

Records are usually available to provide a fairly precise time the death events. In the medical example, the exact time of death is usually recorded in medical records required by law. In the equipment lifetesting example, the time of malfunction or failure is usually known very precisely if the results are catastrophic; and maintenance records give a reasonably precise time even if the failure is not critical to a larger system. In the military example, the event of interest is usually a sensor detection or some other action that is routinely recorded in a log book.

Equation (G-4) is a modification to the product-limit estimator, p_2 , when the times of truncation are known only in grouped form. Herd, reference (2), suggests a similar modification to estimators using the second approach (p_5 or p_6) with aggregated truncation data. Illustrate results for this method based on the sample data base of the main test are given below. Here, of course, we do not know that individual B dropped out of observation at time 2 and that individual D dropped out at time 6. We know only that the two truncations occurred in the interval (1,3) and (3,7), respectively.

Product limit's modification is denoted by $\tilde{p}_2'(t)$ and Herd's modification is denoted by $\tilde{p}_5'(t)$.

Their results on the sample data base are as follows.

<u>t</u>	<u>$\tilde{p}_2'(t)$</u>
0-1	$5/5 = 1.0$
1-3	$4/5 \times 1.0 = 0.8$
3-7	$2.5/3.5 \times 0.8 = 0.571$
(7)	$0.5/1.5 \times 0.571 = 0.190$

<u>t</u>	<u>$\tilde{p}_5'(t)$</u>
0	1.0
1	$5/6 \times 1.0 = 0.833$
3	$3.5/4.5 \times 0.833 = 0.648$
7	$1.5/2.5 \times 0.648 = 0.389$

APPENDIX B

LISTING OF COMPUTER PROGRAM

```

C      TRUNCATED DATA PROGRAM
C
      DIMENSION NN(9),D(14),T(1000),IT(1000),P1(1000),P2(100
*0),P3(1000),P4(1000),P5(1000),P6(1000),PJA4(1000),PJA5
*(1000),PJA6(1000),TJ(1000),ITJ(1000),PZ(1000),S(14,9),
*U(14,9),W(14,9),P11(1000),P12(1000),P13(1000),SL1(50,5
*0),SL2(50,50),SL3(50,50),SBAR1(50),SBAR2(50),SEAR3(50)
*PJ4(50,50),PJ5(50,50),PJ6(50,50),FJ2(50,50),PZ2(1000),
*SL4(50,50),SBAR4(50),PJA2(1000),C(14,9),RMS1(50),RMS2(
*(50),RMS3(50),RMS4(50),UINT1(50),LINT2(50),LINT4(50),U
*INT3(50),RINT1(50),RINT2(50),RINT3(50),RINT4(50),CF(8)
*VARJ2(50),VARJ4(50),VARJ5(50),VARJ6(50),UP1(50),UP2(50
*),UP3(50),UP4(50),RO1(50),RO2(50),RO3(50),RC4(50)
      CALL CVFLOW
C
1  FCRMAT (I5,I10,F10.4,5I5,F8.3)
2  FCRMAT(1X,'NDIE ERROR')
3  FCRMAT(1X,'NTRUNC ERROR')
4  FCRMAT(1X,'NREPL ERROR')
5  FCRMAT(1X,'NEVENT ERROR')
6  FCRMAT(1X,'NWRITE ERROR')
7  FCRMAT (3X,5F8.3)
8  FCRMAT (1X,2I5,14F8.5)
9  FCRMAT (1X,'P',I2,5F8.3,3X,'MEAN')
10 FCRMAT(' ')
11 FCRMAT('1')
12 FCRMAT(1X,I5,10F10.5)
13 FCRMAT (1X,2I5,F10.4,4I5)
14 FCRMAT (1X,'P',I2,9F8.3,3X,'ABS')
15 FCRMAT (5X,9(14,4X))
16 FCRMAT (1X,'P',I2,9F8.3,3X,'RMS')
18 FCRMAT (1X,2I5)
C
C      READ INPUTS AND SET INITIAL VALUES
C
      A=0.01
25 READ(5,1)NDIE,NTRUNC,XTRUNC,NREPL,NEVENT,NWRITE,ISEED,
*NEND,TN
      JEVENT=NEVENT-1
      WRITE(6,13)NDIE,NTRUNC,XTRUNC,NREPL,NEVENT,NWRITE,ISEE
*NEND
      WRITE (6,10)
      NP=14
      SRH=SQRT(.5)
      DO 26 I=1,1000
      T(I)=0.0
      P1(I)=0.0
      P2(I)=0.0
      P3(I)=0.0
      P4(I)=0.0
      P5(I)=0.0
      P6(I)=0.0
      TJ(I)=0.0
      P11(I)=0.0
      P12(I)=0.0
      P13(I)=0.0
26 PZ(I)=0.0
C
C      TEST INPUTS
C
      IF (NDIE.LT.1) GOTO 105
      IF (NDIE.GT.2) GOTO 105
      IF (NTRUNC.LT.1) GOTO 103
      IF (NTRUNC.GT.3) GOTO 103
      IF (NREPL.LT.1) GOTO 104
      IF (NREPL.GT.1000) GOTO 104
      IF (NEVENT.LT.2) GOTO 102
      IF (NEVENT.LE.1000) GOTO 200
C
C      ERROR MESSAGES
C
102 WRITE (6,5)

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```

103 STCP
WRITE (6,3)
STCP
104 WRITE (6,4)
STCP
105 WRITE (6,2)
106 STCP
C
C
C START MAIN CALCULATION
C
200 DC 250 J=1,9
NN(J)=0
DC 250 I=1,NP
S(I,J)=0
U(I,J)=0
C(I,J)=0.0
250 W(I,J)=0
DC 4999 IREPL=1,NREPL
NDI=0
DC 999 IEVENT=1,NEVENT
C
C
C CREATE TTRUNC()
C
CALL RANDOM(ISEED,TTR,1)
GOTO (300,350,400),NTRUNC
GOTO 103
300 TTR=TTR*XTRUNC
GOTO 500
350 TTR=-XTRUNC*ALOG(TTR)
GOTO 500
400 TTR =XTRUNC
C
C
C CREATE TDIE()
C
500 CALL RANDOM(ISEED,TDI,1)
GOTO (600,700),NDIE
GOTO 102
700 TDI=-ALOG(TDI)
C
C
C DETERMINE SMALLER OF TDIE() AND TTRUNC()
C
800 IF (TDI.LE.TTR) GOTO 810
TT=TTR
ITT=0
GOTO 850
810 TT=TDI
NCI=NDI+1
ITT=1
850 T(IEVENT)=TT
IT(IEVENT)=ITT
C
C
C ORDER DATA
C
870 IF (IEVENT.EQ.1) GOTO 999
II=IEVENT-1
CC 890 I=1,II
IF (TT.GT.T(I)) GOTO 890
III=II-I+1
CC 880 J=1,III
JJ=IEVENT-J
IT(JJ+1)=IT(JJ)
880 T(JJ+1)=T(JJ)
IT(I)=ITT
T(I)=TT
GOTO 999
890 CCNTINUE
C
999 CCNTINUE
TN=T(NEVENT)
T7=0
DII=NDI+2.
IF (NDI.GT.0) GOTO 8122

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      P4(NEVENT)=SRH
      P5(NEVENT)=SRH
      TT=0
      DC 8121 J=1,NEVENT
8121 TT=TT+T(J)
      DN=TT/T(NEVENT)
      P6(NEVENT)=SQRT(DN/(DN+1.))
      GOTO 1111
C
C      CALCULATE P1() AND P4() AND P11() VECTORS AND P7 DATA
C
8122 N=NDI
      J=0
      II=0
      III=0
      DC 2199 I=1,NEVENT
      T7=T7+T(I)
      IF(IT(I).EQ.1) GOTO 2150
      J=1
      GOTO 2199
2150 P1(I)=FLOAT(N-1)/FLOAT(NDI)
      P4(I)=FLOAT(N)/FLOAT(NDI+1)
      P11(I)=FLCAT(N)/DII
      TII=T(I)
      III=II
      II=I
      J=0
      N=N-1
2199 CCNTINUE
      T7=-NDI/T7
      IF (J.EQ.0) GOTO 2221
      TII=T(III)
      CTI=T(III)-TII
      IF (NDI.GT.1) GOTO 2200
      TAU=-TII/ALCG(P4(III))
      TTAU=-TII/ALOG(P11(III))
      GOTO 2210
2200 TAL=DTI/ALCG(P4(III)/P4(III))
      TTAU=(T(III)-TII)/ALOG(P11(III)/P11(III))
2210 DT=TN-TII
      IF (DT.GT.150*TAU) TAU=DT/150
      IF (DT.GT.150*TTAU) TTAU=DT/150
      P11(NEVENT)=P11(III)*EXP(-DT/TTAU)
      P4(NEVENT)=P4(III)*EXP(-DT/TAU)
2221 IF (NDI.NE.NEVENT) GO TO 2225
      DC 2224 I=1,NEVENT
      P2(I)=P1(I)
      P3(I)=P1(I)
      P12(I)=P11(I)
      P13(I)=P11(I)
      P5(I)=P4(I)
      P6(I)=P4(I)
2224 GOTO 1111
C
C      CALCULATE P2() AND P5() AND P12() VECTORS
C
2225 PF=1.
      N=NEVENT
      PPP=FLOAT(N+1)/FLCAT(N+2)
      P=1.
      J=0
      DC 2399 I=1,NEVENT
      IF (IT(I).EQ.1) GOTO 2350
      J=1
      GOTO 2399
2350 PP=PP*FLOAT(N-1)/FLOAT(N)
      P=P*FLOAT(N)/FLOAT(N+1)
      PPP=PPP*FLCAT(N)/FLCAT(N+1)
      P2(I)=PP
      P5(I)=P
      P12(I)=PPP
      J=C

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2399 N=N-1
      IF (J.EQ.0) GOTC2425
      IF (NDI.GT.1) GOTO 2400
      TAL=-TII/ALCG(P5(II))
      TTAU=-TII/ALOG(P12(II))
      GOTC2410
2400 TAU=DTI/ALOG(P5(II)/P5(III))
      TTAU=DTI/ALCG(P12(II)/P12(III))
2410 IF (DT.GT.150*TAU) TAU=DT/150
      P5(NEVENT)=P*EXP(-DT/TAU)
      IF (DT.GT.150*TTAU) TTAU=DT/150.
      P12(NEVENT)=PPP*EXP(-DT/TTAU)

C
C
C      CALCULATE P3() AND P6() AND P13() VECTORS
2425 PP=1.
      N=NEVENT
      P=1.
      PPP=FLOAT(N+1)/FLCAT(N+2)
      J=C
      TT=0
      TTT=0
      DC 2599 I=1,NEVENT
      IF (IT(I).EQ.1) GOTO 2550
      J=J+1
      TT=TT+T(I)-TTT
      GOTC 2599
2550 DN=N
      IF (TT.NE.0) DN=DN+TT/(T(I)-TTT)
      PF=PP*(DN-1)/DN
      P=P*DN/(DN+1)
      PPP=PPP*DN/(DN+1.)
      P3(I)=P
      P6(I)=P
      P13(I)=PPP
      J=C
      TT=0
      TTT=T(I)
2599 N=N-1
      IF (J.EQ.0) GO TO 1111
      DTT=TN-TTT
      IF (DTT.GE.1E-70) GOTC 3005
      DN=.5*(J+1)
      GOTC 3010
3005 DN=TT/DTT
3010 P6(NEVENT)=P*SQRT(DN/(DN+1.))
      IF (NDI.GT.1) GO TO 1997
      TTAU=-TII/ALOG(P13(II))
      GOTC 1998
1997 TTAU=DTI/ALCG(P13(II)/P13(III))
1998 IF (DT.GT.150*TTAU) TTAU=DT/150.
      P13(NEVENT)=PPP*EXP(-DT/TTAU)

C
C      SET UP A LOOP FOR ALL JACKKNIFE CALLS (PJ4,PJ5,PJ6)
C
1111 DC 1000 I=1,NEVENT
      DC 1011 J=1,NEVENT
      PJ2(I,J)=0.C
      PJ4(I,J)=0.C
      PJ5(I,J)=0.C
      PJ6(I,J)=0.C
      SL1(I,J)=0.C
      SL2(I,J)=0.C
      SL3(I,J)=0.C
      SL4(I,J)=0.C
1011 CCNTINUE
      PJA2(I)=0.0
      PJA4(I)=0.0
      PJA5(I)=0.0
      PJA6(I)=0.0
      SEAR1(I)=0.C
      SEAR2(I)=0.C

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        SEAR3(I)=0.C
        SEAR4(I)=0.C
        PZ2(I)=0.0
        VARJ2(I)=0.C
        VARJ4(I)=0.C
        VARJ5(I)=0.C
        VARJ6(I)=0.C
1000  CCNTINUE
        DC 1001 I=1, NEVENT
C
C  MOVES DATA INTO TJ() AND ITJ() VECTORS
C
        K=1
        JNEXT=0
        JBEFCR=0
        JAFTR=0
1002  IF(K.EQ.1)GO TO 7003
        TJ(K)=T(K)
        ITJ(K)=IT(K)
        IF(IT(K).EQ.0)GO TO 7001
        JNEXT=JBEFCR
        JBEFCR=K
7001  K=K+1
        GC TO 1002
7002  JAFTR=JBEFCR
        JBEFCR=JNEXT
        GC TO 1010
7003  IF(I.GT.JEVENT) GO TO 7002
1003  IF(K.GT.JEVENT)GO TO 1004
        TJ(K)=T(K+1)
        ITJ(K)=IT(K+1)
        IF(ITJ(K).EQ.0)GO TO 4002
        IF(JAFTR.EQ.0)JAFTR=K
4002  K=K+1
        GC TO 1003
1004  IF(JAFTR.EQ.0)JAFTR=JEVENT
1010  NDIJ=NDI-IT(I)
C
C  CHECK IF ZERO DEATHS
C
        IF(NDIJ.EQ.0)GOTO 1001
        N=NDIJ
        P=1.
        J=0
        III=0
        III=0
        GC TO 1014
C
C  CALCULATE PJ4()VECTORS
C
1014  DC 1010 IJ=1, JEVENT
        IF(ITJ(IJ).EQ.1)GO TO 1015
        J=1
        GC TO 1016
1015  P=FLOAT(N)/FLOAT(NDIJ+1)
        PZ(IJ)=P
        III=II
        II=IJ
        J=0
        N=N-1
1016  CCNTINUE
        IF(J.EQ.0)GC TO 1019
        TII=TJ(II)
        DTI=TJ(III)-TII
        IF(NDIJ.GT.1)GO TO 1017
        TAU=-TII/ALCG(PZ(II))
        GC TO 1018
1017  TAL=DTI/ALOG(PZ(II)/PZ(III))
1018  DT=TJ(JEVENT)-TII
        IF(DT.GT.150.*TAU)TAU=DT/150.
        PZ(JEVENT)=P*EXP(-DT/TAU)
1019  K=1

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1020 IF(K.EQ.I)GC TO 2021
    PJ4(K,I)=ALCG((PZ(K)+A)/(1-PZ(K)+A))
    K=K+1
    GC TO 1020
2021 IF(K.EQ.NEVENT)GO TO 5021
    IF(IT(I).EQ.0)GO TC 1025
5021 TJA=TJ(JAFTER)
    IF(JBEFCR.NE.0)GO TO 1022
    PX=ALOG(PZ(JAFTER))*T(I)/TJA
    PZZ=0.0
    IF (PX.LT.-150) GCTO 7025
    PZZ=EXP(PX)
7025 PJ4(I,I)=ALCG((PZZ+A)/(1-PZZ+A))
    GC TO 1025
1022 TJB=TJ(JBEFCR)
    DTJ=TJA-TJB
    CTE=T(I)-TJB
    PX=DTB*ALOG(PZ(JAFTER)/PZ(JBEFCR))/DTJ
    PZZ=0.0
    IF (PX.LT.-150) GCTO 7025
    PZZ=PZ(JBEFCR)*EXP(PX)
7026 PJ4(I,I)=ALCG((PZZ+A)/(1-PZZ+A))
1025 IF(K.GT.JEVENT)GOTO 3027
    PJ4(K+1,I)=ALOG((PZ(K)+A)/(1-PZ(K)+A))
    K=K+1
    GC TO 1025
3027 IF(NDI.NE.NEVENT)GCTO 1026
    CC 3028 K=1,NEVENT
    CC 3028 L=1,NEVENT
    PJ5(K,L)=PJ4(K,L)
3028 PJ6(K,L)=PJ4(K,L)
    GCTO 1001

C
C  CALCULATE PJ5() VECTORS
C
1026 N=JEVENT
    P=1.
    PP=1.
    J=C
    CC 1028 IJ=1,JEVENT
    IF(ITJ(IJ).EQ.1)GC TO 1027
    J=1
    GC TO 1028
1027 P=P*FLOAT(N)/FLOAT(N+1)
    PP=PP*FLOAT(N-1)/FLOAT(N)
    PZ(IJ)=P
    PZ2(IJ)=PP
    J=J+1
1028 N=N-1
    IF(J.EQ.0)GC TO 1031
    IF(NDIJ.GT.1)GO TC 1029
    TAL=-TII/ALCG(PZ(II))
    GC TO 1030
1029 TAL=DTI/ALCG(PZ(II)/PZ(III))
1030 IF(DT.GT.150*TAU)TAU=DT/150.
    PZ(JEVENT)=P*EXP(-CT/TAU)
1031 K=1
1032 IF(K.EQ.I)GC TO 2033
    PJ5(K,I)=ALCG((PZ(K)+A)/(1-PZ(K)+A))
    PJ2(K,I)=ALCG((PZ2(K)+A)/(1-PZ2(K)+A))
    K=K+1
    GC TO 1032
2033 IF(K.EQ.NEVENT)GO TO 5033
    IF(IT(I).EQ.0)GO TO 1136
5033 IF(JBEFCR.NE.0)GO TO 1135
    PX=T(I)*ALOG(PZ(JAFTER))/TJA
    PZZ=0.0
    IF (PX.LT.-150) GCTO 7027
    PZZ=EXP(PX)
7027 PJ5(I,I)=ALCG((PZZ+A)/(1-PZZ+A))
    GC TO 1136
1135 PX=DTB*ALOG(PZ(JAFTER)/PZ(JBEFCR))/DTJ

```

```

PZZ=0.0
IF (PX.LT.-150) GOTO 7028
PZZ=PZ(JBEFCR)*EXP(PX)
7028 PJ5(I,I)=ALCG((PZZ+A)/(1-PZZ+A))
1136 IF(K.GT.JEVENT)GO TO 1036
PJ5(K+1,I)=ALOG((PZ(K)+A)/(1-PZ(K)+A))
PJ2(K+1,I)=ALOG((PZ2(K)+A)/(1-PZ2(K)+A))
K=K+1
GC TO 1136

```

C
C CALCULATE PJ6() VECTORS
C

```

1036 N=JEVENT
P=1.
J=0
TT=0
TTT=0
DC 1038 IJ=1,JEVENT
IF(ITJ(IJ).EQ.1)GC TO 1037
J=1
TT=TT+TJ(IJ)-TTT
GC TO 1038
1037 CN=N
IF(TT.NE.0)CN=DN+TT/(TJ(IJ)-TTT)
P=P*DN/(DN+1.)
PZ(IJ)=P
J=C
TT=C
TTT=TJ(IJ)
1038 N=N-1
IF(J.EQ.0)GC TO 1041
DT=TJ(JEVENT)-TTT
IF(DT.GT.1E-7)GC TO 1039
DN=.5*(J+1)
GC TO 1040
1039 DN=TT/DT
1040 PZ(JEVENT)=P*SQRT(DN/(DN+1.))
1041 K=1
1042 IF(K.EQ.1)GC TO 2043
PJ6(K,I)=ALCG((PZ(K)+A)/(1-PZ(K)+A))
K=K+1
GC TO 1042
2043 IF(K.EQ.NEVENT)GO TO 5043
IF(IT(I).EQ.0)GO TO 1146
5043 IF(JBEFCR.NE.0)GC TO 1045
PX=T(I)*ALOG(PZ(JAFTER))/TJA
PZZ=0.0
IF (PX.LT.-150) GOTO 7029
PZZ=EXP(PX)
7029 PJ6(I,I)=ALCG((PZZ+A)/(1-PZZ+A))
GC TO 1146
1045 PX=DTB*ALOG(PZ(JAFTER)/PZ(JBEFCR))/DTJ
PZZ=0.0
IF (PX.LT.-150) GOTO 7030
PZZ=PZ(JBEFCR)*EXP(PX)
7030 PJ6(I,I)=ALCG((PZZ+A)/(1-PZZ+A))
1146 IF(K.GT.JEVENT)GOTO 1001
PJ6(K+1,I)=ALOG((PZ(K)+A)/(1-PZ(K)+A))
K=K+1
GC TO 1146
1001 CCNTINUE
DC 8888 I=1,NEVENT
DC 8811 J=1,NEVENT
IF(I.EQ.J)GOTO 8811
SL1(I,J)=FLCAT(NEVENT)*ALOG((P4(I)+A)/(1-P4(I)+A))-FLC
*AT(NEVENT-1)*PJ4(I,J)
SL2(I,J)=FLCAT(NEVENT)*ALOG((P5(I)+A)/(1-P5(I)+A))-FLC
*AT(NEVENT-1)*PJ5(I,J)
SL3(I,J)=FLCAT(NEVENT)*ALOG((P6(I)+A)/(1-P6(I)+A))-FLC
*AT(NEVENT-1)*PJ6(I,J)
SL4(I,J)=FLCAT(NEVENT)*ALOG((P2(I)+A)/(1-P2(I)+A))-FLC
*AT(NEVENT-1)*PJ2(I,J)

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88111 CCNTINUE
88222 CCNTINUE
DC 9999 I=1,NEVENT
DC 9911 J=1,NEVENT
SBAR1(I)=SL1(I,J)/NEVENT+SBAR1(I)
SBAR2(I)=SL2(I,J)/NEVENT+SBAR2(I)
SBAR3(I)=SL3(I,J)/NEVENT+SBAR3(I)
SBAR4(I)=SL4(I,J)/NEVENT+SBAR4(I)
9911 CCNTINUE
IF(SBAR1(I).LT.-180.)SBAR1(I)=-180.
IF(SBAR1(I).GT.174.)SBAR1(I)=174.
IF(SBAR2(I).LT.-180.)SBAR2(I)=-180.
IF(SBAR2(I).GT.174.)SBAR2(I)=174.
IF(SBAR3(I).LT.-180.)SBAR3(I)=-180.
IF(SBAR3(I).GT.174.)SBAR3(I)=174.
IF(SBAR4(I).LT.-180.)SBAR4(I)=-180.
IF(SBAR4(I).GT.174.)SBAR4(I)=174.
EL1=EXP(SBAR1(I))
EL2=EXP(SBAR2(I))
EL3=EXP(SBAR3(I))
EL4=EXP(SBAR4(I))
PJA4(I)=EL1/(1.+EL1)
PJA5(I)=EL2/(1.+EL2)
PJA6(I)=EL3/(1.+EL3)
PJA2(I)=EL4/(1.+EL4)
9999 CCNTINUE
DC 8999 I=1,NEVENT
DC 8911 J=1,NEVENT
IF(SL4(I,J).EQ.0.0)GOTO 3111
VARJ2(I)=(SL4(I,J)-SBAR4(I))*2/FLCAT(NEVENT-1)+VARJ2(
*I)
3111 IF(SL1(I,J).EQ.0.0)GOTO 3112
VARJ4(I)=(SL1(I,J)-SBAR1(I))*2/FLCAT(NEVENT-1)+VARJ4(
*I)
3112 IF(SL2(I,J).EQ.0.0)GOTO 3113
VARJ5(I)=(SL2(I,J)-SBAR2(I))*2/FLCAT(NEVENT-1)+VARJ5(
*I)
3113 IF(SL3(I,J).EQ.0.0)GOTO 8911
VARJ6(I)=(SL3(I,J)-SBAR3(I))*2/FLCAT(NEVENT-1)+VARJ6(
*I)
8911 CCNTINUE
RMS1(I)=SQRT(VARJ2(I)/FLOAT(NEVENT))
RMS2(I)=SQRT(VARJ4(I)/FLOAT(NEVENT))
RMS3(I)=SQRT(VARJ5(I)/FLCAT(NEVENT))
RMS4(I)=SQRT(VARJ6(I)/FLOAT(NEVENT))
UP1(I)=SBAR4(I)+RMS1(I)*TN
RO1(I)=SBAR4(I)-RMS1(I)*TN
UP2(I)=SBAR1(I)+RMS2(I)*TN
RC2(I)=SBAR1(I)-RMS2(I)*TN
UP3(I)=SBAR2(I)+RMS3(I)*TN
RC3(I)=SBAR2(I)-RMS3(I)*TN
UP4(I)=SBAR3(I)+RMS4(I)*TN
RC4(I)=SBAR3(I)-RMS4(I)*TN
IF(UP1(I).GT.174.)UP1(I)=174.
IF(UP1(I).LT.-100.)UP1(I)=-100.
IF(RO1(I).GT.174.)RO1(I)=174.
IF(RO1(I).LT.-100.)RO1(I)=-100.
IF(UP2(I).GT.174.)UP2(I)=174.
IF(UP2(I).LT.-100.)UP2(I)=-100.
IF(RO2(I).GT.174.)RO2(I)=174.
IF(RO2(I).LT.-100.)RO2(I)=-100.
IF(UP3(I).GT.174.)UP3(I)=174.
IF(UP3(I).LT.-100.)UP3(I)=-100.
IF(RO3(I).GT.174.)RO3(I)=174.
IF(RO3(I).LT.-100.)RO3(I)=-100.
IF(UP4(I).GT.174.)UP4(I)=174.
IF(UP4(I).LT.-100.)UP4(I)=-100.
IF(RO4(I).GT.174.)RO4(I)=174.
IF(RO4(I).LT.-100.)RO4(I)=-100.
UINT1(I)=1./(1.+1./EXP(UP1(I)))
RINT1(I)=1./(1.+1./EXP(RO1(I)))
UINT2(I)=1./(1.+1./EXP(UP2(I)))

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      RINT2(I)=1./(1.+1./EXP(RO2(I)))
      UINT3(I)=1./(1.+1./EXP(UP3(I)))
      RINT3(I)=1./(1.+1./EXP(RO3(I)))
      UINT4(I)=1./(1.+1./EXP(UP4(I)))
      RINT4(I)=1./(1.+1./EXP(RO4(I)))
8999 CCNTINUE

C
C   PRINT OUTPUT
C
3050 IF (NWRITE.EQ.0) GOTO 3500
      WRITE(6,8)(IREPL,IT(I),T(I),P1(I),P2(I),P3(I),P4(I),P5
      *(I),P6(I),PJA2(I),PJA4(I),PJA5(I),PJA6(I),P11(I),P12(I
      *),P13(I),I=1,NEVENT)
      WRITE (6,10)

C
C   REDUCE VECTORS
C
3500 K=1
      DO 4000 I=1,NEVENT
      IF (IT(I).EQ.1) GOTO 3900
      IF (I.NE.NEVENT) GOTO 4000
3900 T(K)=T(I)
      P1(K)=P1(I)
      P2(K)=P2(I)
      P3(K)=P3(I)
      P4(K)=P4(I)
      P5(K)=P5(I)
      P6(K)=P6(I)
      PJA2(K)=PJA2(I)
      PJA4(K)=PJA4(I)
      PJA5(K)=PJA5(I)
      PJA6(K)=PJA6(I)
      P11(K)=P11(I)
      P12(K)=P12(I)
      P13(K)=P13(I)
      UINT1(K)=UINT1(I)
      RINT1(K)=RINT1(I)
      UINT2(K)=UINT2(I)
      RINT2(K)=RINT2(I)
      UINT3(K)=UINT3(I)
      RINT3(K)=RINT3(I)
      UINT4(K)=UINT4(I)
      RINT4(K)=RINT4(I)
      K=K+1
4000 CCNTINUE

C
C   CALCULATE DIFFERENCES, MEAN, RMS, AND MEAN ABS
C
      K=1
      TTT=0
      CT=-T(1)
      PP1=1
      PF2=1
      PF3=1
      PF4=1
      PF5=1
      PF6=1
      PF8=1
      PF9=1
      PF10=1
      PF11=(NDI+1.)/DII
      PF12=(NEVENT+1.)/(NEVENT+2.)
      PF13=PP12
      PF14=1.
      PC4=P4(1)
      PC5=P5(1)
      PC6=P6(1)
      PC8=PJA4(1)
      PC9=PJA5(1)
      PC10=PJA6(1)
      CCNF1=1.
      CCNF2=1.

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CCNF3=1.
CCNF4=1.
CCNF5=1.
CCNF6=1.
CCNF7=1.
CCNF8=1.
T4=DT/ALOG(PQ4)
T5=DT/ALOG(PQ5)
T6=DT/ALOG(PQ6)
T8=DT/ALOG(PQ8)
T9=DT/ALOG(PQ9)
T10=DT/ALOG(PQ10)
DC 4998 J=1,9
TT=.1*J
P=1-TT
IF (NDIE.EQ.2) TT=-ALOG(P)
IF (TT.GT.TN) GOTC 4999
4100 NN(J)=NN(J)+1
IF (TT.LE.T(K)) GCTC 4200
PF1=P1(K)
PF2=P2(K)
PF3=P3(K)
PF4=PQ4
PF5=PQ5
PF6=PQ6
PF8=PQ8
PF9=PQ9
PF10=PQ10
PF11=P11(K)
PF12=P12(K)
PF13=P13(K)
PF14=PJA2(K)
CCNF1=UINT1(K)
CCNF2=RINT1(K)
CCNF3=UINT2(K)
CCNF4=RINT2(K)
CCNF5=UINT3(K)
CCNF6=RINT3(K)
CCNF7=UINT4(K)
CCNF8=RINT4(K)
TTT=T(K)
K=K+1
PC4=P4(K)
PC5=P5(K)
PC6=P6(K)
PC8=PJA4(K)
PC9=PJA5(K)
PC10=PJA6(K)
DT=T(K)-TTT
TT1=ALOG(PF8/PQ8)
TT2=ALOG(PF9/PQ9)
TT3=ALOG(PF10/PQ10)
IF (PC8.EQ.0.0.OR.TT1.EQ.0.0) TT1=ALOG((PF8+A)/(1-PC8+A))
IF (PC9.EQ.0.0.OR.TT2.EQ.0.0) TT2=ALOG((PF9+A)/(1-PC9+A))
IF (PC10.EQ.0.0.OR.TT3.EQ.0.0) TT3=ALOG((PF10+A)/(1-PC10+A))
T4=DT/ALOG(PF4/PQ4)
T5=DT/ALOG(PF5/PQ5)
T6=DT/ALOG(PF6/PQ6)
T8=DT/TT1
T9=DT/TT2
T10=DT/TT3
GCTC 4100
4200 D(1)=PF1-P
D(2)=PF2-P
D(3)=PF3-P
D(7)=TTT-TT
D(4)=PF4*EXP(DTT/T4)-P
D(5)=PF5*EXP(DTT/T5)-P
D(6)=PF6*EXP(DTT/T6)-P
D(7)=EXP(TT*T7)-P
D(8)=PF8*EXP(DTT/T8)-P
D(9)=PF9*EXP(DTT/T9)-P

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C(10)=PP10*EXP(DTT/T10)-P
D(11)=PP11-F
D(12)=PP12-F
D(13)=PP13-F
D(14)=PP14-P
CF(1)=CCNF1
CF(2)=CCNF2
CF(3)=CCNF3
CF(4)=CCNF4
CF(5)=CCNF5
CF(6)=CCNF6
CF(7)=CCNF7
CF(8)=CCNF8
DC 4911 I=1,8
CC=CF(I)
C(I,J)=C(I,J)+CC
IF(NWRITE.EQ.0)GOTC 4911
WRITE(6,10)
WRITE(6,12) I,P,(C(I,L),L=1,9)
4911 CCNTINUE
DC 4997 I=1, NP
DD=D(I)
U(I,J)=U(I,J)+DD
W(I,J)=W(I,J)+ABS(DD)
S(I,J)=S(I,J)+DD*DD
IF (NWRITE.EQ.0) GOTO 4997
WRITE (6,10)
WRITE (6,12) I,P ,(U(I,L),L=1,9)
WRITE (6,12) I,P ,(W(I,L),L=1,9)
WRITE (6,12) I,P ,(S(I,L),L=1,9)
4997 CCNTINUE
4998 CCNTINUE
C
4999 CONTINUE
C
C
C PRINT SUMMARY STATISTICS
DC 5100 J=1,9
5100 D(J)=.1*J
WRITE (6,7) (D(J),J=1,9)
WRITE (6,10)
DC 5300 J=1,9
IF (NN(J).GT.0) GOTO 5150
DC 5125 I=1,NP
S(I,J)=1E9
U(I,J)=1E9
C(I,J)=1E9
5125 W(I,J)=1E9
GOTO 5300
5150 XJ=1./NN(J)
5200 DC 5250 I=1,NP
S(I,J)=SQRT(S(I,J)*XJ)
U(I,J)=U(I,J)*XJ
C(I,J)=C(I,J)*XJ
5250 W(I,J)=W(I,J)*XJ
5300 CCNTINUE
WRITE (6,9) (I,(U(I,J),J=1,9),I=1,NP)
WRITE (6,10)
WRITE (6,14) (I,(W(I,J),J=1,9),I=1,NP)
WRITE (6,10)
WRITE (6,16) (I,(S(I,J),J=1,9),I=1,NP)
WRITE (6,10)
WRITE (6,20) (I,(C(I,J),J=1,9),I=1,8)
20 FCRMAT(1X,'C',11,9F8.3,3X,'CCNF')
WRITE(6,10)
WRITE (6,15) (NN(J),J=1,9)
IF (NENC.NE.0) GOTC 108
WRITE (6,11)
GOTO 25
ENC

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