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Estimation in a Marked Poisson Error Recapture Model of Software Reliability

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ESTIMATION IN A MARKED POISSON ERROR RECAPTURE
MODEL OF SOFTWARE RELIABILITY

by

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M.S., University of Manitoba, 1984

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Abstract

Estimation in a Marked Poisson Error Recapture Model of Software Reliability

Rajan Gupta
Old Dominion University, 1991
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Nayak's (1988) model for the detection, removal, and recapture of the errors in a computer program is extended to a larger family of models in which the probabilities that the successive programs produce errors are described by the tail probabilities of discrete distribution on the positive integers. Confidence limits are derived for the probability that the final program produces errors. A comparison of the asymptotic variances of parameter estimates given by the error recapture and by the repetitive-run procedure of Nagel, Scholz, and Skrivan (1982) is made to determine which of these procedures efficiently uses the test time.
To my parents
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Chapter 1

Introduction

A great many papers have appeared concerning software reliability and reliability growth models during the debugging of a program. This is due, in part, to an early realization that the reliability of many systems may depend critically upon the reliability of computer programs; although the hardware component of these systems may be highly reliable, the total system reliability is often limited by that of the software component. Software reliability research is generally aimed at providing the capability to design and build reliable software systems in a cost-effective way.

One of the major factors contributing to the very high levels of reliability that can now be achieved in hardware systems, is the use of component redundancy to provide
tolerance to physical faults. Software redundancy techniques, such as recovery blocks and N-version programming (e.g., [1],[38]) have been proposed which aim to provide tolerance to design faults, the main cause of unreliability in software systems. These techniques are based on the implementation of functionally equivalent, but independently developed modules of diverse design, with provision for either state restoration or replication, and of some means for co-ordinating between the outputs from the replicated modules.

Some highly critical systems have relied on the construction of independently designed versions of the entire software system (e.g., Space Shuttle, A310 Airbus, railway signalling). Many database systems and telephone switching systems employ sophisticated recovery techniques which can prevent corruption of data by certain categories of software faults.

One of the more commonly used techniques for predicting software reliability utilizes reliability growth models. Reliability improves as a result of the process of fault identification and correction known as debugging. These models require failure data and place stringent requirements upon the testing strategy which generates the data.
To obtain more data than is provided by the usual debugging experiment, Nagel, et. al. (1984) proposed a repetitive-run procedure in which a program is restored to its original form and the debugging sequence repeated, perhaps several times, using independently generated series of inputs. The extra information provided by this procedure has been used to check the assumption of exponentially distributed inter-failure times and also to show that distinct errors may have different occurrence rates. Except for the work of Scholz (1986), models that treat specific features of this design do not seem to have been studied in the literature.

In other experiments such as seeding and tagging, inference is not based on the usual observed failure data. The method described in the following paragraphs is based on the one used for many years to estimate the size of animal and fish populations.

Feller (1957) gives a procedure for estimating the number of a certain type of fish in a lake. A new catch is made and the tagged fish as well as untagged fish are counted. Tagged fish are assumed to mix randomly with the untagged fish. The number of fish in the lake can then be estimated by assuming that the proportion of the tagged fish which are re-caught is equal to the proportion of fish in the lake contained within the second catch.
There are two ways of applying these ideas to software reliability. The first is known as seeding, originally suggested by Mills (1970), and consists of inserting a known number of bugs into a program, and monitoring the proportion of inserted bugs found during the debugging process. The second alternative is tagging, suggested by Rudner (1977) which is carried out by giving the program to two programmers. The number of faults found by the first programmer are regarded as tagged, and inferences are drawn from the proportion of the second programmer's faults which have been tagged.

There are a number of drawbacks with these methods, especially with regard to the implicit assumptions which have to be made. For example, both methods assume that all faults are equally likely to be found, seeding assumes that the seeded faults are representative of the indigenous faults, and tagging assumes that the programmers can act independently.

In seeding and tagging procedures, the estimation problem for the number of faults, however, is not onerous. The maximum likelihood estimator is biased, but a modified estimator due to Chapman (1951) has a lower bias and generally better properties, when the the number of faults is more than 50.
Another drawback is that one can not estimate quantities such as failure rates. This method is only useful for bug counting.

The most important criticism for the simple reliability growth models is that they essentially treat software as a 'black box'. No account is taken of internal structure or other known properties of the program under study. There is a need for models that can utilize the large amount of structural information usually available. Hardware reliability theory provides an interesting parallel. One of the most important achievements of this theory is the ability to combine information about component reliability with structural information about the design of the overall system. Unfortunately, software structure tends to be much more complex than hardware structure. Also, the simple component/design dichotomy is less obviously applicable to software, which can be viewed as solely levels of design.

To resolve some of the existing problems associated with models for the usual debugging experiment, Nayak (1988) introduced recapture debugging as a way to get extra information for estimating the number of faults remaining in a system. By placing counters in the software (for an alternative to software testing counters, see p. 25) we observe, in addition to the usual sequence of failure (i.e.,
error detection) times, the hitting frequencies of detected faults. Nayak's (1988) discussion concerns the Jelinski-Moranda (1972) model and procedures useful for estimating the number of remaining faults.

Chapter 2 describes the classification and motivation for a number of software reliability growth models. The main body of new material begins in Chapter 3 and concerns a family of marked Poisson process models for the recapture debugging procedure. In one form, the model describes a stationary event series and an attached Markov chain. An equivalent form, in the sense of giving the same likelihood function, is that of a nonstationary series of main events together with a collection of independent counting processes. The latter form of the model was originally suggested by Nayak (1988), although he considered only the case in which the main event series is a linear pure death process, also known as the Jelinski-Moranda (1972) model. The main contributions of the present work are (i) a procedure for estimating the probability that the final program version produces errors and (ii) a comparison of the error recapture and repetitive-run procedures in terms of the asymptotic variances of parameter estimates obtained by the two procedures. Chapter 4, the final chapter, studies, by using repetitive-run data, the goodness of fit of certain models based on parameterizations introduced in Chapter 3.
Chapter 2

Background

2.1 Introduction

The software segment of a computer system involves instructions or codes used to program the hardware system. Some of the inputs for which a specific job data set or function does not produce the desired output lead to what is termed as software failures. These failures are either due to errors in the coding of the instructions (the program) or an input that is incompatible with the design of the software system.

Early debugging designs consist of detecting and correcting a series of errors during a specified period of testing. If the software is executed on a series of inputs, it may work satisfactorily until time $S_1$, when the first
failure occurs. The programmer then repairs the program, it works satisfactorily until time $S_2$, then it is repaired, and so on. This process is sometimes referred to as a debugging experiment. The models used to describe the failure times $S_1, S_2, \ldots$, are called reliability growth models since, typically the gaps $Y_i = S_i - S_{i-1}$ ($S_0 \equiv 0$), $i=1,2,\ldots$, between failures will increase as faults are removed from the software.

This may not be exactly so due to the fact that the failure times are random, and thus they are subject to statistical fluctuations. A number of models have been proposed in the literature to study such failures. These models assume that failure times have distributions with parameters that depend on the residual faults in the software system. The assumed distributions reflect the software quality as faults are detected and removed from the system.

The models described in Sections 2.2–2.7 are based on assumptions concerning the failure gaps $\{Y_i\}$, the event occurrence times $\{S_1, S_2, \ldots\}$, or the counting process $\{N(t) : t \geq 0\}$ where $N(t)$ is determined by $N(t)=n$ if and only if $S_n < t < S_{n+1}$, with $S_0 \equiv 0$. 
2.2 The Jelinski-Moranda Model

One of the earliest and most widely referenced models is that of Jelinski and Moranda (1972). The model assumes that the failure rate at any point in time is proportional to the current fault content of the program. The initial fault content is denoted by \( v \) and the contribution of each fault by \( \phi \). The failure rate initially is \( v\phi \), and decreases to \( (v-1)\phi \) after the first fault is detected and eliminated, and so on. This model views the occurrence times of the first, second, ..., and \( r \)th failure as the first \( r \) order statistics, in a sample of size \( v \) from an exponential distribution. The times between failures are distributed as independent exponential random variables with rate parameters \( (v-i+1)\phi \), \( i=1,2,...,v \).

Moranda (1979) points out that most models assume that a system is restored to its initial state after repair. For reliability growth models where the failure rate varies, it is generally assumed that this rate changes deterministically or continuously with time. As noted by Littlewood (1981), the failure rate of J-M model changes in discrete steps and this is an essential feature to represent reliability growth.
An inherent assumption of the model is that each bug contributes the same amount \( \phi \) to the overall failure rate. This has been criticized in particular by Littlewood (1981) who argues that the different frequencies of execution of different portions of code will in itself lead to different occurrence rates of faults, all other things being equal. The assumptions of constant failure rate and exponential distributions for the times between successive failures have also been criticized by Schick and Wolverton (1978).

Forman and Singpurwalla (1979) show that the maximum likelihood estimator of \( v \) is unstable, and can be highly misleading when the number of remaining faults is anything but small. Sukert (1977) has observed that the estimate of \( v \) does not always exist, and Littlewood and Verrall (1981) give a condition for its existence which reduces to a requirement that the data exhibit the assumed reliability growth. Littlewood and Verrall (1973) suggest a Bayesian version of the model, but they note that the improvement in estimation properties is marginal in most cases.

### 2.3 The Musa Execution Time Model

This is perhaps the most practical model of all and has been developed to the extent that it takes into account
various personnel and resource constraints.

The model is based on the Jelinski-Moranda (1972) model, but introduces a number of refinements. One of these is an error reduction factor B representing the average ratio of the rate of fault correction to the failure rate. There is an implicit assumption, that the fault correction rate is proportional to the failure rate - not equal to it as assumed by the Jelinski-Moranda (1972), Shooman (1973), Schick-Wolverton (1978) and Littlewood (1980) models. Also introduced is a testing compression factor C which is the average ratio of the failure rate during test to that during operational use.

This model also differs from others in its treatment of time. The program execution time is taken as the time variable, and this in turn is related to calendar time via constraints on fault correction personnel, and computer time. These aspects make the model potentially valuable as a management tool.

The assumptions of this model answer some of the criticisms of the Jelinski-Moranda (1972) model, but the underlying assumption that each fault contributes the same amount to the overall failure rate is often considered to be a weakness. Chenoweth (1981) suggests a generalization where
to allow for improved debugging in the later stages.

Since the basic model structure is identical to that of the Jelinski-Moranda model, the same parameter estimation problems are encountered.

2.4 Poisson Models

If each fault is removed the first time it produces a failure, then the failure rate will decrease as in the previous models, but it may depend on the time of the most recent repair rather than on the total number of repairs that have been made. This is a basic assumption underlying time dependent Poisson process models.

A nonhomogeneous Poisson process \(\{N(t): t \geq 0\}\) with intensity function \(\lambda(x)\) (e.g., Parzen, p. 252) is defined by the following conditions:

(i) \(N(0) = 0\).

(ii) \(N(t), t \geq 0\) has independent increments.

(iii) The number of events \(N(s,t)\) occurring in the interval \((s,t)\) has a Poisson distribution with mean \(\int_s^t \lambda(x)\,dx\).
These assumptions imply that the occurrence time $S_1$ of the first event has distribution

$$P(S_1 > t) = P(N(t) = 0)$$

$$= \exp \left( -\int_0^t \lambda(x) \, dx \right), \quad t \geq 0$$

$$= 1, \quad t < 0$$

By varying $\lambda(x)$ over the entire class of nonnegative functions that are integrable over bounded intervals and which satisfy

$$\lim_{t \to \infty} \int_0^t \lambda(x) \, dx = \infty,$$

we obtain all possible continuous distributions for $S_1$. The importance of the nonhomogeneous Poisson process lies in the fact that the increments of the process are not required to be stationary. Most applications of the model assume that $\lambda(t)$ is either constant, increasing, or decreasing in $t \geq 0$. The mean number of events occurring in the interval $(0,t)$, denoted by $m(t)$, is given by

$$m(t) = \int_0^t \lambda(x) \, dx$$
If events occur at times $S_1, S_2, \ldots$, over a fixed interval $(0,t)$ and follow a non-homogeneous Poisson process, then the joint density function of $S_1, S_2, \ldots, S_{N(t)}$ and $N(t)$ is

$$\prod_{i=1}^{n} \lambda(s_i) \exp \left[ -\int_{0}^{t} \lambda(x) \, dx \right]$$

where if $n=0$, the product factor is equal to one.

The most widely known Poisson model of software reliability growth was proposed by Goel and Okumoto (1979). Their model assumes an intensity function and mean of the form

$$\lambda(x) = \alpha \beta e^{\beta x}, \quad x \geq 0, \quad \alpha, \beta > 0$$

$$m(t) = \alpha(1-e^{\beta t}), \quad t \geq 0, \quad \alpha, \beta > 0$$

Since $\lim m(t) = \alpha$ as $t \to \infty$, the parameter $\alpha$ is the expected total number of failures that may eventually occur, while large values of $\beta$ imply rapid reliability growth.

Under this model the conditional joint density function of $S_1, S_2, \ldots, S_{N(t)}$, given $N(t)=n$, is
Thus $S_1, S_2, ..., S_n$ are conditionally distributed as an ordered sample of size $n$ from a truncated exponential distribution with density function

$$P(x) = n! \prod_{i=1}^{n} \beta e^{-\beta s_i} (1 - e^{-\beta t})^{-1}, \quad 0 < s_1 < s_2 < ... < s_n < t$$

(2.3)

The J-M model has this same property and thus the likelihood functions given by the two models only differ in the probability distribution of $N(t)$. That is, the J-M model implies that $N(t)$ has a binomial distribution with parameters $\nu$ and $(1 - e^{-\beta t})$, whereas any Poisson model implies that $N(t)$ has a Poisson probability distribution.

### 2.5 Littlewood-Verrall Bayesian Model

Littlewood and Verrall (1973) consider the software failure process to be the result of two sources of uncertainty. One pertains to the randomness of input and the other to the state of the program. They assume that a particular subset of the input space that will cause system
failure will be encountered randomly. This leads to the assumption that successive times between failures are independently and exponentially distributed with failure rate $\lambda_i$ for the $i$th failure. The second source of uncertainty is the fault-fixing operation. The uncertainty of fault-fixing is modeled by additionally assuming that $\lambda_i$ is stochastically less than or equal to $\lambda_{i-1}$, that is,

$$P(\lambda_i < x) \geq P(\lambda_{i-1} < x), \text{ for all } x, i.$$  

The growth in reliability is thus stochastic rather than deterministic, the $\lambda_i$ being regarded as random variables. Littlewood and Verrall assume that each $\lambda_i$ has a gamma distribution with parameters $\alpha$ and $\{\psi_i\}$, where the growth function $\psi_i$ is increasing in $i$ and describes the quality of the programmer and the difficulty of the programming task.

Combining the two sources of randomness by Bayesian techniques leads to a decreasing failure rate, and a low failure rate as more time is observed without failure. The assumptions of this model seem reasonable, though Ramamoorthy and Bastani (1980) criticize the model as being too restrictive, because it does not permit perfect debugging (i.e., unlike for the the J-M model, removing a finite number of faults will not, according to the model, produce a perfect
program). One main problem with this model is the choice of \( \{\Psi_i\} \), which determines the stochastic growth pattern. It is not clear how to choose the functional forms of \( \{\Psi_i\} \), however Littlewood (1980) has suggested linear and quadratic functions, which lead to two and three parameter models.

2.6 Order Statistics Models

Besides the Jelinski-Moranda model a number of other models have been derived that belong to the order statistics class of models (e.g., Miller 1986). These derivations are typically based on the assumption that the failure gaps \( \{Y_i\} \) are independent and exponentially distributed with rate parameters \( \lambda_i \), where the \( \lambda_i \) are also random quantities.

Using Bayes' theorem, Littlewood (1973) derived a generalization of the Jelinski-Moranda model in which the ordered failure times \( S_1, S_2, \ldots, S_n \) are distributed as the first \( n \) order statistics in a sample of size \( n \) (\( n \) is a parameter) from a Pareto distribution. Arguing from simpler assumptions, he shows that if the \( i \)th failure occurs at time \( S_i = \tau \), then the failure rate \( \Lambda \) is the sum of \( n-i \) i.i.d random variables, each having a gamma density with scale parameter \( \beta + \tau \) and shape parameter \( \alpha \). Then \( \Lambda \) also has a gamma density.
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\[ g(\lambda|\tau) = (\beta + \tau)^{-1} \lambda^{(\alpha-1)} \exp\left[-(\beta + \tau) \lambda\right] / \Gamma[\alpha(\nu-1)], \]

(\lambda \geq 0, \beta > 0) \quad (2.5)

and the distribution of \( Y_{1:n} \) given \( S_1 = \tau \), is

\[ P(Y_{1:n} > y | S_1 = \tau) = \int_0^\infty e^{-\lambda y} g(\lambda | \tau) \, d\lambda \]

\[ = \left[ (\beta + \tau) / (\beta + \tau + y) \right]^{(\alpha-1)\nu}, \quad y \geq 0 \]

(2.6)

Assuming that the conditional distribution of \( Y_{n+1} \), given the past history \( S_1, S_2, \ldots, S_n \), only depends on the time \( S_n \) of entering the \( n \)th state, Littlewood also obtains the joint density of \( S_1, S_2, \ldots, S_n \), which is identical to that of the first \( n \) order statistics in a sample of size \( \nu \) from a Pareto distribution.

This Pareto order statistics model differs from the model in Section 2.5 in that (a) the gaps \( \{Y_i\} \) are not independent and (b) the Littlewood-Verrall (1973) model of Section 2.5 does not include the total error count parameter \( \nu \). Joe and Reid (1985a,b) give an alternative, and simpler, derivation of the Pareto order statistics model and conjecture that, for any order statistics model, the maximum likelihood estimate of \( \nu \) will be infinite with positive probability.
2.7 Scale Parameter Models

A key assumption made in a number of models is that the failure gaps \( \{Y_i\} \) are independent and have distributions that differ only by scale parameters, the latter being a function of the serial index of events. A scale parameter family of reliability growth models is defined by letting \( \{Y_i\} \) be independent with distribution functions

\[
F_i(y) = G(\psi_i^{-1}y)
\]  

(2.7)

where \( \{\psi_i\} \) are scale parameters and \( G(x) \) is a continuous cdf with \( G(0) = 0 \).

Examples of models within this class, the first two of which were described in Sections 2.2 and 2.5, are the following:

(i) \( \psi_i^{-1} = (v-i+1)\phi, \ i=1,2,...,v, \ \phi > 0 \)

\[
G(x) = 1 - \exp(-x), \ x \geq 0
\]

(i.e., the Jelinski-Moranda (1972), model)

(ii) \( \psi_i = \alpha + \beta i, \ \alpha > 0, \ \beta > 0 \)

\[
G(x) = 1 - (1 + x)^{-\rho}, \ x \geq 0, \ \rho > 0
\]

(Littlewood and Verrall, 1973)
(iii) $\Psi_i^{-1} = \alpha \exp(-\phi i), \alpha, \phi > 0$

$G(x) = 1 - \exp(-x), x \geq 0$

(Cox and Lewis, 1966; Moranda, 1975)

To show that (i) and (ii) imply a limitation on reliability growth, let $Z_i, Z_j, \ldots$, be i.i.d with cdf $G(x)$. Further, let

$$q_{ij} = \log \left( \frac{\Psi_j}{\Psi_i} \right), \ 1 \leq i < j \tag{2.8}$$

and consider the following representation implied by (2.7):

$$\log \left( \frac{Y_j}{Y_i} \right) = q_{ij} + \log \left( \frac{Z_j}{Z_i} \right), \ 1 \leq i < j \tag{2.9}$$

If each $q_{ij}$ is bounded above by a known constant, say $q_{ij}^0$, that depends on $i$ and $j$, then the distribution of $\log(Y_j/Y_i)$ is shifted below that of

$$q_{ij}^0 + \log \left( \frac{Z_j}{Z_i} \right)$$

The latter implies

$$P(Y_j > Y_i, \exp(q_{ij}^0)) = P(\log \left( \frac{Y_j}{Y_i} \right) > q_{ij}^0)$$

$$= P(q_{ij} + \log \left( \frac{Z_j}{Z_i} \right) > q_{ij}^0)$$

$$\leq P(Z_j > Z_i)$$
Thus for \( i < j \), \( Y_j \) can exceed a certain scalar multiple of \( Y_i \) with probability not exceeding 0.5.

To determine the constant \( q_{ij}^0 \) in case (i), write

\[ q_{ij} = \log \left[ (1 - \gamma_i)(1 - \gamma_j)^{-1} \right] \quad (2.11) \]

where, \( \gamma = (\nu + 1)^{-1} \), and \( i, j \leq \nu \) implies \( \gamma_i \leq 1 \) and \( \gamma_j \leq 1 \). Since \( q_{ij} \) is nondecreasing in \( \gamma \), and \( \gamma \leq (j+1)^{-1} \) it follows that \( q_{ij}^0 = \log(j-i+1) \) is an upper bound on \( q_{ij} \).

Similarly, in case (ii),

\[ q_{ij} = \log \left[ (1 + \gamma_j)(1 + \gamma_i)^{-1} \right], \quad 1 \leq i < j \quad (2.12) \]

where \( \gamma = \beta/\alpha \) and \( q_{ij} \) is nondecreasing in \( \gamma \). In this case the maximum value, obtained in the limit as \( \gamma \to \infty \), is \( q_{ij}^0 = \log(j/i) \).

The existence of known upper limits implies a limitation on reliability growth since the intervals between failures are unlikely to increase rapidly if the model giving the upper limit is the true model. However, the simplicity of
these models is attractive and they can be a useful tool for estimating the reliability achieved during debugging.

2.8 Stopping Rules

Stopping rules for deciding when a program is completely debugged have been proposed by Nayak (1988) and Goudie (1990). The context of their discussion, described in more detail in Chapter 3, is a Markov chain \((X_n, Y_n)\) where \(X_n\) is the number of distinct errors detected by the \(n\)th epoch and \(Y_n = n - X_n\). A transition occurs from \((X_n, Y_n) = (x, y)\) to \((x+1, y)\) with probability \((v-x)/v\), or to \((x, y+1)\) with probability \(x/v\). The parameter \(v\) is the initial number of errors in a program and thus debugging must terminate after a fixed number of transitions occur. A well known property of Markov chains is that the waiting times \(W_i\) in states \(i=1,2,...,v-1\) (i.e., between distinct error occurrences or, equivalently, between changes in \(X_n\)) are independent random variables with geometric distributions.

Nayak (1988) proposed deciding that all errors have been eliminated when \(W_i\) first exceeds a positive integer \(k_i\) chosen so that

\[
\alpha \geq P(W_i > k_i; v=i+1) \quad (2.13)
\]
where $a$ is a prechosen error level. Since

$$P(W_i > k_i; \nu = i+1) = \left[ \frac{i}{i+1} \right]^x$$

(2.13) is satisfied if $k_i = \lfloor a_i \rfloor + 1$,

where

$$a_i = \frac{\log a_i}{\log i - \log(i+1)}$$

and $\lfloor x \rfloor$ is the integer part of the real number $x$. Nayak (1988) determined the probability of correcting all errors and tabled these probabilities for various values of $a$ and $\nu$.

Since $P(X_i = 1) = 1$, the likelihood function based on observing $W_1, W_2, ..., W_{x-1}$ and $X_n = x$ is

$$L(\nu) = \left\{ \prod_{j=1}^{x-1} \left( \frac{j}{\nu} \right)^{w_j} \left( 1 - \frac{j}{\nu} \right)^{w_x} \right\} \left( \frac{x}{\nu} \right)^{w_x}$$

(2.15)

where $w_x = n-1-(W_1 + W_2 + ... + W_{x-1})$. This takes the simpler form (Goudie, 1990)

$$L(\nu) = c \binom{\nu}{x} / \nu^n, \quad \nu = x, x+1, ...,$$

(2.16)

where $c$ does not depend upon $\nu$. 

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Goudie's (1990) stopping rule is to decide that all errors have been eliminated when

\[ \frac{L(x)}{L(x+1)} = \frac{(x+1)^{n-1}}{x^n} \]

achieves or exceeds a preassigned level \( A \). Goudie (1990) derives the probability of correcting all errors and shows that the true error level is closely approximated by \( A^{-1} \). In comparison to Nayak's (1988) procedure, the likelihood-based rule yields a small reduction in the average time taken to reach a decision.

For the usual debugging procedure, Forman and Singpurwalla (1977) propose an empirical stopping rule based upon a relative likelihood function. In a similar context Ross (1985) studies a stopping rule which takes into account the error level and provides an upper bound on the proportion of the time testing terminates with the total failure rate exceeding a prior chosen constant.
3.1 Introduction

To describe the detection, removal, and recapture of the errors in a computer program, Nayak (1988) assumed that the first occurrence times follow the Jelinski-Moranda (1972) model and that each of the errors again occur according to independent homogeneous Poisson processes with a common rate parameter \( \phi \). Let \( \pi_1, \ldots, \pi_n \) denote the sequence of programs obtained by correcting errors in an initial program \( \pi_0 \) at times \( S_1, S_2, \ldots, S_n \). A comparative method, known as back-to-back testing (e.g., Vouk, 1990), may be used to observe repeated error occurrences; e.g., if an error is detected
in \( \pi_{i-1} \) at time \( S_i \), then comparing the outputs of \( \pi_{i-1} \) and \( \pi_i \) gives the number of times this error again occurs during the remaining test time. Since \( \pi_{i-1} \) and \( \pi_i \) differ only by the correction made at time \( S_i \), any differences in their outputs are due to the fault that resides in \( \pi_{i-1} \), which has been corrected in \( \pi_i \). This method continuously replicates error detection and thus it is likely to yield more data per unit of test time than other designs.

Empirical evidence (Nagel, Scholz, and Skrivan, 1982) indicates that errors may occur with different probabilities and that errors with the highest occurrence rates are likely to be detected early. Since errors are seldom detected in the final program, inference about its reliability must be based on the error frequencies observed in the previous versions.

In this chapter we consider a family of marked Poisson process models in which the first and subsequent error occurrences are described by a Markov chain. The transition probabilities (Section 3.2) are determined by a discrete distribution \( G \) with tail probabilities \( G(i, 0) \), which are also the probabilities that the successive programs produce errors. Nayak's (1988) model assumes \( G \) is a discrete uniform distribution with mass at \( 1, 2, \ldots, v \) where \( v \) is an integer parameter that represents the number of errors in the initial
Chapter 3: Estimation

The family of models considered in Section 3.2 does not embed Nayak's model, although it is a more robust family since $G(i, \theta)$ may decrease in $i=0,1,2,...$ at different rates for particular cases.

The probability $G(R, \theta)$ that the final program produces errors necessarily depends on the number $R$ of faults eliminated during the period of testing. In this way, the problem of estimating $\tilde{G}(R, \theta)$ is analogous to that of estimating the number $\nu-R$ of remaining errors in Nayak's model. An estimate of $\tilde{G}(R, \theta)$ is $\tilde{G}(R, \hat{\theta})$ where $\hat{\theta}$ is the maximum likelihood estimator of the parameter vector $\theta$. In Section 3.3, a scaled logarithmic function of $\tilde{G}(R, \theta)$ and $\tilde{G}(R, \hat{\theta})$ is shown to have a limiting distribution identical to that of a linear function of $N^{1/2} (\hat{\theta} - \theta)$ where $N$ is the number of events observed during the period of testing. Confidence levels for estimating $G(R, \theta)$ are obtained by simulation and compared with the nominal confidence level given by the limiting normal distribution.

Similar, though usually different models have been used in a biological context (Sandland and Cormack, 1984, Huggins, 1989, Goodman, 1953) to estimate the size of animal populations. The model studied by Sandland and Cormack (1984) seems most closely related to the model studied in this chapter, although it does not describe the time
dependency of subsequent error occurrences. A more recent paper by Nayak (1991) considers a model derived from the superposition of independent homogeneous Poisson processes.

A comparison of the asymptotic variances of parameter estimators given by recapture debugging and by the repetitive-run procedure of Nagel, Scholz, and Skrivan (1982) is made in Sections 3.5 and 3.6 to determine which of these procedures efficiently uses the test time. The comparison assumes identical models for the first occurrence times and that testing is performed for time periods of equal length under the two testing strategies.

3.2 The Model and Likelihood Function

The first and subsequent error occurrences can be modeled by a Markov chain \{\( (R, Z) \) \} where \( R \) is the number of distinct errors detected by the \( i \)th epoch and \( Z \) is the error state at the \( i \)th epoch. At each epoch either a distinct new error is detected or a previously detected error again occurs. From state \( (R, Z) = (r, z), z=1,2,...,r \), a transition occurs to state \( (r+1, r+1) \) (a new error is detected) with probability \( \bar{g}(r, \theta) \) or to state \( (r, i), i \leq r \), whenever a previous error again occurs, with probability \( g(i, \theta) \). The
times $T_1, T_2, \ldots$, at which these events occur are assumed to follow a homogeneous Poisson process, where \{$(R_i, Z_i)$\} and \{$T_i$\} are assumed to be independent. As a consequence of our assumption that \{$T_i$\} is a homogeneous Poisson process, the number $N$ of events occurring in a fixed interval $(0, t)$ has a Poisson distribution with mean $\alpha t$. The model for the event occurrence times and the first and subsequent error occurrences can be specified by the initial error rate $\alpha$ and the distribution function $G(x, \theta)$ of a discrete random variable with mass $g(i, \theta)$ on the positive integers. The survivor function $\bar{G}(x, \theta) = 1 - G(x, \theta)$ determines the probability $\bar{G}(i, \theta)$ that the $i$th program produces errors.

Let $B$ denote the set of epochs at which previously detected errors again occur; that is, $B = \{i: z_i \leq r_{i-1}, i=1, 2, \ldots, n\}$. The likelihood function based on observing $N=n$ and $(R_i, Z_i, T_i)$, $i=1, 2, \ldots, n$ is

$$
\alpha^n e^{-\alpha t} \prod_{i \in B} g(z_i, \theta) \prod_{i=1}^{r} G(i-1, \theta) = L_2(\alpha) L_2(\theta), \quad (3.1)
$$

$$
(0 < t_1 < \ldots < t_n < t, \quad 1 = r_1 \leq r_2 \leq \ldots \leq r_n \leq n,
$$

$$
z_1 = 1, 2, \ldots, r_1, \quad i=1, 2, \ldots, n),
$$

where $L_2(\theta) = 1$ if $n=0$ and where $R_n = r$ is the number of distinct
errors detected during the first $n$ epochs; that is, during
the interval $(0,t)$.

The hitting frequencies $M_1, M_2, \ldots$, of the first, second,
etc., detected faults are given by $M_i = \sum_{j \in B} I(z_j = i)$, where
$I(A)$ is the indicator function of the set $A$. Since $M_i$ does
not include the first time that the $i$th detected error
occurs, we have $\sum_{i=1}^{R} M_i + R = N$. In terms of $M_1, M_2, \ldots, M_r$,
(3.1) can be written

\[ \alpha^n e^{-\alpha t} \prod_{i=1}^{m_i} \frac{1}{G(i-1, \theta)} G(i, \theta). \]  

(3.2)

It should be noted that, (3.2) reduces to the likelihood
function studied by Nayak (1988) when $G$ is a discrete uniform
distribution with mass at $1, 2, \ldots, v$ where $v$ is an integer
parameter.

Let $n_{i_0} < n_{i_1} < \ldots < n_{i,n(i)}$ denote the ordered elements of
\( \{j: z_j = i, j=1,2,\ldots,n\} \), $i=1,2,\ldots,r$. Then $S_{ij} = T_{n(i),j}$ defines
the first occurrence times $S_{10} < S_{20} < \ldots < S_{v0}$, hereafter
denoted by $S_1, S_2, \ldots, S_r$, and also the times $S_{i1} < S_{i2} < \ldots
< S_{in(i)}$ at which the $i$th detected error again occurs during
the interval $(S_i,t)$, $i=1,2,\ldots,r$.

The Jacobian of this transformation is equal to one and
thus the joint density function of $(R, M_1, M_2, \ldots, M_r, S_i,$
Chapter 3: Estimation

\( S_1, \ldots, S_r, S_{11}, S_{12}, \ldots, S_{1 \times (1)}, \ldots, S_{r1}, S_{r2}, \ldots, S_{r \times (1)} \) is

\[
\alpha^n e^{-at} \prod_{i=1}^{r} \left[ g(i, \theta) \right]^{m_i} G(i, \theta) (t-s_i) / m_i!
\]

\((n = r + \sum_{i=1}^{r} m_i, m_i=0,1,2,\ldots, 0<s_1<s_2<\ldots<s_r<t, \)

\(s_1<s_{11}<s_{12}<\ldots<s_{1 \times (1)}<t, i=1,2,\ldots,r)\)

The marginal density function of \((R, M_1, M_2, \ldots, M_r, S_1, S_2, \ldots, S_r)\) is

\[
\alpha^n e^{-at} \prod_{i=1}^{r} \left[ g(i, \theta) \right]^{m_i} G(i-1, \theta) (t-s_i)^{m_i} / m_i!
\]

\((0<s_1<s_2<\ldots<s_r<t, m_i=0,1,2,\ldots, \)

\(i=1,2,\ldots,r, n = r + \sum_{i=1}^{r} m_i, r=1,2,\ldots)\)

and is \(e^{-at}\) if \(r=0\). Since \(N = R + \sum_{i=1}^{r} M_i\), it follows from (3.3) that \((R, M_1, M_2, \ldots, M_r)\) is a sufficient statistic and thus applications of the model do not require observation of the event occurrence times. The latter information, however, may be useful for checking whether the spacings \(Y_i = S_i - S_{i-1}\), \((S_0=0)\) have nonexponential distributions.
Our discussion concerns the family of models defined by

\[
\bar{G}(i, \theta) = \exp(-\theta_1 a_{i1} - \theta_2 a_{i2}), \quad i=0,1,2,\ldots \quad (3.4)
\]

\[
g(i, \theta) = [1-\exp(-\theta_1 b_{i1} - \theta_2 b_{i2})] \exp(-\theta_1 a_{i1-1} - \theta_2 a_{i2-1}),
\]

\[
i=1,2,\ldots,
\]

where \( \theta_1 > 0, i=1,2 \) are unknown parameter values, \( b_{i1} = a_{i1} - a_{i1-1}, \ b_{i2} = a_{i2} - a_{i2-1} \), and \( \{a_{i1}\}, \{a_{i2}\} \) are known constants that satisfy \( a_{i0} = a_{20} = 0, a_{i1}, a_{i2} \) are nondecreasing in \( i=0,1,2,\ldots, \) and \( \lim a_{i1} = \lim a_{i2} = \infty \), as \( i \) tends to infinity. The latter conditions are implied by the requirement that \( \bar{G}(i, \theta) \) be a survivor function. Particular cases of (3.4) are \( a_{i1} = i, a_{i2} = i^2 \) or \( a_{i1} = i, a_{i2} = \log(1+i) \).

In Section (3.4), we note the equivalence of the model defined in the present Section to another form of the model in which the gaps \( Y_i = S_i - S_{i-1}, \ (S_0 = 0) \) between the first occurrence times have independent exponential distributions with rate parameters \( \lambda_i = \alpha G(i-1, \theta) \). In the latter context, setting \( \theta_2 = 0 \) and \( a_{i1} = i \) in (3.4) gives the log linear rate model studied by Moranda (1975) and Cox and Lewis (1966).

Since the second factor of (3.2) does not depend on \( \alpha \), the maximum likelihood estimate of \( \alpha \) is \( \hat{\alpha} = n/t \). With \( g(i, \theta) \)
given by (3.4), the log likelihood function given by the second factor of (3.2) is

\[ l_2(\theta) = \sum_{i=1}^{r} m_i \log[1-\exp(-\theta_1 b_{11} - \theta_2 b_{21})] - \theta_1 \sum_{i=1}^{r} (m_i+1) a_{1,i-1} - \theta_2 \sum_{i=1}^{r} (m_i+1) a_{2,i-1} \]  

(3.5)

If \( n \leq 1 \), then \( r \leq 1 \), \( m_i=0 \) and \( l_2(\theta) \) is constant in \( \theta \). If \( n>1 \) and \( r=1 \), then (3.5) takes its maximum value at \( \theta_1=\infty \), \( i=1,2 \). Otherwise, \( l_2(\theta) \) is concave in \( \theta \) (see Appendix A), and an estimate \( \hat{\theta} \) that maximizes (3.5) is the unique solution to

\[ \sum_{i=1}^{r} m_i b_{j1}[\exp(\theta_1 b_{11} + \theta_2 b_{21})-1]^{-1} = \sum_{i=1}^{r} (m_i+1) a_{j,i-1}, \quad (j=1,2) \]  

(3.6)

3.3 Confidence Limits

An estimate of the probability that the final program produces errors is \( \bar{G}(R, \hat{\theta}) \) where \( \hat{\theta} \) is the maximum likelihood estimate of \( \theta \). Let \( a_i = a_{1i} + a_{2i} \). Then

\[ N^{1/2} a_i^{-1} \left[ \log \bar{G}(R, \hat{\theta}) - \log \bar{G}(R, \theta) \right] \]

\[ = -N^{1/2} a_i^{-1} \left[ (\hat{\theta}_i - \theta_i) a_{1i} + (\hat{\theta}_2 - \theta_2) a_{2i} \right] \]  

(3.7)

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In a later Section we show that (3.7) has a limiting (t → \infty) normal distribution with mean zero and variance \(a' I_0^{-1} a\) where \(a'=(a_1, a_2), a_1=\lim a_{ii}/a_{11}, a_2=\lim a_{2i}/a_{11}\), as \(i \to \infty\), and \(I_0^{-1}\) is the covariance matrix of the joint limiting distribution of \(N^{1/2}(\hat{\theta}_1 - \theta_1)\) and \(N^{1/2}(\hat{\theta}_2 - \theta_2)\). This assumes that the limits \(a_1\) and \(a_2\) are finite, which is true if \(a_{1i}=i\), \(a_{2i}=i^2\) and for other models within the family defined by (3.4). The basic idea behind (3.7) is related to the \(\delta\)-method as discussed, for example, by Rao (1973, pp. 385-388).

Approximate 100(1-\(p\)) percent confidence limits (\(L_1, U_1\)) for \(G(R, \theta)\) are

\[
\begin{align*}
L_1 &= G(R, \hat{\theta}) \exp \left( -N^{-1/2} a_R \hat{\gamma}^{1/2} Z_{1-p/2} R \right) \\
U_1 &= G(R, \hat{\theta}) \exp \left( N^{-1/2} a_R \hat{\gamma}^{1/2} Z_{1-p/2} \right)
\end{align*}
\]

where \(Z_{1-p/2}\) is the upper \(1-p/2\) percentage point of the standard normal distribution. Since \(R\) diverges (\(t \to \infty\)) in probability to infinity, \(\gamma\) can be consistently estimated by

\[
\hat{\gamma} = \hat{b}' \hat{I}_0^{-1} \hat{b}, \text{ where } \hat{b}'=(\hat{b}_1, \hat{b}_2), \text{ and } \hat{b}_1 = a_{12}/a_{11}, \hat{b}_2 = a_{22}/a_{11}.
\]

The probability that the final program produces no errors during any subsequent time period of length \(y\) also depends on \(R\) and is
\[ Q_r = \exp(-\lambda_n y), \quad y > 0 \]

where, \( \lambda_n = \alpha \tilde{G}(i, \theta) \). Letting \( \hat{Q}_r = \exp(-\hat{\alpha} \tilde{G}(R, \hat{\theta}) y) \), we have

\[
N^{1/2} a_r^{-1} \log \left( \frac{\log \hat{Q}_r}{\log Q_r} \right) = N^{1/2} a_r^{-1} (\log \hat{\alpha} - \log \alpha)
- N^{1/2} a_r^{-1} \left[ (\hat{\theta}_1 - \theta_1) a_{1n} + (\hat{\theta}_2 - \theta_2) a_{2n} \right] \tag{3.8}
\]

where since \( a_r \) diverges to infinity in probability, the first term on the right of (3.8) converges in probability to zero and the second term has the same limiting normal distribution as the quantity in (3.7). Confidence limits for \(-\log Q_n\) are

\[
L_2 = (\log \hat{Q}_n) \exp \left[ -N^{-1/2} a_r \hat{\gamma}^{1/2} Z_{1-p/2} \right]
\]

\[
U_2 = (\log \hat{Q}_n) \exp \left[ N^{-1/2} a_r \hat{\gamma}^{1/2} Z_{1-p/2} \right]
\]

Table 3.1 shows simulated percentages of the time that the confidence limits cover \( \tilde{G}(R, \theta) \) and \( Q_n \). The simulated percentages fall close to the nominal 95 percent level when \( E(N) \) is large, and are sometimes about 10 percentage points below the 95 percent level when \( E(N) \) is small. Since \( \tilde{G}(R, \theta) \) and \( Q_n \) will tend to take values at the extreme endpoints of the interval \((0, 1)\) whenever \( E(N) \) is large, the quantities being estimated are necessarily extreme values as \( t \to \infty \). As
Table 3.1: Empirical confidence levels for estimating \( \bar{G}(R, \theta) \) and \( Q_x \), \( y=100 \); the nominal level is 0.95 with 1,000 replications; \( a_1 = i \), \( a_2 = i^2 \); \( E(N), E(R), Q_x \), and \( \bar{G}(c, \theta), c = E(R) \) are theoretical values not obtained by simulations.

<table>
<thead>
<tr>
<th>( E(N) )</th>
<th>( \theta_1 )</th>
<th>( \theta_2 )</th>
<th>( E(R) = c )</th>
<th>( \bar{G}(c, \theta) )</th>
<th>( Q_x )</th>
<th>( \bar{G}(R, \theta) )</th>
<th>( Q_R )</th>
<th>( \bar{G}(R, \theta) )</th>
<th>( Q_R )</th>
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<td>0.997</td>
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<td>94.7</td>
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</table>
shown in Table 3.1, the widths of confidence limits are similar in magnitude to $G(c, \theta)$ and $Q_c$ where $c=E(R)$, although the confidence limits are not designed to cover the latter quantities.

The simulations were performed by generating 1,000 realizations of the sufficient statistic $(N, R, M_1, M_2, \ldots, M_n)$. Each replicate requires generating a realization $N=n$ of a Poisson random variable having mean $\alpha t$. If $R_n \leq 1$, then $\hat{\theta}_n$ is undefined; if this event occurs, it is counted as one trial for which the confidence limits do not include the quantity being estimated. If $n > 1$, the sufficient statistic can be written as a function of i.i.d random variables $X_1, X_2, \ldots, X_n$ with density function $g(i, \theta)$. That is,

$$R_0 = 0, \quad R_1 = 1, \quad R_i = R_{i-1} + I(X_i > R_{i-1}), \quad i=2, 3, \ldots, n$$

$$M_i = \sum_{j=1}^{n} I(X_j \leq R_{j-1}, \quad X_j = i), \quad i=1, 2, \ldots, R_n.$$ 

From this it follows that the sufficient statistic can be computed sequentially from independent random variables having a uniform distribution on the interval $(0, 1)$. 

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3.4 An Equivalent Form of the Model

We now show that the following assumptions give a likelihood function identical to (3.3).

(i) The spacings \( Y_i = S_i - S_{i+1}, \) \((S_0 = 0)\) between the first occurrence times are independent random variables with density functions \( \lambda_i \exp(-\lambda_i y), y \geq 0 \) where, \( \lambda_i = \alpha G(i-1, \theta). \)

(ii) Counts \( M_i = M_i(S_i, t) \) of subsequent error occurrences have distributions determined by a collection \{\( M_i(t) \}\} of independent homogeneous Poisson processes with rate parameters \( \xi_i = \lambda_i - \lambda_{i+1}. \)

(iii) \{\( S_i \}\} and \{\( M_i(t) \}\} are independent collections of random variables.

This form of the model, originally suggested by Nayak (1988), explicitly describes the first occurrence times and thus it directly relates error recapture models to the more common reliability growth models.

By interpreting \( \lambda_i \) as the hitting rate of the remaining faults after \( i-1 \) faults have been corrected, \( \xi_i = \lambda_i - \lambda_{i+1} \) is then the change in this rate due to correcting the \( i \)th
detected error and $\bar{\xi}_1, \bar{\xi}_2, \ldots$ can be interpreted as the hitting rates of the first, second, etc., detected faults. These are average rates in that the errors that are detected first, second, etc., may vary from one repetition of the experiment to another. The model assumes that the effect of correcting faults is additive in that $\bar{\xi}_1 + \bar{\xi}_2 + \ldots + \bar{\xi}_r + \lambda_{r+1} = \lambda_i$. The parameterization $\lambda_i = \alpha G(i-1, \theta)$ assumes $\lambda_i$ decreases to zero. However, $\lambda_i$ must decrease to model reliability growth and $\bar{\xi}_i$ is not a meaningful quantity otherwise.

Let $M_i = M_i(S_i, t)$ denote the number of times the error detected at time $S_i$ again occurs during the interval $(S_i, t)$. The joint density function of $(R, S_1, S_2, \ldots, S_r, M_1, M_2, \ldots, M_r)$ is easily obtained from the fact that $M_i, M_2, \ldots, M_r$ are conditionally, given $(R, S_1, S_2, \ldots, S_r)$, independent Poisson random variables with means $\bar{\xi}_i(t-S_i)$. Since this joint density function is identical to (3.3), the model described by (i)-(iii) is equivalent to the model in Section 3.2. The intervals $(S_i, t)$ have random length and thus the unconditional distribution of $M_i$ is not Poisson; this seems to be the precise way that (i)-(iii) differ from the model studied by Sandland and Cormack (1984).
3.5 The Distributions of $R$ and $S_k$

If the $\lambda_i$ are all distinct, then $H_k(x) = P(S_k \leq x)$ can be written (Cox, 1962, p.17)

$$H_0(x) = 1, \quad H_1(x) = 1 - \exp(-\lambda x), \quad x \geq 0$$

$$H_k(x) = \sum_{i=1}^{k} \pi_{ik} \left[1 - \exp(-\lambda_i x)\right], \quad x \geq 0, \quad \pi_{ik} = \prod_{j \neq i}^{k} \frac{\lambda_j}{(\lambda_j - \lambda_i)}$$

where the weights $\pi_{ik}$ (possibly negative) have a sum equal to one. Since $P(R \geq k) = H_k(t)$, the distribution of $R$ is given by $P(R=k) = H_k(t) - H_{k+1}(t)$, $k=0,1,2,\ldots$. By noting that $R \geq k$ and $S_k \leq t$ are identical events, we also have

$$P(S_k \leq x | R \geq k) = \frac{H_k(x)}{H_k(t)}, \quad 0 < x < t$$

$$= 1, \quad t \leq x.$$ 

3.6 A Comparison of the Error Recapture and Repetitive-Run procedures.

Nagel, Scholz, and Skrivan (1982, 1984) proposed a repetitive-run procedure where, after restoring a program
to its original form, the debugging sequence is repeated, perhaps several times, using independently generated series of inputs. A run is initiated by randomly selecting an input according to a usage distribution, defined as part of the original problem specification. Each run consists of testing the program on a random series of inputs and correcting errors whenever they are detected. The order of detecting errors as well as the gaps between the error detection times may vary from one replication to another. The repetitive-run procedure has been used by Nagel, Scholz, and Skrivan (1982, 1984) and also by Dunham and Pierce (1985) to study the effect of debugging on the reliability of several programs. In this Section we compare the asymptotic variances of parameter estimates obtained under the error recapture and repetitive-run procedures.

Let $V_1, V_2, \ldots, V_n$ denote independent random vectors having the same distribution as $V=(R, S_1, S_2, \ldots, S_n)$. As in Section 3.4, the gaps $Y_i = S_i - S_{i-1}$ between the first occurrence times are assumed to be independent and have exponential density functions $\lambda_i \exp(-\lambda_i y), y \geq 0$ where, $\lambda_i = \alpha G(i-1, \theta)$. If testing in each replicate extends over a time period of length $s$, the total test time is then $ms$. To compare the error recapture and repetitive-run procedures, we assume test
periods of equal length (i.e., ms=t) and that the first occurrence times have the same distribution under both models. That is, $\lambda_i$ is assumed to be given by $\lambda_i = a \exp[-\theta(i-1)]$, where $a$ and $\theta$ are positive scalar parameters.

Since $V_1, V_2, ..., V_n$ are i.i.d random vectors, it suffices to consider the log likelihood function based on one observation of $V$, namely,

$$
\ell_j(\alpha, \theta) = \sum_{i=1}^{\infty} \log \lambda_i I(R \geq i) - \sum_{i=1}^{\infty} (\lambda_i - \lambda_{i-1}) S_i I(R \geq i) - s \sum_{i=1}^{\infty} \lambda_{i-1} I(R=i)
$$

$$
\ell_j(\alpha, \theta) = \log a \sum_{i=1}^{\infty} i I(R=i) + \alpha s \sum_{i=1}^{\infty} \left[ e^{-\lambda_i} - e^{-\lambda_{i-1}} \right] (S_i/s) I(R \geq i)
$$

$$
\ell_j(\alpha, \theta) = \theta \sum_{i=1}^{\infty} \frac{(i(i-1)}{2} I(R=i) - \alpha s \sum_{i=1}^{\infty} e^{-\lambda_i} I(R=i)
$$

The information on $(\alpha, \theta)$ given by $V$ is

$$
b_{11} = -E(\partial^2/\partial \alpha^2 \ell_j) = E(R)/\alpha^2
$$

$$
b_{12} = -E(\partial^2/\partial \alpha \partial \theta \ell_j) = -sE(R e^{-\theta R}) + s \sum_{k=1}^{\infty} C_{1k} E[(S_k/s) I(R \geq k)]
$$

$$
b_{22} = -E(\partial^2/\partial \theta^2 \ell_j) = \alpha s E(R^2 e^{-\theta R}) - \alpha s \sum_{k=1}^{\infty} C_{2k} E[(S_k/s) I(R \geq k)]
$$

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Chapter 3: Estimation

\[ C_{\nu} = k e^{-\nu} - (k-1) e^{-\nu(k-1)}, \quad C_{\nu^2} = k^2 e^{-\nu^2} - (k-1)^2 e^{-\nu^2(k-1)}, \quad k=1,2,\ldots, \]

\[ E[(S_k/s) \mathbb{I}(R \geq k)] = (1/s) \int_0^x dH_k(x) \quad (3.11) \]

This last expression follows from (3.9) and is also given by

\[ E[(S_k/s) \mathbb{I}(R \geq k)] = \sum_{i=1}^k \pi_{ik} \left( \frac{1}{\lambda_i s} \right) [1 - \exp(-\lambda_i s)] - \sum_{i=1}^k \exp(-\lambda_i s) \]

Let \((\tilde{\alpha}, \tilde{\theta})\) denote the maximum likelihood estimator of \((\alpha, \theta)\) given by \(m\) replicates in the repetitive-run procedure, and let \((\hat{\alpha}, \hat{\theta})\) denote similar estimators given by the error recapture procedure. The asymptotic variances are

\[ \text{Var}(\hat{\alpha}) = (\alpha t)^{-1} \alpha^2 \quad \text{Var}(\hat{\theta}) = (\alpha t)^{-1} e^{\theta(1-e^{-\theta})^2} \]

\[ \text{Var}(\tilde{\alpha}) = (mb)^{-1} \alpha^2 b_{22} \quad \text{Var}(\tilde{\theta}) = (mb)^{-1} E(R) \]

\[ b = b_{22} E(R) - (\alpha b_{12})^2 \]

where, in the expressions given for the variances of \(\hat{\alpha}\) and \(\hat{\theta}\), \(N\) has been replaced by \(\alpha t = E(N)\).

By substituting \(t = ms\), the asymptotic relative efficiencies are
Table 3.2: Asymptotic relative efficiencies of the repetitive-run ($\tilde{\alpha}$, $\tilde{\theta}$) and error recapture ($\hat{\alpha}$, $\hat{\theta}$) estimators; efficiencies are variance ratios with $\text{var}(\tilde{\alpha})$ and $\text{var}(\tilde{\theta})$ in the numerators.

<table>
<thead>
<tr>
<th>$\alpha$</th>
<th>$\theta$</th>
<th>$\text{E}(R_\alpha)$</th>
<th>$e(\tilde{\alpha}, \hat{\alpha})$</th>
<th>$e(\tilde{\theta}, \hat{\theta})$</th>
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<td>0.34</td>
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<td>0.28</td>
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<td>0.24</td>
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<td>0.30</td>
<td></td>
</tr>
<tr>
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<td>0.09</td>
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<td>9.5</td>
<td>0.05</td>
<td>0.09</td>
<td></td>
</tr>
</tbody>
</table>
\begin{align*}
\text{Var}(\hat{\alpha})/\text{Var}(\bar{\alpha}) &= b (\alpha s b_2)^{-1} \\
\text{Var}(\hat{\theta})/\text{Var}(\bar{\theta}) &= b [\alpha s E(R)]^{-1} e^\theta (1-e^{-\theta})^2
\end{align*}

where these quantities depend only on \( \alpha s \) and \( \theta \).

The calculated efficiencies in Table 3.2 are based on (3.11), (3.12), and the distribution of \( R \) given in Section 3.5. The relative efficiency varies from about 1 to 13 percent and thus the time period of testing for the repetitive-run procedure may need to be more than eight times that of error recapture to obtain the same amount of information. The greatest gain in information occurs when \( E(R) \) is small and this corresponds to programs for which the error detection rate is small. The low efficiencies in Table 3.2 are due to the fact that the repetitive-run procedure permits each error to be observed at most one time during each replicate.

3.7 Limiting Distributions

Let \( \hat{\theta}_n \) denote the maximum likelihood estimator of \( \theta \) based on observing \((R_1, Z_1), (R_2, Z_2), \ldots, (R_n, Z_n)\). In this section we
show that the limiting \((n \to \infty)\) distribution of \(n^{1/2}(\hat{\theta}_n - \theta)\) is identical to the limiting distribution that would be obtained if \(\hat{\theta}_n\) were computed from i.i.d random variables \(X_1, X_2, \ldots, X_n\) that have density function \(g(i, \theta)\). The asymptotic covariance matrix is then \(I_\theta^{-1} I_\theta = (I_{ij})\), where \(I_{ij} = E_{\theta} \partial^2 / \partial \theta_i \partial \theta_j [\log g(X, \theta)]\) and \(X\) has density function \(g(i, \theta)\).

In terms of \(X_1, X_2, \ldots, X_n\), the log likelihood function based on the second factor of (3.1) is

\[
l_2(\theta) = \sum_{i=1}^{n} \log g(X_i, \theta) I(X_i \leq R_{i-1}) + \sum_{i=1}^{n} \log \frac{g(X_i, \theta)}{g(i, \theta)} I(X_i > R_{i-1})
\]

where \(R_0 = 0, R_1 = 1,\) and \(R_k = R_{k-1} + I(X_k > R_{k-1}), k=2,3,\ldots\)

Let \(l(\theta) = \sum_{i=1}^{n} \log g(X_i, \theta)\) and note that

\[
l_2(\theta) - l(\theta) = \sum_{i=1}^{n} I(X_i > R_{i-1}) [\log g(X_i, \theta) - \log \frac{g(X_i, \theta)}{g(i, \theta)}]
\]

To simplify this last expression, let \(i=n_1 < n_2 < \ldots < n_r \leq n\) denote the epochs at which \(X_i > R_{i-1}\). Conditionally, given \(N_1 = n_1, N_2 = n_2, \ldots, N_r = n_r,\) and \(R_n = r,\) the set \(X_{n(1)}, X_{n(2)}, \ldots, X_{n(r)}\), consists of independent random variables with density functions \(h_k(i, \theta) = g(i, \theta) / g(k-1, \theta), i=k,k+1,\ldots\)
Since the conditional distribution of $X_{N(1)}, X_{N(2)}, \ldots, X_{N(r)}$, depends on $N_1, N_2, \ldots, N_r$ and $R_n$ only through $R_n$, it is simpler to let $Y_1, Y_2, \ldots, Y_r$ be independent random variables with density functions $h_k(i, \theta), k=1, 2, \ldots, r$. Then

$$l'_2(\theta) - l(\theta) = \sum_{k=1}^{R_n} \log h_k(Y_k, \theta)$$

(3.13)

and

$$-\frac{\partial^2}{\partial \theta_p \partial \theta_q} \log h_k(i, \theta) = b_{p1} b_{q1} \left[ \exp(\theta_1 b_{11} + \theta_2 b_{21}) - 1 \right]^{-2} \exp(\theta_1 b_{11} + \theta_2 b_{21}) \quad (p, q=1, 2, \ i=k, k+1, \ldots)$$

By using the relations $e^x(e^x - 1)^{-2} = (e^x + e^{-x} - 2)^{-1}$ and $e^x + e^{-x} - 2 \geq x^2$, we obtain

$$| -\frac{\partial^2}{\partial \theta_p \partial \theta_q} \log h_k(i, \theta) | \leq (\theta_p \theta_q)^{-1} \quad (p, q=1, 2) \quad (3.14)$$

Our remaining discussion requires the notation

$$U_1 = \partial / \partial \theta_1 l_2(\theta), \quad V_1 = \partial / \partial \theta_1 l(\theta),$$

$$w_{ik} = \partial / \partial \theta_i \log h_k(Y_k, \theta), \quad (k=1, 2, \ldots, r)$$
Since $E(W_{ik}) = 0$ and $E(W_{ik}^2) = -E[\partial^2/\partial \theta_i \partial \theta_i \log h_k(Y_k, \theta)]$
we have from (3.13) and (3.14) that

$$Var(U_i - V_i) = E(\sum_{k=1}^{R_n} W_{ik}^2)$$

$$\leq \theta_i^{-2} E(R_n), \quad (i=1,2)$$

Thus $n^{-1/2}(V_i - U_i)$ converges ($n \to \infty$) in probability to zero providing limit $n^{-1} E(R_n) = 0$.

The representation of $R_n$ given earlier in Section 3.3 implies

$$R_n = 1 + \sum_{i=2}^{n} I(X_i > R_{i+1})$$

$$E(R_n) = 1 + \sum_{i=1}^{n-1} E[G(R_i, \theta)] \quad (3.15)$$

where, since $n^{-1} E(R_n)$ is an average of the terms on the right of (3.15), it suffices to show that $\lim E[G(R_n, \theta)] = 0$. This limit is easily obtained from the fact that $R_n$ tends to infinity in probability, and thus further details are omitted.

Since $V=(V_1, V_2)$ is a linear function of i.i.d random variables, the preceding discussion shows that $n^{-1/2} V$ and $n^{-1/2} U$ are asymptotically equivalent and that $n^{-1/2} U$ has a limiting bivariate normal distribution with mean vector zero.
and covariance matrix $I_\theta$. The usual consistency and Taylor series arguments (e.g., Cox and Hinkley, 1974) imply that $n^{1/2}(\widehat{\theta}_n - \theta)$ has a limiting bivariate normal distribution with covariance matrix $I_\theta^{-1}$. Since $\{T_n\}$ and $\{R_i, Z_i\}$ are independent, $n^{1/2}(\widehat{\theta}_n - \theta)$ and $n^{1/2}(n/T_n - \alpha)$ are also independent. By Theorem 8.1 of Serfozo (1975), (see also Karr, 1986, p. 406) $N^{1/2}(\widehat{\alpha} - \alpha)$ and $N^{1/2}(\widehat{\theta} - \theta)$ have asymptotically ($t \to \infty$) independent normal distributions with variance $\alpha^2$ and covariance matrix $I_\theta^{-1}$, respectively.
Chapter 4

Application to Repetitive-Run Data

4.1 Introduction

Nagel, Scholz and Skrivan (1984) observed that many models have been introduced in the literature, but without a clear statement about the mathematical and statistical foundations that motivated the model. They pointed out that for a consistent theoretical foundation, a deeper understanding of the process is needed. As an attempt in this direction, they conducted a series of experiments consisting of simulations conducted on code prepared according to a set of requirements and executed with randomly selected inputs. The code is initialized to an original state then tested on randomly generated inputs. Errors are corrected as they are encountered until a stopping rule is satisfied. Replication is introduced by repeating the entire process from
initialization. The following section gives further details concerning the testing process.

4.2 Description

The simulations are initiated by generating random inputs according to a distribution, called the usage distribution, defined as part of the original problem specification. After correcting an error in the program, other inputs are generated independently and the process is repeated. If for some execution an error is indicated, the error is recorded together with the number of executions since the last error, and the error is corrected.

The simulation begins with the program in its initial state. This state is reached when the program successfully compiles and correctly executes a number of predetermined test cases. These tests are defined as static tests for a given specification and the program must pass these static tests as well as successfully execute the input causing failure before simulation can be reinitiated. Once reinitiated, the process is repeated error by error until a stopping rule is satisfied. Termination of the experiment occurred when an error is detected that is too costly to fix.
or an upper bound on the number of inputs is reached, whichever comes first.

Traditional tests on software force the experimenter to predict the reliability of the program from a single manifestation of the error process. To observe different realizations of the order of detecting errors the program is restored to its original state and testing is repeated, using another randomly selected input sequence. The experimental flow for each run is exactly the same except for the consequences of using different inputs. Each run may generate different random errors in different orders and with varying spacings between the error detection times.

4.3 Software Error Categories

The following list describes the category of different possible software errors in a program:

**Computational Errors**

- Incorrect operand in logical sequence
- Incorrect use of parenthesis
- Sign convention error
- Unit or data conversion error
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Computation produces an over/under flow
Incorrect/inaccurate equation used
Precision loss due to mixed mode
Missing computation
Rounding or truncation error

Logical Errors

Incorrect Operand in logical expression
Logic activities out of sequence
Wrong variable being checked
Missing logic or condition tests
Too many/few statements in loop
Loop iterated incorrect number of times
Duplicate logic

Data Input Error

Invalid input read from correct data file
Input read from incorrect data file
Incorrect input format
Incorrect format statement referenced
End of file encountered prematurely
End of file missing
Data Handling Errors

Data file not rewound before reading
Data initialization not done
Data initialization done improperly
Variable referred to by the wrong name
Bit manipulation done incorrectly
Incorrect variable type
Data packing/unpacking error
Sorting error
Subscripting error

Data Output Error

Data written on wrong file
Data written according to the wrong format statement
Data written in wrong format
Data written with wrong carriage control
Incomplete or missing output
Output field size too small
Line count or page eject problem
Output garbled misleading
Interface Errors
  wrong subroutine called
  Call to subroutine not made or made in wrong place
  Subroutine arguments not consistent in type, units
  Subroutine called is nonexistent
  Software/data base interface error
  Software/Software interface error

Data Definition Errors
  Data not properly defined/dimensioned
  Data referenced out of bounds
  Data being referenced at incorrect location
  Data pointers not incremented properly

Data Base Errors
  Data not initialized in data base
  Data initialized to incorrect value
  Data units are incorrect

Operation Errors
  Operating system error
  Hardware error
  Operator error
Test execution error
User misunderstanding/error
Configuration control error

Documentation Errors
User manual
Interface specification
Design specification
Requirements specification
Test documentation

4.4 The Likelihood Function

The repetitive-run procedure yields several realizations of \((R, S_1, S_2, \ldots, S_n)\), where \(R\) is the number of failures observed during an interval \((0, s)\) and \(S_1, S_2, \ldots, S_n\) are the ordered failure times. As before, the gaps \(Y_i = S_i - S_{i-1} (S_0=0)\), \(i=1, 2, \ldots\) are assumed to be independent random variables with exponential density functions

\[
f(y) = \lambda_i \exp(-\lambda_i y), \quad y \geq 0 \quad (4.1)
\]

where

\[
\lambda_i = \alpha G(i-1, \theta) \quad (4.2)
\]
Although this parameterization is identical to the one studied in Chapter 3, the two models differ in fundamental ways; e.g., except for the fact that \( R \) has a discrete distribution, the observations here have continuous distributions whereas the sufficient statistic for the error recapture model has a discrete distribution.

Let \( s_i \) denote the length of the period of testing in the \( i \)th replicate and let \( V_i = (R_{i1}, Y_{i1}, Y_{i2}, \ldots, Y_{iR_{i1}-1}) \), \( i = 1, 2, \ldots, m \) denote the observations obtained in \( m \) replicates, where \( Y_{iR_{i1}-1} \) is the spacing between \( S_{i1} \) and \( S_{iR_{i1}} \) with \( S_{i1}, S_{i2}, \ldots, S_{iR_{i1}} \) being the times at which errors are detected in the \( i \)th replicate. Under the assumption that \( V_1, V_2, \ldots, V_n \) are independent random vectors and that \( Y_1, Y_2, \ldots, \) have independent exponential distributions, the full likelihood function is

\[
L(\alpha, \theta) = \prod_{i=1}^{m} \prod_{j=1}^{R_{i}} \prod_{j=1}^{R_{i}+1} \exp(-\lambda_j y_{ij}) \tag{4.3}
\]

where

\[
\lambda_j = \alpha \exp(-\theta_1 a_{1,j-1} - \theta_2 a_{2,j-1}), \quad j = 1, 2, \ldots, \tag{4.4}
\]

and

\[
a_{1,j} = j, \quad a_{2,j} = j^2
\]
Let $D_k = \{i : R_i \geq k\}, \ k = 1, 2, \ldots$ and note that $y_{ik}$ is an observed quantity whenever $i \in D_k$. By substituting from (4.4), the log likelihood can be written as

$$\ln L(\alpha, \theta) = A_1 \ln \alpha - \theta_1 B_1 - \theta_2 B_2 - \alpha C_1 - \alpha \bar{G}(1, \theta) C_2 - \alpha \bar{G}(2, \theta) C_3 - \ldots$$

(4.5)

where

$$A_1 = \sum_{i=1}^{m} R_i$$

$$B_1 = \sum_{i=1}^{m} \sum_{j=1}^{R_i} a_{i,j-1}$$

$$B_2 = \sum_{i=1}^{m} \sum_{j=1}^{R_i} a_{i,j-1}$$

$$C_k = \sum_{D_k} y_{ik} \ \ (k = 1, 2, \ldots)$$

$$\bar{G}(k, \theta) = \exp(-\theta_1 a_{1k} - \theta_2 a_{2k})$$

$$\bar{G}(0, \theta) = 1. \quad k = 1, 2, \ldots, \ (4.6)$$

An estimate $(\hat{\alpha}, \hat{\theta})$ that maximizes (4.5) is the solution to the system of equations.
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\[ A_1/\alpha - \left[ C_1 + G(1, \theta) C_2 + G(2, \theta) C_3 + \ldots \right] = 0 \quad (4.7) \]

\[ - B_1 + \frac{A_1 \left[ a_{11} G(1, \theta) C_2 + a_{12} G(2, \theta) C_3 + \ldots \right]}{[C_1 + G(1, \theta) C_2 + G(2, \theta) C_3 + \ldots]} = 0 \quad (4.8) \]

\[ - B_2 + \frac{A_1 \left[ a_{21} G(1, \theta) C_2 + a_{22} G(2, \theta) C_3 + \ldots \right]}{[C_1 + G(1, \theta) C_2 + G(2, \theta) C_3 + \ldots]} = 0 \quad (4.9) \]

Since (4.8) and (4.9) depend only on \( \theta \), \( \hat{\theta} \) can be obtained by solving the last two equations and then substituting \( \hat{\theta} \) into (4.7) to get \( \hat{\alpha} \).

Under the stopping rule described earlier in this chapter, the period of observation in each replicate ends with a failure. A common data modification in such cases (e.g., Pedersen, 1979) is to treat the period of observation as being a fixed interval \((0, s_i)\), where \( s_i \) is actually the occurrence time of the last observed event.

4.5 A Test of Fit

Each replicate may produce a different final program, so the main parameter of interest is less clear than for the
error recapture procedure. However, the constants \( \{a_{1i}\} \) and \( \{a_{2i}\} \) must be selected prior to fitting a model.

Consider the case \( a_{1i} = i \), \( a_{2i} = i^2 \) and the following two submodels of (4.4)

\[
\lambda_{1i} = \alpha \exp(-\theta_1 i), \quad \alpha > 0, \quad \theta_1 > 0 \tag{4.10}
\]

\[
\lambda_{1i} = \alpha \exp(-\theta_2 i^2), \quad \alpha > 0, \quad \theta_2 > 0 \tag{4.11}
\]

and we determine whether an adequate fit can be achieved with the latter.

Let \( L(\alpha, \theta_1, 0) \) denote the likelihood function corresponding to (4.10) and similarly let \( L(\alpha, 0, \theta_2) \) be the likelihood function corresponding to (4.11). Then

\[
\ln L(\alpha, \theta_1, 0) = A_1 \ln \alpha - \theta_1 B_1 - \alpha C_1 - \alpha G(1, \theta_1, 0) C_2 - \alpha G(2, \theta_1, 0) C_3 - \ldots \tag{4.12}
\]

\[
\ln L(\alpha, 0, \theta_2) = A_1 \ln \alpha - \theta_2 B_2 - \alpha C_1 - \alpha G(1, 0, \theta_2) C_2 - \alpha G(2, 0, \theta_2) C_3 - \ldots \tag{4.13}
\]

The estimating equations corresponding to (4.10) and (4.11) are obtained by setting \( \theta_2 = 0 \) in (4.8) and \( \theta_1 = 0 \) in (4.9).
Consider the following subsets of the total set of parameter values,

\[ \Omega' = \{ (\alpha, \theta_1, \theta_2) : \alpha > 0, \theta_1 > 0, \theta_2 > 0 \} \]

\[ \Omega_{10} = \{ (\alpha, \theta_1, \theta_2) : \alpha > 0, \theta_1 > 0, \theta_2 = 0 \} \]

\[ \Omega_{01} = \{ (\alpha, \theta_1, \theta_2) : \alpha > 0, \theta_1 = 0, \theta_2 > 0 \} \]

where \( \Omega_{10} \) and \( \Omega_{01} \) denote the parameter sets corresponding to models (4.10) and (4.11), respectively. In this context, a test of fit of the model defined by (4.10) is a test of the composite null hypothesis \( \theta_2 = 0 \) versus the one sided alternative \( \theta_2 > 0 \).

Let \( W \) be the likelihood ratio statistic for the simpler model (4.10). For a large number of replicates and under suitable regularity conditions (Cox and Hinkley, 1974), the null distribution of the likelihood ratio statistic \( W \) is approximately a chi-square distribution with 1 degree of freedom where \( W \) is given by

\[ e^{-1/2} \hat{W}_i = \frac{\max_{(\alpha, \theta_1, \theta_2) \in \Omega_{10}} L(\alpha, \theta_1, \theta_2)}{\max_{(\alpha, \theta_1, \theta_2) \in \Omega} L(\alpha, \theta_1, \theta_2)} \quad (4.14) \]
and \( \Omega = \Omega_0 + \Omega' \) is parameter space for the full model. Similarly, with \( \Omega = \Omega_0 + \Omega' \) the likelihood ratio statistic \( W_2 \) corresponding to (4.11) can be defined in a form like (4.14).

4.6 Numerical Example

The data in Appendix B was collected by Nagel, Scholz, and Skrivan (1984) and are times between failures observed in 50 replicates when using the repetitive-run procedure. A program run consists of testing the program on a randomly selected input series and correcting errors whenever they are detected. The runs are replicated by restoring the program to its original form and testing it again on another randomly selected input series. Each run terminates when an error is detected that is too costly to fix or when an upper bound on the length of an input series is reached.

Part of the analysis of the data in Appendix B given by Nagel, Scholz, and Skrivan (1984) assumes that the waiting times (i.e., numbers of executions) between failures have independent geometric distributions with parameters \( \{p_i\} \). They define \( p_i \) as the conditional probability that a random execution of the program will result in an error given that \( i-1 \) errors have been corrected. By plotting estimates of log
Chapter 4: Application 63

$p_i, \ i=1,2,\ldots$, they conclude that \( \log p_i \) is linear in \( i, i=1,2,\ldots \)

As noted by Miller (1988), the geometric distributions can be approximated by exponential distributions whenever the quantities \( p_i \) are small. A continuous approximation to the geometric model assumes that the waiting times have independent exponential distributions with rate parameters \( \lambda_i = p_i \). Then \( \log p_i \) is linear in \( i, i=1,2,\ldots \), only if \( \lambda_i \) takes the form \( \lambda_i = \alpha \exp[-\theta(i-1)], \alpha > 0 \), where \( \theta \) is positive whenever reliability improves as faults are removed from a program.

The log linear rate model was proposed by Moranda (1975) and also appears in Cox and Lewis (1966).

Using the data in Appendix B and fitting the more general form of the model described in Sections 4.4 and 4.5, we obtain the estimates \((\hat{\alpha}, \hat{\theta}_1, \hat{\theta}_2) = (0.37, 1.07, 0.0)\). The maximum of the log likelihood (4.5) is -2232.92. The estimates \((\hat{\alpha}, \hat{\theta}_1) = (0.37, 1.07)\) and \((\hat{\alpha}, \hat{\theta}_2) = (0.064, 0.132)\) maximize the likelihood functions corresponding to the simpler models (4.10) and (4.11), respectively. The respective maximum values of the log likelihood are -2232.92 and -2356.41.

Further the necessary calculations for \( W_1 \) and \( W_2 \) yield their respective values as \( W_1 = 0.0 \) and \( W_2 = 246.98 \). The upper tail value of the chi-square distribution with one degree of freedom at the 5 percent level of significance is 3.84. Since
$W_2$ exceeds this value, the full model (4.4) is preferred over the corresponding simpler model (4.11), and the small value of $W_1$ indicates no difference in the fit of models (4.4) and (4.10). Further, since (4.11) is rejected and (4.10) is accepted by the likelihood ratio test, (4.10) (i.e., the log linear rate model) is the best fitting model.

The Newton-Raphson method was used to solve the nonlinear estimating equations (4.8) and (4.9). We began with an initial approximation $(\theta_{10}, \theta_{20})$ in the positive quadrant close to the origin and repeatedly improved it. At the $i$th stage when there is no further improvement, $(\theta_{1i}, \theta_{2i})$ is the approximation to $(\theta_1, \theta_2)$.

To check whether the estimates found above actually maximize the likelihood, the values of likelihood function were calculated inside a fairly large grid of $(\theta_1, \theta_2)$ values. These values, not shown here, indicate that the calculated values of the estimates yield the maximum of the likelihood function.

Our purpose in this section is not that of proving that any particular model will give a better fit than another model. Since the rate at which errors are detected and eliminated is likely to vary from one program to another, the goodness of fit of a model must generally be examined to avoid overly optimistic prediction of reliability.
In comparison to the log linear rate model (4.10), the above results indicate that no improvement is realized by including the extra term $a_{ii} = i^2$. This conclusion was also reached by Nagel, Scholz, and Skrivan (1984), not only for the data in Appendix B, but also for other programs they tested. Miller (1988) has pointed out, however, that the log linear rate model is unlikely to be a universal model. The example he gives is a system consisting of two software modules which, if tested separately, the failure times are assumed to follow a log linear rate model. If the two modules are then combined and tested as one system, he argues that the failure rates cannot then exhibit a log-linear pattern.
Appendix A

Lemma A: Show that $l_i(\theta)$ in equation (3.5) is concave in $\theta_i > 0$, $i = 1, 2$.

Proof: Let $U = (u_{pq})$, $p, q = 1, 2$ where

$$U_{pq} = - \frac{\partial^2}{\partial \theta_p \partial \theta_q} l_i(\theta)$$

$$= \sum_{i=1}^{r} b_{pi} b_{qi} C_i$$

$$C_i = m_i \exp(\theta_1 b_{1i} + \theta_2 b_{2i}) \{\exp(\theta_1 b_{1i} + \theta_2 b_{2i}) - 1\}^{-2}$$

We have $u_{ii} \geq 0$ and

$$(u_{11} u_{22} - u_{12}^2) \left( \sum C_i \right)^{-2} = \left( \sum b_{1i}^2 f_i \right) \left( \sum b_{2i}^2 f_i \right) - \left( \sum b_{1i} b_{2i} f_i \right)^2$$

where $f_i = C_i / \sum C_i$. Since $b_{1i} > 0$ and $b_{2i} > 0$, Holder's inequality implies $u_{11} u_{22} - u_{12}^2 \geq 0$. Thus $U$ is positive semidefinite and the latter implies that $l_i(\theta)$ is concave in $\theta_i > 0$, $i = 1, 2$, (e.g., Beltrami, 1970, p. 74).

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Appendix B

Table B.1: Times between the occurrence of program errors in m=50 replicates. (Nagel, Scholz, Skrivan, 1984)

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Bibliography


Biography

Rajan Gupta was born on August 22, 1956 in Bijnor town of Uttar Pradesh province of India. He graduated from University of Delhi in May 1975 with a B.Sc. (Hons.) in Mathematics, and M. Sc. Mathematical Statistics in May 1977. Soon after he began his career as Junior Research Scientist at Delhi Telephones. In 1982, he emigrated to Canada, where he carried out graduate studies at the University of Manitoba, and received a Master’s degree in Applied Statistics in May 1984. During 1990-91, he was Asst. Professor of Statistics, at Embry-Riddle Aeronautical University, Daytona Beach. His former position include Biostatistician with McNeil Pharmaceuticals. He was awarded a Ph.D. in Statistics from Old Dominion University in December, 1991.

He has coauthored two papers on software reliability. This research was sponsored by the Fault-Tolerant Systems Branch at NASA Langley Research Center, grant number NAG-1-835.

His future interests include statistical methodology in biopharmaceutical research such as oncology, cardiovascular and AIDS research.

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