CHAPTER 16

SOFTWARE RELIABILITY SIMULATION

Robert C. Tausworthe
Jet Propulsion Laboratory
California Institute of Technology

Michael R. Lyu
Bellcore

1  INTRODUCTION

Previous chapters discuss the opportunities and benefits of SRE throughout the entire life cycle, from requirements determination through design, implementation, testing, delivery, and operations. Properly applied, SRE is an important positive influence on the ultimate quality of all life cycle products. The set of life cycle activities and artifacts, together with their attributes and interrelationships, that are related to reliability comprise what we here refer to as the reliability process. The artifacts of the software life cycle include documents, reports, manuals, plans, code, configuration data, test data, ancillary data, and all other tangible products.

Software reliability is dynamic and stochastic. In a new or upgraded product, it begins at a low figure with respect to its new intended usage and ultimately reaches a figure near unity in maturity. The exact value of product reliability, however, is never precisely known at any point in its lifetime.

The software reliability models described in Chapter 4 attempt to assess expected reliability or future operability using observed failure data and statistical inference techniques. Most of these only treat the exposure and handling of failures during testing or operations. They are restricted in their life cycle scope and adaptability to general use for a number of reasons, including their foundation on oversimplified assumptions and their primary focus on testing and operations phases.

Modelers have traditionally imposed certain simplifying assumptions in order to obtain closed-form, idealized approximations of software reliability. Some modelers may have relaxed an assumption here or there in attempts to provide more generality, but as models become more and more realistic, the likelihood of obtaining simple analytic solutions plunges to impossibility.

This situation is not a roadblock to software reliability modeling, but perhaps a boon, in that it forces us to the apply modern technology to the problem. Computer models are not subject to the oversimplifications required to obtain closed-form results. Numerical methods can cope with models having very realistic and complex representations of project processes, software artifacts, and the development and operational environments.

Reliability models attempt to capture the structure and interrelationships among artifacts, activities, resources, quality, and time. However, this Chapter is mainly about computational techniques for modeling reliability behavior. It does not present a tool for operational situations that you may immediately apply, off-the-shelf. It does present concepts for generalized tools that mirror reliability processes. We hope that the material presented will demonstrate the power, flexibility, and potential benefits that simulation techniques offer, together with methods for representing artifacts, activities, and events of the process, and techniques for computation.

2 RELIABILITY SIMULATION

A simulation model describes a system being characterized in terms of its artifacts, events, interrelationships, and interactions in such a way that one may perform experiments on the model, rather than on the system itself, ideally with indistinguishable results.

1 Suitable extensions of the concepts of this Chapter may also apply to simulation of other quality profiles, such as availability.
Simulation presents a particularly attractive computational alternative for investigating software reliability because it averts the need for overly restrictive assumptions and because it can model a wider range of reliability phenomena than mathematical analyses can cope with. Simulation does not require that test coverage be uniform, or that a particular fault-to-failure relationship exist, or that failures occur independently, if these are not actually the case.

But power and generality are ineffective where ignorance reigns. Scientific philosophy teaches us to seek the simplest models that explain poorly understood phenomena. For example, when we do not understand how fault attributes relate to consequent failures, we may as well simplify the model by assuming that faults produce independent failures, at least until our experiments prove otherwise.

But objective validation of even a simple reliability model may be problematic, because controlled experiments, while easy to simulate, will be impossible to conduct in practice. However, if we can build an overall model upon simple and plausible submodels that together integrate cleanly to simulate the phenomenon under study, then we may gain some aggregate trust from the combined levels of confidence we may have in the constituent submodels.

2.1 The Need for Dynamic Simulation

Reliability modeling ultimately requires good data. But software projects do not always collect data sets that are comprehensive, complete, or consistent enough for effective modeling research or model application. Additionally, industrial organizations are reluctant to release their reliability data for use by outside parties. Further, data required for software reliability modeling in general, and execution time models in particular, seem to be even more difficult to collect than other types of software engineering data. Even when data are available, they are rarely suitable for isolation of individual reliability drivers.

In practicality, isolating the effects of various driving factors in the life cycle requires exploring a variety of scenarios “with other factors being the same.” But no real software project can afford to do the same project several times while varying the factors of interest. Even if they could, control and repeatability of factors would, at best, be questionable. A project may attempt, of course, to utilize data from past experiences, “properly adjusted” to appear as if earlier realizations of the current project were available. However, in view of the current scarcity of good, consistent data, this may not be realistic.

Reliability modelers thus never have the real opportunity to observe several realizations of the same software project. Nor are they provided with data that faithfully match the assumptions of their models. Nor are they able to probe into the underlying error and failure mechanisms in a controlled way. Rather, they are faced not only with the problem of guessing the form and particulars of the underlying random processes from the scant, uncertain data they possess, but also with the problem of best forecasting future reliability using that data.

Since good data sets are so scarce, one purpose of simulation is to supply carefully controlled, homogeneous data or software artifacts having known characteristics for use in evaluating the various assumptions upon which existing reliability models have been built. Since actual software artifacts (such as faults in computer programs) and processes (such as failure and fault removal) often violate the assumptions of analytic software reliability models, simulation can perhaps provide a better understanding of such assumptions and may even lead to a better explanation of why some analytic models work well in spite of such violations.

But while simulation may be useful for creating data sets for studying other, more conventional reliability models, it cannot provide the necessary attributes of the phenomena being modeled without real information derived from real data collected from real projects, past and present.

A second use of simulation, then, is in forecasting the driving influences of a real project. Models that can faithfully portray the relative\(^2\) consequences of various proposed alternatives can potentially assess the relative advantages of the candidates. Once a project sufficiently characterizes its processes, artifacts, and utilization of resources, then trade-offs can indicate the best hopes for project success.

Simulation can mimic key characteristics of the processes that create, validate, and revise documents and code. It can mimic faulty observation of a failure when one has, in fact, occurred, and, additionally, can mimic system outages due to failures. Furthermore, simulation can distinguish faults that have been removed from those that have not, and thus can readily reproduce multiple failures due to the same as-yet unrepaired fault.

\(^2\)Absolute accuracy is not required for many trade-off studies. Factors which remain the same for all alternative choices do not affect the relative advantage analyses.
Some reliability subprocesses may be sensitive to the passage of execution time (e.g., operational failures), while others may depend on wall-clock, or calendar, time (e.g., project phases); still others may depend on the amount of human effort expended (e.g., fault repair) or on the number of test cases applied. A simulator can relate model-pertinent resource dependencies to a common base via resource schedules, such as workforce loading and computer utilization profiles.

2.2 Dynamic Simulation Approaches

_Simulation_ in this Chapter refers to the technique of imitating the character of an object or process in a way that permits one to make quantified inferences about the real object or process. A _dynamic_ simulation is one whose inputs and observables are events and parameter values, either continuous or discreet, that vary over time. The formal characterization of the object or process is the _model_ under study.

When the form of the model changes over time, adapting to actual data from an evolving project, the simulation is _trace-driven_. If parameters and interrelationships are static, without trace data, the simulation is _self-driven_.

The observables of interest in reliability engineering are usually discrete integer-valued quantities (e.g., counts of errors, defects, faults, failures, lines of code) that occur, or are present, as time progresses. Studies of reliability in this context belong to the general field of discrete-event process simulation. Readers wishing to learn more about discrete-event simulation methods may consult [?].

One approach to simulation produces actual physical artifacts and portions of the environment according to factors and influences believed to typify these entities within a given context. The artifacts and environment are allowed to interact naturally, whereupon one observes the actual flow of occurrences of activities and events. We refer to this approach as _artifact-based_ process simulation, and discuss it in detail in Section 16.??.

The other reliability simulation approach [?, ?] produces time-line imitations of reliability-related activities and events. No artifacts are actually created, but are modeled parametrically over time. The key to this approach is a _rate-based_ architecture, in which phenomena occur naturally over time as controlled by their frequencies of occurrence, which depend on driving factors such as numbers of faults so far exposed or yet remaining, failure criticality, workforce level, test intensity, and execution time.

Rate-based event simulation is a form of modeling called _system dynamics_, with the distinction that the observables are discrete events randomly occurring in time. Systems dynamics simulations are traditionally non-stochastic and non-discrete. But as will be shown, extension to a discrete stochastic architecture is not difficult. For more information on the systems dynamics technique, see [?].

The use of simulation in the study of software reliability is still formative, experimental, speculative, controversial, and in the proof-of-concept stage. Although simulation models conceptually seem to hold high promise both for creating data to validate conventional models and for generating more realistic forecasts than analytic models do, the evidence to support these hypotheses is currently rather scant and arguable. There are some favorable indications of potential, however, to be discussed.

3 THE RELIABILITY PROCESS

Because of the lack of good data, past efforts in modeling the reliability process have perhaps been, to some, daunting tasks with uncertain benefits. However, as projects are now becoming increasingly better instrumented, data availability will eventually make this modeling entirely feasible and accurate. Some simulations of portions of the reliability process where measurements are routinely taken are practical.

The reliability process, in generic terms, is a _model_ of the reliability-oriented aspects of software development, operations, and maintenance. Since every project is different, describing an “average” case requires characterizing behavior typical of a class, with variations according to product, situation, environmental, and human factors. In this Section, we shall attempt to describe some of the more qualitative aspects of the software reliability process. Quantitative profiles will then follow in subsequent Sections.

3.1 The Nature of the Process

Quantities of interest in a project reliability profile include artifacts, errors, inspections, defects, corrections, faults, tests, failures, outages, repairs, validations, retests, and expenditures of resources, such as CPU time,
staff effort, and schedule time.

A number of factors hold varying degrees of influence over these interrelated elements. Influences include relatively static entities, such as product requirements, as well as other, more dynamic factors, such as the order and concurrency among activities in the process. One would hope to quantify trends, correlations, and perhaps causal factors from data gathered from previous similar projects that would be of current use. Even when formal data are not available, project personnel may often be able to apply their experiences to estimate many of the parameters of the reliability profile needed for modeling.

We will aggregate activities relating to reliability into typical classes of work, such as

1. **CONSTRUCTION** generates new documentation and code artifacts, while human mistakes inject defects into them. Activities divide into separate documentation and coding subphases, and perhaps further divide into separate work packages for constructed components.

2. **INTEGRATION** combines reusable documentation and code components with new documentation and code components, while human mistakes may create further defects. Integration activities divide into separate documentation and code integration subphases, and perhaps further divide into separate work packages according to the build architecture.

3. **INSPECTION** detects defects through static analyses of software artifacts. Inspections also divide into separate document and code subphases mirroring construction. Inspections may fail to recognize defects when encountered.

4. **CORRECTION** analyzes and removes defects, again in document and code correction subphases. Corrections may be ineffective, and may inject new defects.

5. **PREPARATION** generates test plans and test cases, and readies them for execution.

6. **TESTING** executes test cases, whereupon failures occur. Some failures may escape observation, while others may initiate system outages. Failure criticality determinations are made.

7. **IDENTIFICATION** makes failure-to-fault correspondences and fault category assignments. Each fault may be new or previously encountered. Identification may erroneously identify the cause of a failure.

8. **REPAIR** removes faults (not necessarily perfectly) and possibly introduces new faults.

9. **VALIDATION** performs inspections and static checks to affirm that repairs are effective, but may err in doing so; it may also detect that certain repairs were ineffective (i.e., the corresponding faults were not totally removed), and may also detect other faults.

10. **RETEST** executes test cases to verify whether specified repairs are complete. If not, the defective repair is marked for re-repair. New test cases may be needed. Retests may err in qualifying a fault as repaired.

### 3.2 Structures and Flows

Work in a project generally flows in the logical precedence of tasks listed above. But some activities may take place concurrently and repeatedly, especially in rapid prototyping, concurrent engineering, and spiral models of development. Models of behavior cannot ignore project paradigms, but must adapt to them.

Events occur and activities take place through the application of resources over intervals of time. No progress in the life cycle results unless activities consume resources. As examples, a code component of 500 lines of code (LOC) may require an average of $W_c$ work hours and $H_c$ CPU hours per LOC to develop, to be expended between the schedule times $t_1$ and $t_2$; testing the component may require $W_t$ work hours to generate and apply test cases and $H_t$ CPU hours per test case to execute, scheduled for the time interval between $t_3$ and $t_4$; and a repair activity may require $W_r$ work hours and $H_r$ CPU hours to complete, during the interval between times $t_5$ and $t_6$.

The project resource schedule is essential for managing the reliability process. It defines the project activities, products, flow of work, and allocation of resources. It thus reflects the planned development methodology, management and engineering decisions, and environment constraints.
Projects may, of course, measure failure profiles and other reliability data without recording the schedule and resource actual performance details. However, they will not be able to extract quantitative relationships among reliability and management parameters without these details.

The data essential to a process schedule define the resources and resource levels that are applied throughout the project duration. Schedule items may, for example, appear as tuples, such as

\[(\text{resource, event \_process, rate, units, } t_{\text{begin}}, t_{\text{end}})\]

Together, the items relate the utilization of all resources at each instant of time throughout the process. The tuple specifies that the named \text{event \_process} activity uses the designated \text{resource} at the given application \text{rate} during the designated time interval \((t_{\text{begin}}, t_{\text{end}})\), not to exceed the allocated \text{units} limit.

The \text{rate} defines the amount of \text{resource} consumed per \(dt\) of the \text{event \_process}. If \(dt\) is calendar time and the resource is human effort, the rate is staff level; if the resource is CPU and \(dt\) is in CPU hours, the rate is CPU hours per calendar day. When an \text{event \_process} expends its allocated \text{units}, the effective \text{rate} becomes zero.

Projects typically express schedule information in units of calendar time. If a project includes weekends, holidays, and vacations, then the schedule must either exclude these as inactive periods or else provide compensating rate factors between resource days and calendar time. For example, if a project is idle for two weeks during the winter holiday season, the schedule should not allocate any resources during this time. If a project works only 5 days a week, the resource utilization rate should allocate only \(\frac{5}{7}\)-ths of a workday per calendar day. (However, if the project allocates time and resources in weeks, then 5 days is a normal work week, and no rate adjustment is necessary in this case.)

### 3.3 Interdependencies Among Elements

Causal relationships exist between a project’s input reliability drivers and its resulting reliability profile, in that all development subprocesses consume resources and are driven, perhaps randomly, by other factors of influence.

Quantification of relationships is tantamount to modeling, and is required for simulation. As examples, the degree to which code and documents are inspected correlates with the number and seriousness of faults discovered in testing; the correctness of specifications relates to the correctness of ensuing code; and the seriousness of failures influences when a project will schedule the causal faults for repair.

Some relationships may be generic, while others may be unique to a given project. Some interrelationships may be subtle, while others may seem more axiomatic. Some of the typical axiomatic generic relationships among reliability profile parameters are:

1. All activities (including outages) consume resources.
2. Code written to missing, incorrect, or volatile specifications will be more faulty than code written to correct, complete, and stable specifications.
3. Tests rely on the existence of test cases. Old test cases rarely expose new faults.
4. The number of faults removed will be less than the number attempted. The attempted removal activity may also create new faults.
5. The number of validated fault removals will not exceed the number of attempts. Validation may erroneously report an fault removed.
6. Retesting usually encounters only failures due to bad fixes.

### 3.4 Software Environment Characteristics

Software in a test environment performs differently than it will in an operational environment. There are many reasons, more adequately addressed in Chapter 14, why this is the case. Principal reasons among them, however, are differences in configuration, execution purpose, execution scenarios, attitudes toward failures, and orientation of personnel. In brief, testing and operations are different environmentally.
Reliability profiles during test and operations depend on the environments themselves, not just on the test case execution scenarios and values of environment parameters. Although testing may attempt to emulate real operations in certain particulars, we must recognize that the characters of testing and operations are apt to differ significantly.

![Figure 16-1: Faults vs. execution time in test and operational facilities for a system of about 100,000 lines of assembly-language code.](image)

Figure ?? illustrates the effects that environment can have on the test profile. It depicts best-fit curves to actual cumulative failure data taken over three portions of a test activity. The software under test was part of the ground data system used in NASA’s deep-space missions.

The testing marked “Preliminary” was preparatory in nature, performed while the software was being installed in a “compatibility test area,” or test facility configured to match identically the operational environment to the extent economically feasible. The facility contained essentially everything electronic except the large-aperture antennas, low-noise maser receiver amplifiers, high-power transmitters, and spacecraft data system, that the later operational environment would have. Interfaces with missing operational subsystems were simulated.

Testing during the pre-transfer period proceeded until the failure rate leveled off at about 200 total faults (a predicted 18 faults remained). Then, activity transferred to deep-space tracking stations using similar scenarios, but in operational situations. Testing during this period found another 95 faults (considerably more than the 18 predicted). Despite the fact that the configurations and scenarios were essentially identical, there were about 306 – 218 = 88 faults that would never have been found had testing continued only in the test facility!

Predicting failures in differing environments requires, at the least, adaptability of the reliability model to fit configuration, scenario, and previous failure data. Such adaptations must accommodate for differences in hardware, test strategies, loading, database volume, and user training.

4 ARTIFACT-BASED SIMULATION

Software developers have long questioned the nature of relationships between software failures and program structure, programming error characteristics, and test strategies. Von Mayrhauser et al. [?, ?, ?] have performed experiments to investigate such questions, arguing that the extent to which reliability depends merely on these factors can be measured by generating random programs having the given characteristics,
and then observing their failure statistics. It is not important, in this respect, that the programs actually execute to perform useful functions, but merely that they possess the hypothesized properties that “real” programs would have in a given environment.

If the hypothesis is true, then the effects of the various controlled elements under study would be readily discovered. For example, by adjusting code structural characteristics (e.g., size, ratio of branching decisions to loop decisions, and fault distribution) in a controlled set of experiments, one may observe the contributory effects to failure behavior. One may also learn something about the sensitivities of reliability models to their founding assumptions. Such studies would lead practitioners to the best model(s) to use in given situations.

To explore the conjecture, they identified program properties and test strategies to be investigated. Then they performed experiments using automatically generated programs having the given properties, subjected these to the selected test strategies, and measured the reliability results.

Their investigations proceeded using only single-module programs (i.e., ones with no procedure calls), assumed that faults are only of a single type and severity, distributed uniformly throughout the program, and considered only a constant likelihood that a failure results when execution encounters a statement containing a fault.

There is no fundamental limitation in the artifact simulation technique that excludes procedure calls, multiple fault types, and time-dependent statistics. They were excluded in these early experiments to establish basic relationships. Their architecture will support multiple subprograms, faults of various types, severities, and distributions, and time-varying parameters at the later stages of experimentation.

4.1 Simulator Architecture

The reported simulation covers the coding, testing, and debugging portions of the software life cycle. The simulator consists of the following components (see Figure 16-2).

![Simulator Architecture Diagram](image)

Figure 16-2: Artifact-based software process simulator program.

The Code Generator uses program design and/or code structural and error characteristics to produce executable code with faults. Code generation is discussed more fully later (Section 16.2). The faults injected into the program cause actual execution failures during testing to occur in such a manner as to be detected by the Test Harness module, discussed below.

The Compiler is an ordinary compiler, the same as an actual project would use. The compiler generates executable code from the generated code and from updates following each of the fault repairs.

The Test Data Generator uses the generated code, together with parameters that select testing strategy, testing criteria, and phase (unit test, integration test, system test, etc.) to produce test input data and testing procedure parameters.

The Test Harness module applies test data to the simulated system in accordance with the selected test procedures, then detects each failure as it occurs, and categorizes it according to predetermined fault exposure and severity criteria.
The Debugger and Code Repair function of the simulation locates and repairs faults recognized by the test harness, and then reschedules the program for compilation and retesting. The debugger may fail to locate the fault and may either completely or incompletely remove the fault, when located. It may, at times, even introduce new faults. The debugger parameters include inputs to control locatability, severity, completeness, and fault detectability.

Reliability Analysis combines the failure data output by the test harness with the residual fault data from the debugger (undetected errors and incorrect repairs) to assess the reliability of the simulated code. This assessment compares failure results with the output of a conventional software reliability model.

4.1.1 Simulation Inputs

Artifact simulation experiments can vary many aspects of program construction and testing to investigate the effect of static properties on dynamic behavior. Inputs may include those which characterize code structure, coding errors, test input data, test conduct, failure characteristics debugging effectiveness, and computing environment.

The investigated code structure parameters pertained to control flow, data declaration, structural nesting, and number and size of subprograms. Statement type frequencies represented the structural dependencies of a program. The experiments assumed four types of program statements: assignments, looping statements, if statements, and subprogram calls. Data structure declaration characteristics were not simulated because faults in such structures tend to be caught by the compiler, and because the effects of faults in such statements would be included among those of the four chosen types.

Type, distribution, density, and fault-to-failure relationship parameters influenced the insertion of coding errors. The generator used type information to select the kinds of statements composing the program. Distribution information controlled where faults were located, either clustered in specified functions or scattered randomly throughout the program. Fault-to-failure relationships defined the frequency of failures when faults are encountered at run time.

Test input data depend on the testing environment, operational scenario, testing strategies, test phase, desired coverage, and resources available for testing. In the simulations reported, resource considerations were not addressed. Provisions for test strategies included features for random, directed, functional, mutation, sequencing, and feature testing. Test coverage selection included parameters designating node, branch, and data flow modes.

Projects classify failure attributes by type, severity, and detection status. Investigations so far have treated only failures of a single type and severity.

Debugging effectiveness depended on parameters associated with fault detection, identification, severity, and repair. Each of these, except severity, has correctness and resource dimensions. For example, identification establishes a fault-to-failure correspondence and the time required to make that correspondence.

Computing environment parameters included all the data required to run the test harness and analyze the failure data. The experiment environment data included machine, language, and workload parameters.

4.1.2 Simulated Code Generation

A code generator may produce simulated code using measured parameters of the actual project (trace-driven), or from generic data taken from a wide variety of project histories (self-driven).

Figure ?? illustrates a self-driven code generator architecture. The reported code simulator operates approximately as follows: Given the number of modules and the set of module sizes, the generator creates statements of the specified types according to their given occurrence frequencies, sometimes followed by code that represents a fault of the specified character.

The means for invoking multiple modules and for controlling the depth and length of nested structures were not explicitly revealed in the source references. Readers wishing to duplicate the simulator for their own experiments will have to decide upon appropriate models for these characteristics.

It is not necessarily assumed that real faults are uniformly distributed over the program; rather, the program generator can seed faults either on a function-by-function basis or statement-by-statement, according to a distribution by statement type and nesting level. Further, the fault exposure ratios of each fault need not be the same.
The presence of a real fault in a normal program corresponds to an ersatz element within the simulated program that can cause a failure contingent on a supposed failure characteristic. As the simulated program executes, a statement containing a fault will, with a given probability (the fault exposure ratio), cause a failure.

But the ersatz failure, when it occurs, is not meant to duplicate the appearance the real failure; only its occurrence and location are of importance. The injected fault thus needs to raise an exception with information that will identify where, when, and what type of failure has occurred. The reported simulator used a divide-by-zero expression to trigger the fault, which was then trapped by the execution harness via the \texttt{signal} capability of C.

An important step toward extending the utility of the simulation technique would be to make it trace-driven, or adaptive to project measurements as they emerge dynamically, rather than using only the static historic data of the self-driven simulation described above. Figure 16-3 illustrates how trace data would replace the random selection of statements during program execution.

4.1.3 The Execution Harness

The reported execution harness contains a test driver program that creates an interface between the generated program and its test data, spawns a “child” process to execute the generated program, and collects execution-time data on the child process separately from that of its ancillary functions, which may not have the same structural characteristics. The returned value of the child process indicated whether the test resulted in failure, or terminated naturally.

Simulated faults in the generated program could have been represented in any of a number of ways. The reported experiments used arithmetic overflow. Having detected the failure, the test harness enters the returned information into the failure log for use in locating and removing the fault. The updated program then recompiles and re-executes.

4.1.4 Reliability Assessment

The output log provides the execution time of each test run and indicates which runs experienced failures; it also identifies which fault caused the failure. Tools, such as those described in Appendix A, can use this data to generate tabular and graphical analyses of the failures. Such analyses may include application of any of a number of reliability growth models, maximum likelihood estimates and confidence limits for model parameters, and visual plots of important reliability attributes, such as cumulative failures or present failure intensity versus time.
The reliability assessment function also has available to it the output of the debugging function, which tells which faults were correctly, incorrectly, or incompletely made. At any time, then, the status of remaining faults in the generated program is visible.

4.2 Results

Reliability investigations using artifact simulation are currently in their formative stage. The fundamental, first-order validations of the equivalence hypotheses are yet in progress. Consequently, the process of evolution has imposed some limitations that will disappear, with time. The fundamental question has been: Do simulated programs in simulated environments exhibit reliability profiles representative of “real” programs in “real” environments that have the same parameters?

A software artifact simulation study, [?], compared (Figure ??) the results of testing a 5000 line C program with the predicted performance using the basic execution time model. These experiments demonstrated that the order in which failures occurred among statements containing faults closely matched the execution counts for those statements, and that the failure counts correlated with the types of program structures surrounding the faults.

These and other early results tend to confirm that static measures of program structure, error characteristics, and test strategies influence the reliability profiles of simulated and “real” programs in the same ways.

(Mike: TBD The reviewer asked “Can you relate this artifact-based simulation to work on FINE (at Univ of Illinois?) or other such fault-injection methods and tools?” Do you know of this work, and can you supply the information asked for?)

Artifact simulation studies of the future will continue to quantify the extent to which static parameters relate to reliability dynamics. As the software simulation art evolves, the effects of size, multiple-procedure program structures, multiple failure types, nonuniform fault distributions, and nonstationary parameters on reliability will increasingly become known.
5 RATE-BASED SIMULATION

The fundamental basis of rate-controlled event process simulation is the representation of a stochastic phenomenon of interest by a time series \( x(t) \) whose behavior depends only on a rate function, call it \( \beta(t) \), where \( \beta(t) \, dt \) acts as the conditional probability that a specified event occurs in the infinitesimal interval \( (t, t + dt) \).

A number of the analytic reliability growth models discussed in Chapter 4 echo this assumption and further assume that events in non-overlapping time intervals are independent. The processes modeled are thereby Markov processes \([?]\), or non-homogeneous Poisson processes (NHPP), which are also Markov processes. These include the models proposed by Jelinski and Moranda \([?]\), Goel and Okumoto \([?]\), Musa and Okumoto \([?]\), Duane \([?]\), Littlewood and Verrall \([?]\), and Yamada \([?]\). Rate functions for these appear in Section 16.??

The algorithms described here not only apply to simulating Markov processes, but are capable of simulating processes having time-dependent event count dependencies and irregular rate functions. These algorithms can simulate a much more general and realistic reliability process than has ever been hypothesized for any analytic model.

The mathematics presented in this Section treat general statistical event processes and rate-driven event processes, not merely those believed to describe software failures. As in the analytic models mentioned above, it is only the form of the rate functions and interpretation of parameters that set these models apart as pertaining to software. We begin this specialization formally in the next Section and continue it through Section 16.??.

First, however, we derive the forms of the simulation algorithms.

5.1 Event Process Statistics

If \( S_0 \) and \( S_1 \) denote the states of an event \( E \), \( S_0 \) in effect before the event and \( S_1 \) after its occurrence, then a particular member of the stochastic time series defined by \( \{ \beta(t), S_0, S_1 \} \) beginning at time \( t = 0 \) is a sample function, or realization, of the general rate-based discrete-event stochastic process. The zero subscript on \( \beta(t) \) signifies the \( S_0 \), or zero occurrences, starting state.

The statistical behavior of this process is well-known: The probability that event \( E \) will not have occurred prior to a given time \( t \) is given by the expression

\[
P(E_{t < \tau}) = \exp \left( - \int_0^\tau \beta(s) \, ds \right)
\]
\[ P_0(t) = e^{-\lambda_0(t, 0)} \]  

(16-1)

where

\[ \lambda(t, t_0) = \int_{t_0}^{t} \beta_0(\tau) \, d\tau \]  

(16-2)

The form of \( \beta_0(t) \) is unrestricted, but generally must satisfy

\[ \beta_0(t) \geq 0 \quad \text{and} \quad \lambda_0(\infty, 0) = \infty \]  

(16-3)

The first of these prevents the event from occurring at a negative rate, and the second stipulates that the event must eventually occur. If the second condition is violated, there will be a finite probability that the event will never occur.

When the events of interest are failures, \( \beta_0(t) \) is often referred to as the process hazard function and \( \lambda_0(t, 0) \) is the total hazard. The probability distribution function and probability density for the time of an occurrence are then

\[ F_1(t) = 1 - P_0(t) \]  

(16-4)

\[ f_1(t) = \beta_0(t) e^{-\lambda_0(t, 0)} \]  

(16-5)

The mean time of occurrence is

\[ E(t) = \int_{0}^{\infty} t \beta_0(t) e^{-\lambda_0(t, 0)} \, dt \]  

(16-6)

If \( \lambda_0(t, 0) \) is known in closed form, we may sometimes be able to write down and analyze the event probability and mean time of occurrence functions directly. In all but the simplest cases, however, we will require the assistance of a computer. When we cannot express the integrals in closed form, we can still evaluate them using straightforward numerical analysis.

5.2 Single-Event Process Simulation

It is rather easy and straightforward to simulate the rate-based single-discrete-event process, as illustrated in the following computer algorithm (expressed in the C programming language) which returns the occurrence time:

```c
double single_event(double t, double dt, double (*beta)(double))
{
    int event = 0;

    while (event == 0)
    {
        if (occurs(beta(t) * dt))
        {
            event++;
            t += dt;
        }
    }
    return t;
}
```

Above, the C language syntax defines a function named `single_event()` that will eventually return a double-precision floating-point value of the time of event occurrence. Starting at time \( t \), and continuing as long as the event value remains 0, the function monitors the event status; at the occurrence, event increases by 1, as signified by the ++ operation, which stops the iteration. Time augments by \( dt \) units each iteration, denoted by the “+=” operation.

We have programmed the `occurs(x)` operation as a macro that compares a random() value over \([0, 1)\) with the formal parameter \( x \), which must be less than unity, thus attaining the specified conditional probability function. (The `extern double` designation declares that `random()` is in an external library that returns a double precision floating-point value.) The interested reader may wish to consult [?] for a discussion of random number generation techniques.
extern double random(void);
#define occurs(x) (random() < x)

The particular application determines the form of the user-supplied rate function \( \beta(t) \). Any required initialization takes place in the `main()` program prior to invocation of the `single_event()` function. Figure ?? depicts the basic data flow of the overall program.

![Simulation program structure for a single event occurrence.](image)

We must choose the \( dt \) in simulations to satisfy the following conditions:

1. \( dt \) is smaller than the desired time-granularity of the reliability profile,
2. variation in \( \beta(t) \) over the incremental time intervals \( (t, t + dt) \) is negligible,
3. the chance of multiple event occurrences within a \( dt \) interval is negligible, and
4. the magnitude of \( \beta(t) \) \( dt \) is less than unity at each \( t \) in the interval of interest.

The time-complexity of the algorithm is \( O(\beta t/dt) \), where the \( \beta \) component represents the maximum complexity of computing \( \beta(t) \).

We may also simulate the behavior of a non-stochastic\(^3\) rate-based single-event process merely by altering the algorithm for `occurs()`. If \( \beta(t) \) represents the occurrence rate, then the event occurs when its integral reaches unity.

```c
double accumulated_rate;
#define occurs(x) (if ((accumulated_rate += x) < 1.) \n    then FALSE else TRUE)
```

The construction above increments `accumulated_rate` by \( x \) prior to checking its value; the expression then switches from a false to a true state when the value reaches unity. (The "\" at the end of the line signifies continuation on the next line of the macro.)

### 5.3 Recurrent Event Statistics

If we permitted the iteration in the previous algorithm to continue throughout a given time interval \( (0, t) \), then the simulated event could occur a random number of times, which could be counted. We may compute the probability distribution function \( F_n(t) \) that the \( n \) occurrence lies in the interval \( (0, t) \) as follows: If \( t_{n-1} \) has just been observed as the \( (n-1) \)st event occurrence, then we may treat the interval immediately after \( t_{n-1} \) as a new experiment. Translating Eq. (??) to the \( n \)th occurrence interval produces the occurrence distribution function conditioned on \( t_{n-1} \).

\(^3\)This technique also approximates the calculation of the mean occurrence behavior of a stochastic process; however, the method is exact only for the constant hazard case.
Simulation offers a relatively economical alternative in the evaluation of rate-based performance over the more complex numeric integrations of the previous Section. The recurrent events algorithm below is a simple extension of the single-occurrence event code that returns the number of occurrences over the time interval \((t_n, t)\). Its computational complexity to the \(n\)th occurrence is only \(O(\beta t_n/dt)\), in constant space:

\[
\begin{align*}
P_{n-1}(t | t_{n-1}) & = e^{-\lambda_{n-1}(t, t_{n-1})} \\
F_n(t | t_{n-1}) & = 1 - P_{n-1}(t | t_{n-1}) \\
\lambda_k(t, t_k) & = \int_{t_k}^{t} \beta_k(\tau) d\tau
\end{align*}
\]

The time dependency retained in Eq. (16-7) reflects the possible nonstationary nature of the event process. Each of the \(\beta_k(t)\) functions is subject to the restrictions given in condition (16-6); otherwise \(F_n(t | t_{n-1})\) above must be divided by \(1 - P_{n-1}(\infty | t_{n-1}) = 1 - e^{-\lambda(\infty, t_{n-1})}\). We shall assume these requirements in the remainder of this Chapter.

The \(n\)th occurrence probability densities then follow from differentiation of Eq. (16-7),

\[
\begin{align*}
f_n(t | t_{n-1}) & = \beta_{n-1}(t) e^{-\lambda_{n-1}(t, t_{n-1})} \\
f_n(t) & = \beta_{n-1}(t) \int_{0}^{t} e^{-\lambda_{n-1}(t, \tau)} f_{n-1}(\tau) d\tau
\end{align*}
\]

the latter being recursively defined, with \(t_0\) for the \(n = 1\) case defined as 0. The conditional probability displays the same type of statistical behavior seen in Eq. (16-7) for the single occurrence case above, but operates piecewise on successive intervals between occurrences.

Finally, \(F_n(t)\) follows by integration,

\[
F_n(t) = \int_{0}^{t} f_n(\tau) d\tau
\]

When events are modeled as Markov occurrences, the probability \(P_n(t)\) that exactly \(n\) occurrences appear in the interval \((0, t)\) is known [3] to be of the form

\[
\begin{align*}
P_0(t) & = e^{-\lambda_0(t, 0)} \\
P_n(t) & = \int_{0}^{t} \beta_{n-1} P_{n-1} e^{-\lambda_n(t, \tau)} d\tau
\end{align*}
\]

Mathematically closed-form solutions for these probability functions are rarely\(^4\) known. General solutions thus require simple, but perhaps time consuming, recursive numerical methods: The time-complexity of \(f_n(t | t_{n-1})\) is of order \(O(\beta(t - t_{n-1})/dt)\); \(f_n(t)\) and \(P_n(t)\) are of order \(O(\beta nt/\int dt)\), and \(F_n(t)\) is also of order \(O(\beta nt/\int dt)\).

The expected time of the \(n\)th occurrence follows directly from Eq. (16-7) as a recursive expression,

\[
\tau_n = \int_{0}^{\infty} tP_{n-1}(t) \int_{0}^{t} e^{-\lambda_{n-1}(t, \tau)} f_{n-1}(\tau) d\tau
\]

with time-complexity \(O(\beta nt_\infty/\int dt)\) and space complexity \(O(t_\infty/\int dt)\).

5.4 Recurrent Event Simulation

Simulation offers a relatively economical alternative in the evaluation of rate-based performance over the more complex numeric integrations of the previous Section. The recurrent events algorithm below is a simple extension of the single-occurrence event code that returns the number of occurrences over the time interval \((t_n, t)\). Its computational complexity to the \(n\)th occurrence is only \(O(\beta t_n/\int dt)\), in constant space:

\(^4\)Closed-form solutions for \(P_n(t)\) and \(f_n(t)\) are known to exist when the process is of the non-homogeneous Poisson variety, namely \(P_n(t) = \lambda^n(t, 0) \exp[-\lambda(t, 0)]/n!\) and \(f_n(t) = \beta(t) \lambda^n(0)/n!\).
```c
void recurrent_event(double ts, double t, double dt,
                     double (*beta)(int, double), int *events)
{
    while (ts < t)
    {
        if (occurs(beta(events, ts) * dt))
            *events += events;
        ts += dt;
    }
}
```

The calling program must initialize the `events` parameter to the actual number of occurrences prior to time $t_a$; `events` will contain the new count after the function returns. (Note that we renamed `events` in the plural to acknowledge that multiple occurrences are being counted.) Figure 16-7 depicts the program data flow structure.

![Program Data Flow Structure](image)

Figure 16-7: Simulation program for recurrent events.

Mathematically, $\beta_n(t)$ is valid only in the interval $t_n \leq t < t_{n+1}$ and signifies that $n$ occurrences of the event have occurred prior to $t$. The use of `beta(events, t)` in the algorithm acknowledges that the event may recur from time to time and that the occurrence rate function may not only change over time, but also may be sensitive to the number of event occurrences (as well as possibly other influences). The simulation algorithm observes the event occurrence times and may change the `beta()` function as required by the application.

We may also simulate non-stochastic rate-based recurrent-event processes by making the single-occurrence `occurs()` function recognize unit crossings of the rate accumulator, as follows:

```c
#define occurs(x) (if ((accumulated_rate +/- x) < 1.) \ 
               then FALSE else (accumulated_rate +/- 1., TRUE))
```

Note that `accumulated_rate` decrements by unity at each occurrence, signified by the “=+-” operation.

### 5.5 Secondary Event Simulation

Another type of event process of interest is when a primary event triggers the occurrence of a secondary event of a different type. For example, producing a unit of code may create a fault in the code.

Notationally, if $p_i$ denotes the probability that the $i$th occurrence of the primary event causes the occurrence of the secondary event, then we may express the probability $P_m^{(2)}(t)$ that $m$ such secondary events have occurred in the interval $(0, t)$ as

$$P_m^{(2)}(t) = \sum_{n=m}^{\infty} p_m p_n(t) P_n(t)$$  \hspace{1cm} (16-16)

---

5This technique, as before, approximates the average occurrence behavior of stochastic recurrent-event processes.
with

\[ p_{m|n} = \text{Prob}\{m \text{ secondary events}|n \text{ primary events in } (0, t)\} \]
\[ = \sum_{i \in I} p_i \ldots p_m (1 - p_{i+m}) \ldots (1 - p_n) \]

where the index vector \( i = (i_1, i_2, \ldots, i_n) \) is a permutation of \((1, 2, \ldots, n)\) such that \((i_1, \ldots, i_m)\) extends over all combinations of \( m \) out of \( n \) primary events, a set \( I \) of size \( \binom{n}{m} \). The computational complexity of \( P_{m|n}(t) \) is thus of combinatorial order, and not practical to evaluate in general cases of practical interest. In the special case that \( p_i = p \) is constant, \( p_{m|n} \) reduces to the binomial function,

\[ p_{m|n} = \binom{n}{m} p^m (1 - p)^{n-m} \]

The simulation algorithm for a dependent secondary event process, however, can remain quite general; one merely adds a mapping array secondary_event that relates the primary event to its secondary event and a function \( p(i, \text{ events}, t) \) that returns the probability that the primary event triggers the secondary event.

```c
if (occurs(beta(i, events, ta) * dt))
{  ++events[3];
  if ((j = secondary_event[i]) && occurs(p(j, events, t)))
  {  ++events[j];
  }
}
```

We may similarly treat multiple secondary events emanating from a single primary event, at only a moderate increase in algorithm complexity.

### 5.6 Limited Growth Simulation

When the final number \( N \) of occurrences that an event process may reach is prespecified, the normal growth of the event count over time must stop after the \( N \)th occurrence. For example, if there are \( N \) faults to repair and repairs proceed reasonably, then effort ceases after the last one is fixed.

Simulating this behavior is simple, but must include steps to prevent the event count from overshooting \( N \) when multiple occurrences occasionally take place within a \( dt \) interval. This may be done by altering the event counting functions not to exceed prespecified maxima `max_events` as follows:

```c
if (events[i] < max_events[i])
  ++events[i];
```

### 5.7 The General Simulation Algorithm

You may have already guessed the form of a general rate-based discrete-event process simulator. It is merely the recurrent-event algorithm augmented to accommodate multiple simultaneous events, multiple event categories, secondary events, and growth limits.

The general algorithm below incorporates all of these features. It simulates \( f \) event processes over a time interval \( ta \) to \( t \) using time slices of duration \( dt \); an initialized input array `events` counts the occurrences, which may not exceed corresponding values in the `max_events` array; an array `event_categories` contains the mapping of event occurrences into categories, which counts occurrences up to the maxima specified in the `max_categories` array; and a `secondary_event` array and `p()` function control secondary occurrences, as described in Section 16.22. For readability, the control function name `beta()` becomes `rate()`. We also add `action()` and `display()` functions, described below.

```c
void simulate(int f, double ta, double t, double dt,
  double (*rate)(int, int *, double),
  int events[], int max_events[],
  int categories[], int max_categories[],
  int event_category[],
  int secondary_event[], double (*p)(int, int *, double),
```

16
The `action()` function specifies what takes place when an event occurs. For example, if one category of events represents identified faults and another represents repairs, then `action()` may compute an unrepaird fault parameter for `display()`, or it may recompute appropriate `max_events` or `max_categories` bounds. The `action()` functions may well also pass additional parameters, such as `i`, `j`, `k`, and `m`, should these local values be needed to effect the proper change in state.

The `display()` function outputs the simulation status monitors as a profile in time. It may publish only certain parameters of interest, or it may detail the entire reliability state at each `dt`, depending on the time-line information desired by the user.

Figure ?? shows the overall simulation program data flow.

![Diagram of simulation program data flow](image)

**Figure 16-8: General rate-controlled process simulator program.**

We must choose the `dt` for the simulation experiments simultaneously to satisfy earlier-stated constraints imposed by each of the event rates. As a consequence, execution may be very slow. Alternatively, we could speed up the algorithm by choosing larger values of `dt` and computing the numbers of multiple events that may occur during each of the larger intervals, as determined by the probability functions of primary and
secondary events. It is known, when event occurrences in non-overlapping intervals are independent (see, e.g., [?]), that primary events are Poisson distributed and secondary events are binomially distributed. Generally, however, the probability functions are unknown, even when the rate functions are fairly simple.

But if these probability functions were known, there would only be slight changes required in the algorithm above: \( \text{occurs}(\text{rate}() \ast \text{dt}) \) would be replaced by a \( \text{primary} (\text{rate}() \ast \text{dt}) \) function that counts the random number \( n \) of primary event occurrences in the \( \text{dt} \) interval; \( \text{occurs}(\text{p}()) \) gets replaced by \( \text{secondary}(n, p()) \), which counts the number \( m \) of occurrences of secondary events; and \( \text{events[]} \) augments by \( n \) and \( m \), rather than unity, respectively.

If we desire, for execution-time reasons, a value of \( \text{dt} \) that is too large for use in the general algorithm above, but is yet small enough that primary and secondary event statistics over \( \text{dt} \) intervals are approximately Poisson and binomially distributed, respectively, then the modified algorithm can be applied. We refer to this configuration as the “piecewise-Poisson” approximate simulation. Piecewise-Poisson simulations, of course, are valid for the all the usual NHPP models, because no approximations are actually made. We have not yet studied the validity of the approximation applied to other processes.

6 RATE-BASED RELIABILITY

Rate-based reliability simulation is a natural extension of techniques for analyzing conventional models, because many of these are also are rate (or hazard) based. The underlying processes assumed by these models are thus the same.

Because of algorithmic simplicity, simulation serves as a powerful tool not only for analyzing the behaviors of processes assumed to have complex rate functions, but also for investigating whether the stochastic nature of a project’s measured failure data is typical of that obtained by simulation. One may vary the modeling assumptions until profiles reach a satisfactory alignment.

The challenge in life cycle simulation is finding rate functions that satisfactorily describe all of the activities, not just testing. Such a model enables optimum planning through trade-offs among allocated resources, test strategies, etc.

6.1 Rate Functions of Conventional Models

Several published analytic models treat (or approximate) the overall growth in reliability during the test and fault removal phases as non-homogeneous Poisson processes in execution time, while others focus on Markov execution-time interval statistics. While these may differ significantly in their assumptions about underlying failure mechanisms, they differ mathematically only in the forms of their rate functions. Some examples are the following:

1. The Jelinski-Moranda model [?] describes statistics of failure time intervals under the presumption that \( \beta_0(t) = \beta_0(1 - n/n_0) \), where \( n_0 \) is the estimated (unknown) number of initial software faults and \( \beta_0 \) is initial failure rate.

2. The Goel-Okumoto model [?] treats an overall reliability growth process with \( \beta(t) = n_0 \phi e^{-\beta t} \), where \( n_0 \) and \( \phi \) are input parameters, \( n_0 \phi \) being the initial failure rate, and \( \phi \) the rate decay factor. Strictly speaking, this rate function violates the conditions on \( \lambda(t, 0) \) imposed in (??), because \( \lambda_0(\infty, 0) = n_0 \) and \( P_0(\infty) = e^{-\rho_0} \). In practicality, \( n_0 \) is usually fairly large, so the consequences may be negligible.

3. The Musa-Okumoto model [?] posits an overall reliability growth process in which \( \beta(t) = \beta_0/(1 + \theta t) \), where \( \beta_0 \) is the initial failure rate and \( \theta \) is a rate decay factor. Both \( \beta_0 \) and \( \theta \) are input parameters.

4. The Duane model [?] deals with another overall reliability growth model, with \( \beta(t) = kbt^{k-1} \), where \( k \) and \( b \) are input parameters. Condition (??) requires that \( 0 < \beta < 1 \).

5. The Littlewood-Verrall inverse linear model [?] is an overall reliability growth model with \( \beta(t) = \beta_0/\sqrt{1 + \theta t} \) where \( \beta_0 \) is the initial failure rate and \( \theta \) is a rate decay factor.

6. The Yamada delayed S-shape model [?] represents still another overall reliability growth model, with \( \beta(t) = \phi \gamma t^{-1-\gamma} \), where \( \phi \) (the maximum failure rate) and \( \gamma \) are input parameters. This rate function,
too, violates condition (??), as \( \lambda_0(\infty, 0) = e^{\phi}/\gamma \) and \( P_0(\infty) = e^{-e^{\phi}/\gamma} \); again, when the large number of faults is large, the effect is negligible.

You may find discussions of these models elsewhere in Chapter 4.

### 6.2 Simulator Architecture

We have already discussed the algorithm for rate-based simulation. The remaining architectural considerations are characterized by input parameters, event rate functions, event-response actions, and output displays. The scope of user requirements should set the level of detail being simulated.

A reliability process simulator should be able to respond to schedules and work plans and to report the performance of subprocesses under the plan. By viewing simulated results, users may then replan as necessary. The simulator described here therefore does not assume specific relationships involving staff, resource, or schedules, but expects these as inputs, in the form described in Section 16.??.

Simulations should also embody interrelationships among project elements. For example, defective specifications should lead to faults in the code unless defects are corrected before coding takes place; missing specifications should introduce even more coding errors; testing should not take place without test cases to consume; repair activity should follow fault identification and isolation; and so on.

A more comprehensive simulation model ([??]) of the reliability process uses about 70 input parameters describing the software project and development environment, together with a project plan of arbitrary length containing activities, resources allocated, and application schedules. This simulator displays time-line profiles of almost 50 measures of project reliability status and the resources consumed, by activity. Its experimental use is described later in Section 16.??.

We shall illustrate the principles of reliability process simulation in a somewhat more simplified example—only 25 input parameters and a project resource schedule are required. You should not regard this example necessarily as a tool ready for industrial use, but as a framework and means for experimentation, learning, and extension.

In the example, we simulate only a single category of events for each reliability subprocess. Further, simulations produce only two types of failure events, namely, defects in specification documents and faults in code, all considered to be in the same severity category.

We also simplify the example reliability process not to include document and code reuse and integration, test preparations and the dependencies between testing and test-case availability, outages due to test failures, repair validation, and retesting.

Other, more detailed, simplifications appear in the discussions below.

#### 6.2.1 Environment Considerations

We know that characteristics of the programming, inspection, test, and operational environments can influence the rates at which activities take place. For simplicity, however, we have eliminated as many of these from the example simulator as seemed reasonable to our goals here. A more refined tool for general-purpose industrial use would, of course, probably include more definitive environmental inputs.

Events, of themselves, carry no intrinsic hazard values. The rates at which events occur depend on a number of environmental and other factors, including the nature of the events themselves. The model must treat event hazards differently in different situations.

Some faults may be easier to discover by inspection than by testing, while for others, the opposite may be true. The fault discovery rate in testing normally depends on such parameters as the CPU instruction execution rate, the language expansion factor, the failure-to-fault relationship, and the scheduled CPU hours per calendar day that are applied. During inspections, on the other hand, fault discovery depends on the discovery-to-fault relationship, the fault density, the inspection rate, and applied effort.

A fault is independent of its means of discovery. The model must therefore realize different hazard-per-fault rates in differing discovery environments, rather than merely assign a specific hazard rate to the fault itself.
6.2.2 Subprocess Representation

In the example simulator, each activity produces occurrences of one or more uncategorized event types, either primary or secondary. Table 16-1 lists the simulated primary events. Except for test failures, all are goal-oriented processes with limiting values, as shown. Test failures are limited by the current fault hazard function.

Table 16-1: Reliability process primary events and limits.

<table>
<thead>
<tr>
<th>Primary Event</th>
<th>Rate Control</th>
<th>Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>doc unit created</td>
<td>build workforce</td>
<td>doc size</td>
</tr>
<tr>
<td>doc unit inspected</td>
<td>document insp workforce</td>
<td>doc insp goal</td>
</tr>
<tr>
<td>doc defect treated</td>
<td>document corr workforce</td>
<td>defects recognized</td>
</tr>
<tr>
<td>code unit created</td>
<td>coding workforce</td>
<td>code size</td>
</tr>
<tr>
<td>code unit inspected</td>
<td>code insp workforce</td>
<td>code insp goal</td>
</tr>
<tr>
<td>test failure</td>
<td>size, faults, cpu, exposure</td>
<td>faults created</td>
</tr>
<tr>
<td>failure analyzed</td>
<td>analysis workforce</td>
<td>test failures</td>
</tr>
<tr>
<td>fault repair attempt</td>
<td>repair workforce</td>
<td>faults found</td>
</tr>
</tbody>
</table>

Table 16-2 defines the secondary events that occur with a primary event, controlled by an occurrence probability that may depend on a number of combined factors. For example, the number of defects or faults recognized during inspections will not only depend on the inspection efficiency (the fraction of defects recognized when inspected), but also on the density of defects in the material being inspected. All secondary event occurrences are naturally limited in number to the occurrences of their primary events; no other limits are imposed.

Table 16-2: Reliability process secondary events, correspondences, and controls.

<table>
<thead>
<tr>
<th>Secondary Event</th>
<th>Primary Event</th>
<th>Rate Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>defect created</td>
<td>doc unit created</td>
<td>defect density</td>
</tr>
<tr>
<td>defect recognized</td>
<td>doc unit inspected</td>
<td>latent defects, efficiency</td>
</tr>
<tr>
<td>defect corrected</td>
<td>defect treated</td>
<td>correction efficiency</td>
</tr>
<tr>
<td>fault created</td>
<td>code unit created</td>
<td>fault density, missing/faulty doc</td>
</tr>
<tr>
<td>fault recognized</td>
<td>code unit inspected</td>
<td>latent fault density, efficiency</td>
</tr>
<tr>
<td>fault corrected</td>
<td>fault treated</td>
<td>fault correction efficiency</td>
</tr>
<tr>
<td>fault identified</td>
<td>test failure</td>
<td>id efficiency, fault density</td>
</tr>
<tr>
<td>fault repaired</td>
<td>failure analyzed</td>
<td>repair efficiency</td>
</tr>
<tr>
<td></td>
<td>repair attempt</td>
<td></td>
</tr>
</tbody>
</table>

6.2.3 Document Construction

Document generation occurs presumably at a constant mean number of units per workday, not to exceed the document size goal, modeled as a Poisson random value whose mean is the given size. Defects occur at a constantaverage rate per produced unit.

The general simulation algorithm requires that the average number of defects committed per $dt$ interval be made less than unity by choice of $dt$. For example, if one chooses the documentation unit as the page, then there should be fewer than one defect in the number of pages produced in the time $dt$, on average. If one expects a greater defect rate, then a smaller $dt$ must be chosen. (Use of the piecewise-Poisson approximation model would relax this restriction to the $dt$ over which the approximation is valid).

Input parameters are...
6.2.4 Document Inspection

Document inspection is a goal-limited process similar to document construction. Inspections take place at constant average rates per workday, encountering defects in proportion to the defect density and applied inspection effort, but recognizing only a fraction of those defects encountered.

The number of inspected units at any time may not exceed the number of units so far created, nor the document inspection goal, a binomially distributed value determined from the document size goal and the input inspection fraction. The number of defects recognized cannot, of course, exceed the number created.

Because known defects may not yet have been removed at the time of an inspection discovery, we count the event as a new defect in proportion to the fraction of as-yet undiscovered defects.

The salient input parameters are

\[ \text{doc\_inspection\_fraction} \]
\[ \text{doc\_inspected\_per\_workday} \]
\[ \text{fraction\_defects\_recognized} \]

6.2.5 Document Correction

Staff resources, correction resource requirements per defect, and defect correction efficiency determine the rate at which defects get treated and thereby corrected. Attempted corrections may also inject new defects. Corrections decrease the defect count, while new defects increase it.

The number of defects treated at any time is less than the number so far recognized, plus the number of bad corrections (i.e., defects treated but not corrected). The number of corrections cannot exceed either the number of defects recognized or treated.

Generally, defect corrections could change the overall amount of required documentation; however, we have not modeled this effect here.

The input values needed are

\[ \text{defects\_treated\_per\_workday} \]
\[ \text{fraction\_defects\_fixed} \]

6.2.6 Code Construction

Code production follows the same general routine as document construction. However, the faults injected depend not only on normal human fallibility, but also on the amount of defective and missing specifications, all three of which could cause faults of different classifications. We assign each injected fault in the example simulator to the same category, however.

As with documentation, we must assure that the number of faults per \( dt \) interval not exceed unity over the duration of interest. For example, if one chooses the code unit as a line of code, then there should not be more than one fault injected into the number of lines of code produced in \( dt \) time units, on average.

The number of units created is limited by the code size goal, a Poisson-distributed value whose mean is the input size.

External inputs required are

\[ \text{code\_size} \]
\[ \text{code\_per\_workday} \]
\[ \text{faults\_per\_unit} \]
\[ \text{faults\_per\_defect} \]
\[ \text{faults\_per\_missing\_doc\_unit} \]

Fault injection is a secondary event to code unit construction; the number of faults per code unit has three sources: the faults per code unit produced from perfect specifications; faults per document defect times the average number of defects per unit code; and the faults per missing document unit times the number of missing documentation units per code unit.
6.2.7 Code Inspection

Code inspection mirrors the document inspection process. Inspections take place at constant average rates per workday, encountering faults in proportion to fault density and applied inspection effort, but recognizing only a fraction of those encountered.

The number of units inspected cannot exceed the number of code units created so far, nor the code inspection goal, a binomially distributed value determined from the code size goal and the input inspection fraction. The number of faults recognized, of course, cannot exceed the number injected.

External input values are

\[
\begin{align*}
\text{code\_inspection\_fraction} \\
\text{code\_inspected\_per\_workday} \\
\text{fraction\_faults\_recognized}
\end{align*}
\]

Since previously found faults may not yet have been removed at the time of an inspection discovery, we count a fault discovery as a new fault in proportion to the fraction of as-yet undiscovered faults.

6.2.8 Code Correction

Code correction simulation follows the document correction pattern, translated to code units. Fault correction attempts reduce the open fault count when successful and may increase it if unsuccessful.

The number of faults treated cannot exceed the number recognized, plus the number treated but not fixed. The number of faults corrected is limited to the number found by inspection and the number treated so far.

In general, code corrections may require document changes. The simulator consumes resources for such changes, but does not alter the documentation size and defect status.

The input parameters that apply to code correction are

\[
\begin{align*}
\text{faults\_treated\_per\_workday} \\
\text{fraction\_faults\_fixed}
\end{align*}
\]

6.2.9 Testing

In simulated test activities, failures occur in proportion to the test hazard per fault, the current fault density, and applied CPU resources. The resource schedule specifies the rates of CPU consumption, so the only additional input parameters needed is

\[
\text{test\_hazard\_per\_fault}
\]

The test hazard per fault parameter depends on the CPU execution rate, the compiler code expansion factor, and the fault exposure ratio.

The number of test failures is unlimited; as long as there are faults, failures will occur at the hazard rate.

6.2.10 Fault Identification

The simulator presumes that projects analyze failures at constant average rates per workday, not to exceed the number of failures observed. The number of faults identified must remain less than the number created, but unrecognized by inspection. The fraction of undiscovered faults and the probability of correct isolation regulate the fault identification process.

Pertinent inputs are

\[
\begin{align*}
\text{failures\_analyzed\_per\_workday} \\
\text{fraction\_failures\_isolated}
\end{align*}
\]
6.2.11 Fault Repair

Attempts to remove faults consume constant average resources per fault. Only a fraction of attempted repairs are actually successful; the rest will mistakenly be reported as repaired.

The number of attempted repairs cannot exceed the number of faults so far found (in inspections, as well as in testing), plus the number bad repair attempts (those that did not result in fault removal). The number of faults removed in this process must not exceed the number found, less those that were corrected after inspections, and also may not exceed the number of repair attempts.

Resources consumed by attempted code repairs include resources for changes in code and documents. The amounts and makeup of code and documentation do not change, however.

Input parameters are

- code_repairs_per_workday
- fraction_faults_repaired

6.3 Display of Results

Internally, a process simulator carries very detailed, fine-grained information on the activities and events under study, of types that are both visible and latent in real projects. In the spirit of simulation, the profiles viewed by humans should appear as if taken from reality.

However, a simulation user may well desire visibility into latent values, such as the numbers of unfound defects and faults, in order to make decisions on subsequent actions. When real project profiles match their corresponding simulation profiles, then the user probably expects that the latent behaviors will also agree. But one must not expect latent, model-internal behaviors to be accurate, because they can never be matched with reality.

To some extent, real profiles depend on how projects instrument and organize themselves for reliability measurement. They may record the status of documents and development code only at certain milestones. Other parameters, such as failures, may be logged automatically by the operating system, if detected, or by humans on a daily or weekly basis.

Visible project parameters include (1) the input facts (or assumptions) that define the environment and (2) the measured profiles, such as pages of documentation, lines of code, defects and faults found by inspections, failures, test faults identified, repairs, resources expended, and schedule time.

7 APPLICATIONS

7.1 Example 1: The Galileo Project

This Section describes simulating a real-world project, based on data and parameters taken from a subsystem of the Galileo project at the Jet Propulsion Laboratory ([?]).

Galileo is an outer planet spacecraft project that began at the start of fiscal year 1977, a mission that was originally entitled “Jupiter Orbiter and Probe,” or JOP. Unlike previous outer solar system missions, the Galileo orbiter was intended to remain in Jovian orbit for an extended interval of time. This would allow observations of variations in planetary and satellite features over time to augment the information obtained by single-observation opportunities afforded by previous fly-by missions. Galileo was launched in October of 1989, to reach the Jovian system in 1995.

There are two major on-board flight computers in the Galileo spacecraft: The Attitude and Articulation Control Subsystem (AACS), and the Command and Data System (CDS). A significant portion of each of these systems is embodied in software. This case study focuses on the CDS software reliability profile.

The CDS performs such critical functions as command and control of the spacecraft and the acquisition and transmission of flight data. CDS software selects among the many available telemetry rates and modes, and commands and controls all on-board experiments involving instruments.

The CDS flight software is characterized as a real-time embedded subsystem having high reliability requirements in a project where the mission design was redone several times. The software consists of about 17,000 lines of assembly language code, with about 1400 pages of documentation, produced over a period of

Redesigns were necessitated by launch delays due to congressional actions and the Challenger disaster.
approximately 300 calendar weeks. The project spent 1200 days (in 5-day workweeks) in pre-test activities and 420 days (in 7-day workweeks) in test preparation, tests, and rework, for a total of 1620 total days. The actual test period lasted only 280 of the 420 days; the project recorded the failure profile only during this 280-day subsystem software testing period.

Legend:

- DU Document Units
- DI Document Units Inspected

Figure 16-9: Simulated Galileo CDS documentation status profile.

Some of the CDS project parameters needed for simulation were calculated from project records; other values were estimated by project personnel; we chose the remaining values as probably typical of this project, but for which we had no immediately available data. We assigned believed-typical values, for example, to parameters relating to the injection and removal of defects and faults. Thus, even though only a few verifiable parameters were available outside the software testing phase, we nevertheless formed an entire plausible hypothetical model in order to illustrate an end-to-end reliability process.

For lack of better development life cycle information, we presumed that all CDS events occurred at uniform rates per event, that all activities took place without resource and schedule variations, and that testing required applied CPU resources according to the basic execution time model.

Observing experiments using the simulator described in [?] led to regressive adjustments of the estimated project rate input parameters. Each experiment profiled the status of documents, code, defects, and faults as random streams; the final parameter values resulted in event profiles typified by those shown in the Figures that follow. The Figures depict the results of a single experiment in simulating documentation, code, defect, and fault profiles of the CDS software, sampled at 2 week intervals.

Note in particular the profiles of documentation, code, injected defects, and injected faults (precisely those activities where no real project data was available to aid in regressive adjustments). The smoothness that appears in the rise of these curves is due to the regularity of the schedule, not randomness in performance. Performance deviations seem invisible not because they are small, but because they are relatively small, as a result of the law of large numbers.\(^7\)

Although we have no CDS data to refute this behavior, we doubt that the assumed constant resource levels reflect reality. A more realistic extension to the case study would have been to introduce irregular schedules, since we know that people rarely dedicate their time exclusively to one single activity at a time. If actual CDS schedule information had been available, we could have input this data into the simulation, whereupon the process statistics would probably have appeared more irregular.

\(^7\) The law of large numbers governs the rate at which the sample mean of an experiment converges to the distribution mean. For many processes, this is of order \(O(1/\sqrt{n})\).
Figure 16-10: Simulated Galileo CDS code status profile.

Figure ?? shows that the simulated documentation units (DU) goal (dotted line) was actually a little less (1380) than the 1400 pages predicted (in this particular experiment), and shows that the project did reach this goal. We set a 90% documentation inspection goal, which was reached (DI). There was an rms deviation of about 37 pages measured across many experiments.

Figure ?? shows that the experimental code units (CU) goal was actually (dotted line) slightly more (17093) than the 17000 LOC predicted (in this simulation). Again, the amount of code inspected (CI) attained the 90% inspection goal we had set. Experimental rms deviation was about 130 LOC.

Figure ?? displays the experimental defect behavior: injected document errors (E), detected defects (D), remaining defects (E-d), and remaining detected defects D-d. These profiles appear a little more irregular than do those of documentation and code production, but not much. The documentation appears to contain a sizable number of latent defects; even many of the detected defects appear to have been left uncorrected. Experimental rms deviation in the final defect counts were about 25.

Figure ?? shows the experimental fault activity. These profiles exhibit visible randomness, only partially masked by the law of large numbers. The rise in total faults (F) during the code construction activity appears almost linear, again a consequence of a constant effort schedule. We chose project parameters that prevented the creation of faults by imperfect corrections and repairs.

Correction of faults found in inspections began in week 150 and continued until week 240, removing 162 of them. The plateau in total repaired faults (R) and in total remaining faults (F-R) between weeks 240 and 260 occurred due to staff preparations for testing. At week 260, the test phase began, and failures (f) rose to 341 by week 300. Of these, 284 were repaired (r).

By the end of the 300 week simulated project, almost 10 faults per kiloline of code had been found in inspections and corrected, and another 20 faults per kiloline of code had been uncovered by testing and removed. The latent fault density was about 2 faults per kiloline of code.

Standard deviations computed after conducting many such experiments were about 22 in fault count (F), 30 in faults remaining (F-R), 32 in discovered faults (f), and 25 in remaining discovered faults (f-r).

If the simulation parameters were typical of the CDS project, and had the real CDS project been conducted a number of times, these same ranges of variations in the final status of the CDS artifacts would have been observed. The simulation did not replicate the CDS project behavior, but mirrored the behavior of a CDS-type project.

Although the final fault discovery count seems typical of a CDS-type project, the time profile of the simulation results shown in Figure ?? does not quite seem to match the character of the actual project data.
The failure rate seems to be high during early tests and too low during later tests. The actual test resource schedule was certainly not as simple as that used in the simulation.

On the basis of these experiments, it appears that realistic simulation of the general reliability process will require that detailed resource and schedule information will have to be provided to the model. It is important to remember that an actual project will probably not proceed as smoothly as its simulation. Consequently, projects will have to plan and measure their achievements as carefully for simulation as they will for the actual production.

### 7.1.1 Comparisons with Other Software Reliability Models

Figure ?? compares the actual CDS subsystem failure data with that obtained from a constant-test-hazard-per-fault simulation. We detailed the testing phase into five subactivities with constant staffing, but having irregular CPU and schedule allocations, as shown in Table ???. We obtained these schedule parameters using “eyeball regression” of the simulator output against the project data. The fit appears adequate to describe the underlying nature of the failure process (we did not expect an exact fit, since the failure process is considered random).

Table 16-3: Simulator schedule for CDS testing phase.

<table>
<thead>
<tr>
<th>activity</th>
<th>accumulated failures</th>
<th>begin week</th>
<th>end week</th>
<th>staffing</th>
<th>CPU rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 functional test</td>
<td>90</td>
<td>0</td>
<td>5</td>
<td>2.0</td>
<td>0.4</td>
</tr>
<tr>
<td>2 feature test</td>
<td>150</td>
<td>5</td>
<td>13</td>
<td>2.0</td>
<td>0.4</td>
</tr>
<tr>
<td>3 operation test 1</td>
<td>300</td>
<td>13</td>
<td>23</td>
<td>2.0</td>
<td>1.2</td>
</tr>
<tr>
<td>4 operation test 2</td>
<td>325</td>
<td>23</td>
<td>33</td>
<td>2.0</td>
<td>1.0</td>
</tr>
<tr>
<td>5 operation test 3</td>
<td>341</td>
<td>33</td>
<td>40</td>
<td>2.0</td>
<td>2.0</td>
</tr>
</tbody>
</table>

A comparison of failure profile simulation results with predictions of three other models, Jelinski-Moranda (JM), Musa-Okumoto (MO), and Littlewood-Verrall (LV), appears in Figure ???. For better amplification of model differences, the Figure displays failures per week, rather than cumulatively. The JM, MO, and LV
Figure 16-12: Simulated Galileo CDS fault status profile.

statistics are “one-week-ahead” predictions, in which all failure data up to a given week are used to predict the number of failures for the next week.

The Figure shows that the CDS simulation behavior is very similar to the noisy actual profile in variability and trends; the simulation could have been used for assessing the CDS reliability status and trends during the entire testing phase. The simulated profile could even have been calculated well prior to the start of testing from schedule and resource plans. The other models above could not adequately predict even one week ahead.
Figure 16-13: Simulated Galileo CDS testing fault density profile (newly discovered faults per 2-week interval), constant CPU resource.

Mike: TBD. You need to label the axes on this figure.

Figure 16-14: Cumulative failure data for Galileo CDS.
Figure 16-15: Comparison of actual, model-predicted, and simulated (variable CPU resource) Galileo CDS test fault densities.
8 SUMMARY

In many ways, the methods for software reliability assessment reported in this Chapter are very satisfying, and in other ways, most frustrating. The techniques provide quantitative measures of software quality that can be used by management to guide progress of a project. The frustration is that progress unveils the depth of our ignorance about system reliability and the dearth of experimental reliability data. We have not addressed the means for obtaining the best simulation structures nor the number of past examples needed to validate them. Nor have we addressed means for forecasting parameter values for a project from historical data. But ignorance, frustration, and the challenge they inspire are potent motivations for research.

The modeling assumptions required by the two simulation approaches addressed in this Chapter are certainly less restrictive, but perhaps more demanding than those underlying analytic models. Simulation solves software reliability prediction problems by producing programs and data conforming precisely to reliability process assumptions. If simulation profiles differ from actual performance, then the user can adjust the simulation model until an “acceptable” match with reality obtains. Simulation thus enables investigations of questions too difficult to be answered analytically.

Tools and environments supporting simulations may offer significant assistance and insight to researchers and practitioners in understanding the effects of various software reliability drivers, in evaluating sensitivities of behavior to various modeling assumptions, and in forecasting software project status profiles, such as time-lines of work products and the progress of testing, fault isolation, repair, validation, and retest efforts.

Attempts to reproduce reliability signatures of real-world projects using simulation have, so far, been encouraging. The results tend to coincide intuitively with how “real” programs behave, and strengthen the hope that such methods in the future will enable fuller investigations into the relationships between static measures and dynamic performance. Such relationships foretell reliability profiles of programs not yet written.
PROBLEMS

1. Sketch a C program for the self-driven simulated code generator of Section 16.??. What method did you use to represent the injected faults?

2. Use Eq. (??) to derive a closed-form expression for the conditional mean occurrence times \( \hat{t}_n \) when the rate function is independent of time, but depends on the number \( n \) of events (i.e., \( \beta_n(t) = \beta_n \)).

3. Write a program to calculate the mean by Eq. (??) and conditional mean occurrence profiles for the simple, rate-driven recurrent event process in Problem ??, above.

4. Develop a formula for calculating the variance \( \sigma_n^2 = E[(t_n - \hat{t}_n)^2] \) of a rate-driven process. Extend the program in Problem ?? to compute the standard deviation, \( \sigma_n \).

5. Use the program of Problem ?? to analyze the behavior of a Jelinski-Moranda rate function \( \beta_n = 1 - n/25 \). Run the program and compare the conditional mean profile with the true mean occurrence time. Is there a significant difference? Plot the mean profiles and \( \pm 1-\sigma_n \) event envelopes. Is the expected deviation from the mean significant?

6. Write a program to simulate the event process in Problem ??, above. Plot several simulated random event profiles. How significant are the differences among the profiles?

7. Rewrite the rate function of the program in Problem ?? and Problem ?? to analyze the performance of the Musa-Okumoto model. How significant are the differences between the output profiles? How do these results compare with those of Problem ?? and Problem ???

8. Gather environmental and project data (including the resource schedule) of a simple reliability process in your organization (such as software failures) and simulate it. Compare simulated and measured results. Was it possible to make the two appear as sample functions of the same random process? What changes in the simulator inputs were required?

9. Based on comparisons of simulated and actual project profiles, what conclusions can be made on the accuracy of simulation experiments?

10. Write the C code for the `rate(process, events, t)` function of the example simulator from the descriptions given in the discussions of reliability subprocess architectures. The formal parameters are `process`, the integer index of the event; `events`, a pointer to the integer array of event counts; and `t`, the time of the simulation. Indicate how the program consumes staff and computer resources as activities unfold.
REFERENCES


Answers to Selected Problems

We could structure the code simulation program as

```c
struct program {
    int number_functions;
    ...
}

struct function {
    char *name;
    int number_parameters;
    int size;
    int nesting_level;
    double fault_density;
    double fault_exposure;
    double type_assign;
    double type_loop;
    double type_if;
    double type_call;
}

void main(argc, argv)
char **argv;
{
    int i, j;
    program p;
    function f[MAX_FUNCTIONS];
    char statement[MAX_STATEMENT_LENGTH];
    double choice;

    initialize_parameters(argv[1], &p, f);

    for (i = 0; i < p.number_functions; i++)
    {
        generate_function_header(statement, &f[i]);
        for (j = 0; j < f[i].size; )
        {
            choice = random();
            if (choice < f[i].type_assign)
                generate_assignment(statement, &f[i]);
            else if (choice < f[i].type_loop)
                generate_loop(statement, &f[i]);
            else if (choice < f[i].type_if)
                generate_if(statement, &f[i]);
            else /* choice < f[i].type_call */
                generate_function(statement, &f[i]);
            j++;
        }
        output(statement);
    }
}
```

Each of the `generate_type()` functions would contain the logic to generate the proper type of statement, followed by code that seeds faults of the proper character, as

```c
if (random()) < f->fault_density)
    inject_fault(statement, f->fault_exposure);
output(statement);
```

One way of injecting the fault would be to use code such as

```c
if (random()) < fault_exposure_ratio)
    raise(SIGFPE);
```

with an appropriate entry in the test harness to catch the failure event,
The sequence of conditional mean occurrence times is

\[ \hat{t}_n = \sum_{i=0}^{n-1} \frac{1}{\beta_i} \]

If we set \( Q_n(t) = \int_0^t f_{n-1}(\tau)e^{-\lambda_{n-1}(t, \tau)} d\tau \), then Eq. (??) leads to the recursive form

\[ Q_n(t) = e^{-\lambda_{n-1}(t, \tau)}Q_n(t - \Delta t) + \int_{t-\Delta t}^t e^{-\lambda_{n-1}(t, \tau)} f_{n-1}(\tau) d\tau \]

which can be easily be integrated numerically using the trapezoidal rule, from which \( f_n(t) \) and \( \hat{T}_n \) follow, as illustrated in the following code:

```c
#include <stdio.h>
#include <stdlib.h>

#define MAX_CELLS 3000  /* Maximum number of dt intervals */
#define TMAX 300.        /* Maximum time value */
#define DT (TMAX / (int) MAX_CELLS)

#define double pn[2][MAX_CELLS];

double main(int argc, char **argv)
{
    double b, c, et, Qn, t, t_bar_n, t_hat_n;
    int i, n, n0, this, last;

    n0 = initialize(argc, argv);
    et = 1.0;
    t_hat_n = 0.0;
    for (n = 0; n < (int) n0; n++)
    {
        b = beta(n);
        c = exp(-b * DT);
        et = 1.0;
        this = n % 2;
        last = '!'this;
        t = t_bar_n = 0.0;
        for (i = 0; i < MAX_CELLS; ++i, t += DT)
        {
            if (n == 0)
            {
                pn[this][i] = b * et;
                et *= c;
            }
            else
            {
                if (i == 0)
                { Qn = 0.0;
                } else
                { Qn = c * Qn + 0.5 * DT * (pn[last][i] + c * pn[last][i - 1]);
                    pn[this][i] = b * Qn;
                }
                t_bar_n += t * pn[this][i];
            }
        }
    }
}
```
The expression for the variance $\sigma_n^2$ of the occurrence time is

$$\sigma_n^2 = \int_0^\infty t^2 \beta_n(t) \int_0^t e^{-\lambda_n(t, \tau)} f_n(\tau) d\tau - t_n^2$$

Thus, one may calculate the variance by making the following alterations to the program in Problem ??:

```c
    double b, e, et, Qn, q, sigma_n, t, t_bar_n, t_hat_n;
    ...
    t = t_bar_n = sigma_n = 0.;
    ...
    q = t * pn[this][i];
    t_bar_n += q;
    sigma_n += t * q;
    ...
    sigma_n = sqrt(sigma_n * DT - t_bar_n * t_bar_n);
    ...
    if (e = t_bar_n - sigma_n < 0.)
        e = 0.;
    printf("%d, \%1.1f, \%1.1f, \%1.1f, \%1.1f\n",
           n + 1, t_bar_n, t_hat_n, sigma_n, e, t_bar_n + sigma_n);
```
Figure 16-17: Comparison of sample functions of a Jelinski-Moranda process with $\beta(n) = 1 - n/25.$

```c
#define N 6
#define occurs(x) (random() < x)
#define random() ((double) rand() / (double) RAND_MAX)

void main(int argc, char **argv)
{
    double dt = 0.1, t = 0.0;
    int change, events[N], i;

    initialize(argc, argv);
    for (i = 0; i < N; i++)
        events[i] = 0;
    while (t < 100.0)
    {
        change = 0;
        for (i = 0; i < N; i++)
            if (occurs(beta(events[i]) * dt))
            {
                events[i]++;
                change++;
            }
        t += dt;
        if (change)
            {
                printf("%.2f", t);
                for (i = 0; i < N; i++)
                    printf(" %d", events[i]);
                printf("\n");
            }
    }
}

static double beta_0, n_0;

void initialize(int argc, char **argv)
{
    time_t t;

    if (argc < 3)
        {
            fprintf(stderr, "Usage: jm <n_0> <beta_0>\n");
            exit(1);
        }
}
```
double beta(int n)
{
    return (beta_0 * (1. - (double) n / n_0));
}

It is merely necessary to change initialization and $\beta_n(t)$ functions to the Musa-Okumoto form. Figure ?? and Figure ?? below depict the mean and typical sample functions from the process with $\beta(t) = 1/(1 + 0.08t)$. Note that deviations from the mean are much wider than in the Jelinski-Moranda process above.

Figure 16-18: Comparison of mean and $\pm 1-\sigma$ deviations in a Musa-Okumoto process with $\beta(t) = 1/(1 - 0.08t)$.

Comparisons of actual and simulated results will probably tell you which model best fits your data and the accuracy of fit. Based on calculations or observations of the standard deviations of simulated results, you will probably discover that a range of model parameters could describe the actual project. Monte Carlo runs of the simulator using the range of parameters will reveal what confidence you may attribute to the predictive power of models.

The rate() function governs the occurrence of primary events only. Secondary events are controlled by a separate function, p(). In the code segment below, the workforce() and CPU_resource() functions return the resource rates currently scheduled for the given process.

double rate(int process, int *events, double t)
{
    double d, pace, w;
    if ((w = workforce(phase[process])) <= 0.)
        return 0.;
Figure 16-19: Comparison of sample functions of a Musa-Okumoto process with $\beta(n) = 1/(1 + 0.08t)$.

```c
switch (process)
{
    case DOC_UNIT_CREATED:
        pace = doc_per_workday * w;
        break;
    case DOC_UNIT_INSPECTED:
        pace = doc_inspected_per_workday * w;
        break;
    case DEFECT_TREATED:
        pace = defects_treated_per_workday * w;
        break;
    case CODE_UNIT_CREATED:
        pace = code_per_workday * w;
        break;
    case CODE_UNIT_INSPECTED:
        pace = code_inspected_per_workday * w;
        break;
    case FAULT_TREATED:
        pace = faults_treated_per_workday * w;
        break;
    case TEST_FAILURE:
        if (d = (double) events[CODE_UNIT_CREATED])
            pace = test_hazard_per_fault
                * (REMAINING_FAULTS / d)
                * CPU_resource(phase[process]);
        else
            pace = 0.0;
        break;
    case FAULT_ANALYZED:
        pace = failures_analyzed_per_workday * w;
        break;
    case FAULT_REPAIR_TRY:
        pace = code_repairs_per_workday * w;
        break;
}
return pace;
```

One may simulate consumption of resources by making the `workforce()` and `CPU_resource()` functions subtract the amount of resource (rate times $\Delta t$) from the number of remaining units allocated in the schedule.
tuple regulating the given process.