

Distributed Computation of Passage Time Quantiles and Transient State Distributions in Large Semi-Markov Models

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Abstract

Semi-Markov processes (SMPs) are expressive tools for modelling concurrent systems; they are a generalisation of Markov processes that allow for arbitrarily distributed sojourn times. This paper presents an iterative technique for passage time and transient analysis of large structurally unrestricted semi-Markov processes. Our method is based on the calculation and subsequent numerical inversion of Laplace transforms and is amenable to a highly scalable distributed implementation. Results for a distributed voting system model with up to 1.1 million states are presented and compared against simulation.

1. Introduction

Traditional techniques for the analytical performance modelling of parallel and distributed systems are predominantly based on the steady-state analysis of Markov chains. This is restrictive for two main reasons. Firstly, the Markov property imposes the (often unrealistic) limitation that all time delays must be exponentially distributed. Secondly, steady-state measures are adequate to determine mean passage time values but not to determine passage (response) time quantiles. This is an especially serious problem since passage time quantiles are assuming increasing importance as key quality of service and performance metrics.

The aim of the present study is to investigate the use of semi-Markov processes for the purposes of system description and analytical passage time calculation. By using SMPs we can specify more realistic models with generally distributed delays while still maintaining some of the analytical tractability associated with Markovian models.

Our specific contribution is an iterative algorithm for large structurally unrestricted SMPs that generates passage time densities and quantiles, as well as transient state distributions. The algorithm is based on the calculation and subsequent numerical inversion of Laplace transforms. One of the biggest problems involved in working with semi-Markov processes is how to store the Laplace transform of state sojourn times in an effective way, such that accuracy is maintained but representation explosion does not occur. We address this issue with a constant-space representation of a general distribution function based on the evaluation demands of the numerical inversion algorithm employed.

We implement our technique in a scalable, distributed and checkpointed analysis pipeline and apply it to instances of a distributed voting model. The high-level model description is given in the form of a semi-Markov Stochastic Petri net – our own preliminary proposal for a non-Markovian Stochastic Petri net formalism – and is textually described in an extended semi-Markovian version of the high-level DNAmaca Markov chain specification language [7]. Our results are validated against a simulation derived from the same high-level model.

The rest of this paper is organised as follows. In Section 2, we briefly detail the background theory behind semi-Markov processes, and show how to derive first passage times and transient distributions. Our iterative passage time procedure is presented and formalised in Section 3. Section 4 describes the practical implementation issues in numerically inverting Laplace transforms as well as storing and manipulating general distributions. Section 5 briefly introduces the semi-Markov stochastic Petri net formalism and DNAmaca specification system. Passage time and transient results are produced for systems with up to $\sim 10^6$ states which are validated by simulations. Section 6 concludes and considers future work.

2. Definitions and Background Theory

2.1. Semi-Markov Processes

Consider a Markov renewal process $\{(X_n, T_n) : n \geq 0\}$ where T_n is the time of the n th transition ($T_0 = 0$) and $X_n \in \mathcal{S}$ is the state at (just after) the n th transition. Let the kernel of this process be:

$$R(n, i, j, t) = \mathbb{P}(X_{n+1} = j, T_{n+1} - T_n \leq t \mid X_n = i)$$

for $i, j \in \mathcal{S}$. The continuous time semi-Markov process (SMP), $\{Z(t), t \geq 0\}$, defined by the kernel R , is related to the Markov renewal process by:

$$Z(t) = X_{N(t)}$$

where $N(t) = \max\{n : T_n \leq t\}$, i.e. the number of state transitions that have taken place by time t . Thus $Z(t)$ represents the state of the system at time t . We consider time-homogeneous SMPs, in which $R(n, i, j, t)$ is independent of any previous state except the last. Thus R becomes independent of n :

$$\begin{aligned} R(i, j, t) &= \mathbb{P}(X_{n+1} = j, T_{n+1} - T_n \leq t \mid X_n = i) \\ &= p_{ij} H_{ij}(t) \end{aligned}$$

where $p_{ij} = \mathbb{P}(X_{n+1} = j \mid X_n = i)$ is the state transition probability between states i and j and $H_{ij}(t) = \mathbb{P}(T_{n+1} - T_n \leq t \mid X_{n+1} = j, X_n = i)$, is the sojourn time distribution in state i when the next state is j .

2.2. First passage times

Consider a finite, irreducible, continuous-time semi-Markov process with N states $\{1, 2, \dots, N\}$. Recalling that $Z(t)$ denotes the state of the SMP at time t ($t \geq 0$), the first passage time from a source state i at time t into a non-empty set of target states \vec{j} is:

$$P_{i\vec{j}}(t) = \inf\{u > 0 : Z(t+u) \in \vec{j} \mid Z(t) = i\}$$

For a stationary time-homogeneous SMP, $P_{i\vec{j}}(t)$ is independent of t and we have:

$$P_{i\vec{j}} = \inf\{u > 0 : Z(u) \in \vec{j} \mid Z(0) = i\} \quad (1)$$

$P_{i\vec{j}}$ is a random variable with an associated probability density function $f_{i\vec{j}}(t)$ such that the passage time quantile is defined as:

$$\mathbb{P}(t_1 < P_{i\vec{j}} < t_2) = \int_{t_1}^{t_2} f_{i\vec{j}}(t) dt$$

In general, the Laplace transform of $f_{i\vec{j}}$, $L_{i\vec{j}}(s)$, can be computed by solving a set of N linear equations:

$$L_{i\vec{j}}(s) = \sum_{k \notin \vec{j}} r_{ik}^*(s) L_{k\vec{j}}(s) + \sum_{k \in \vec{j}} r_{ik}^*(s) \text{ for } 1 \leq i \leq N \quad (2)$$

where $r_{ik}^*(s)$ is the Laplace-Stieltjes transform (LST) of $R(i, k, t)$ from Section 2.1 and is defined by:

$$r_{ik}^*(s) = \int_0^\infty e^{-st} dR(i, k, t)$$

Eq. (2) has a matrix-vector form, $A\tilde{x} = \tilde{b}$, where the elements of A are arbitrary complex functions; care needs to be taken when storing such functions for eventual numerical inversion (see Section 4). For example, when $\vec{j} = \{1\}$, Eq. (2) yields:

$$\begin{pmatrix} 1 & -r_{12}^*(s) & \cdots & -r_{1N}^*(s) \\ 0 & 1 - r_{22}^*(s) & \cdots & -r_{2N}^*(s) \\ 0 & -r_{32}^*(s) & \cdots & -r_{3N}^*(s) \\ \vdots & \vdots & \ddots & \vdots \\ 0 & -r_{N2}^*(s) & \cdots & 1 - r_{NN}^*(s) \end{pmatrix} \tilde{x} = \begin{pmatrix} r_{11}^*(s) \\ r_{21}^*(s) \\ r_{31}^*(s) \\ \vdots \\ r_{N1}^*(s) \end{pmatrix} \quad (3)$$

where $\tilde{x} = (L_{1\vec{j}}(s), L_{2\vec{j}}(s), \dots, L_{N\vec{j}}(s))^T$. When there are multiple source states, denoted by the vector \vec{i} , the Laplace transform of the passage time distribution at steady-state is:

$$L_{\vec{i}\vec{j}}(s) = \sum_{k \in \vec{i}} \alpha_k L_{k\vec{j}}(s) \quad (4)$$

where the weight α_k is the probability at equilibrium that the system is in state $k \in \vec{i}$ at the starting instant of the passage. If $\tilde{\pi}$ denotes the steady-state vector of the embedded discrete-time Markov chain (DTMC) with one-step transition probability matrix $P = [p_{ij}, 1 \leq i, j \leq N]$, then α_k is given by:

$$\alpha_k = \begin{cases} \pi_k / \sum_{j \in \vec{i}} \pi_j & \text{if } k \in \vec{i} \\ 0 & \text{otherwise} \end{cases} \quad (5)$$

The row vector with components α_k is denoted by $\tilde{\alpha}$.

2.2.1 Transient state distributions

Another useful modelling result is the transient state distribution, $T_{ij}(t)$, of a stochastic process:

$$T_{ij}(t) = \mathbb{P}(Z(t) = j \mid Z(0) = i)$$

From Pyke's seminal paper on SMPs [10], we have the following relationship between passage time densities and

transient state distributions, in Laplace form:

$$T_{ij}^*(s) = \begin{cases} \frac{1}{s} \frac{1-h_i^*(s)}{1-L_{ii}(s)} & \text{if } i = j \\ L_{ij}(s)T_{jj}^*(s) & \text{if } i \neq j \end{cases} \quad (6)$$

where $T_{ij}^*(s)$ is the Laplace transform of T_{ij} and $h_i^*(s) = \sum_j r_{ij}^*(s)$ is the LST of the sojourn-time distribution in state i . For multiple target states, this becomes:

$$T_{i\vec{j}}^*(s) = \sum_{k \in \vec{j}} T_{ik}^*(s) = \frac{1}{s} \left(\Lambda_i \delta_{i \in \vec{j}} + \sum_{k \in \vec{j}, k \neq i} \Lambda_k L_{ik}(s) \right) \quad (7)$$

where $\Lambda_n = (1 - h_n^*(s))/(1 - L_{nn}(s))$ and δ_B is 1 if condition B is true and 0 otherwise.

To construct $T_{i\vec{j}}^*(s)$, for a vector of target states \vec{j} we require $2|\vec{j}| - 1$ passage time quantities, $L_{ik}(s)$, which we can be computed from $|\vec{j}|$ matrix calculations of the form of Eq. (3).

As for passage times, for multiple source states, \vec{i} , we weight the transient distributions accordingly:

$$T_{\vec{i}\vec{j}}^*(s) = \sum_{k \in \vec{i}} \alpha_k T_{k\vec{j}}^*(s).$$

3. Iterative Passage time Analysis

In this section, we describe a passage time generation method that creates successively more accurate approximations to the SMP passage time quantity of Eq. (2).

The iterative passage time technique considers the r th transition passage time of the system, $L_{ij}^{(r)}(s)$. This is the conditional probability density of the system being in any of the specified target states after r state transitions. The unconditioned passage time density, $L_{i\vec{j}}(s)$, is then obtained in the limit as $r \rightarrow \infty$. We calculate $L_{i\vec{j}}^{(r)}(s)$ for a sufficiently high value of r to give an approximation to within a specified degree of accuracy.

This iterative method bears a loose resemblance to the well-known *uniformization* technique [9, 8, 5] which can be used to generate transient-state distributions and passage time densities for Markov chains. However, as we are working with semi-Markov systems, there can be no *uniformizing* of the general distributions in the SMP. The general distribution information has to be maintained as precisely as possible throughout the process. We achieve this by using the representation technique described in Section 4.

Once an $L_{i\vec{j}}(s)$ quantity has been created, it can be used to generate $L_{i\vec{j}}^{(r)}(s)$ passage times (c.f. Eq. (4)) or transient distributions (c.f. Eq. (7)).

3.1. Technical Description

Recall the semi-Markov process, $Z(t)$, of Section 2.2, where $N(t)$ is the number of state transitions that have taken place by time t . We define the r th transition first passage time to be:

$$P_{i\vec{j}}^{(r)} = \inf\{u > 0 : Z(u) \in \vec{j} \mid N(u) \leq r, Z(0) = i\} \quad (8)$$

which is the time taken to enter a state in \vec{j} for the first time having started in state i at time 0 and having undergone up to r state transitions. $P_{i\vec{j}}^{(r)}$ is a random variable with associated probability density function, $f_{i\vec{j}}^{(r)}(t)$, which has Laplace transform $L_{i\vec{j}}^{(r)}(s)$.

$L_{i\vec{j}}^{(r)}(s)$ is, in turn, the i th component of the vector

$$\tilde{L}_{\vec{j}}^{(r)}(s) = (L_{1\vec{j}}^{(r)}(s), L_{2\vec{j}}^{(r)}(s), \dots, L_{N\vec{j}}^{(r)}(s))$$

which may be computed as:

$$\tilde{L}_{\vec{j}}^{(r)}(s) = U(I + U' + U'^2 + \dots + U'^{(r-1)}) \tilde{e} \quad (9)$$

Here U is a transition matrix with elements $u_{pq} = r_{pq}^*(s)$. U' is a modified transition matrix with elements $u'_{pq} = \delta_{p \notin \vec{j}} u_{pq}$, where states in \vec{j} have been made absorbing. The column vector \tilde{e} has entries $\tilde{e}_k = \delta_{k \in \vec{j}}$.

We include the initial U term in Eq. (9), so as to generate cycle times for cases such as $L_{ii}^{(r)}(s)$ which would otherwise register as 0, if U' were used instead.

From Eqs. (1) and (8):

$$P_{i\vec{j}}^{(r)} = P_{i\vec{j}}^{(\infty)} \quad \text{and thus} \quad L_{i\vec{j}}^{(r)}(s) = L_{i\vec{j}}^{(\infty)}(s).$$

Now, $L_{i\vec{j}}^{(r)}(s)$ can be generalised to multiple source states \vec{i} using the normalised steady-state vector, $\tilde{\alpha}$, of Eq. (5):

$$\begin{aligned} L_{\vec{i}\vec{j}}^{(r)}(s) &= \tilde{\alpha} \tilde{L}_{\vec{j}}^{(r)}(s) \\ &= (\tilde{\alpha} U + \tilde{\alpha} U U' + \tilde{\alpha} U U'^2 + \dots \\ &\quad \dots + \tilde{\alpha} U U'^{(r-2)} + \tilde{\alpha} U U'^{(r-1)}) \tilde{e} \end{aligned} \quad (10)$$

Calculating the individual $\tilde{\alpha} U U'^m$ vectors of Eq. (10) is a sparse matrix-vector multiplication operation, which can be achieved iteratively by multiplying the previous vector, $\tilde{\alpha} U U'^{(m-1)}$, by the transition matrix U' . The worst-case

time complexity for the calculation of Eq. (10) is $O(N^2r)$ versus the $O(N^3)$ of typical matrix inversion techniques.

Convergence of the sum in Eq. (10) is said to have occurred at a particular r , if for a given s -point:

$$\begin{aligned} |\operatorname{Re}(L_{ij}^{(r+1)}(s) - L_{ij}^{(r)}(s))| &< \epsilon \text{ and} \\ |\operatorname{Im}(L_{ij}^{(r+1)}(s) - L_{ij}^{(r)}(s))| &< \epsilon \end{aligned} \quad (11)$$

where ϵ is chosen to be a suitably small value (e.g. 10^{-8}).

4 Laplace Transform Inversion

The key to practical analysis of semi-Markov processes lies in the efficient representation of their generally distributed functions. Without care the structural complexity of the SMP can be recreated within the representation of the distribution functions. This is especially true with the manipulations performed in the iterative passage time calculation of Section 3.

Many techniques have been used for representing arbitrary distributions – two of the most popular being *phase-type distributions* and *vector-of-moments* methods. These methods suffer from, respectively, exploding representation size under composition and containing insufficient information to produce accurate answers after large amounts of composition.

As all our distribution manipulations take place in Laplace-space, we link our distribution representation to the Laplace inversion technique that we ultimately use. Our implementation supports two Laplace transform inversion algorithms: the Euler technique [2] and the Laguerre method [1] with modifications summarised in [6].

Both algorithms work on the same general principle of sampling the transform function $L(s)$ at n points, s_1, s_2, \dots, s_n and generating values of $f(t)$ at m user-specified t -points t_1, t_2, \dots, t_m . In the Euler inversion case $n = km$, where k typically varies between 15 and 50, depending on the accuracy of the inversion required. In the modified Laguerre case, $n = 400$ and, crucially, is independent of m .

The choice of inversion algorithm depends on the characteristics of the density function $f(t)$. If the function is continuous, and has continuous derivatives (i.e. it is “smooth”) then the Laguerre method can be used. If, however, the density function or its derivatives contain discontinuities – for example if the system exclusively contains transitions with deterministic or uniform holding-time distributions – then the Euler method must be employed.

Whichever inversion algorithm is used, it is important to note that calculating $s_i, 1 \leq i \leq n$ and storing all the distribution transform functions, sampled at these points, will be sufficient to provide a complete inversion. Storing our distribution functions in this way has three main advantages. Firstly, the function has constant storage space, independent of the distribution-type. Secondly, each distribution has, therefore, the same constant storage even after composition with other distributions. Finally, the function has sufficient information about a distribution to determine the required passage time or transient density (and no more).

Our implementation employs a distributed master-slave architecture similar to that of the Markovian passage time calculation tool of [6]. The master processor computes in advance the values of s at which it will need to know the value of $L_{ij}^{(r)}(s)$ in order to perform the inversion. The s -values are then placed in a global work-queue to which the slave processors make requests. On making a request slave processors are assigned the next available s -value and use this to construct the matrices U and U' . The iterative algorithm is then applied to calculate the truncated sum of Eq. (10) for that s -value. The result is returned to the master and cached (both in memory and on disk so that all computation is checkpointed), and once all values have been computed and returned, the final Laplace inversion calculations are made by the master. The resulting t -points can then be plotted on a graph. As inter-slave communication is not required, the algorithm exhibits excellent scalability (see Section 5.3.3).

5. Distributed System Modelling

We demonstrate the SMP analysis techniques of the previous sections with a semi-Markov model of a distributed voting system. As there is a rich tradition of modelling distributed systems with stochastic Petri nets [4, 11], we propose and then make use of a semi-Markov extension of GSPNs to generate the model.

The model is specified in a semi-Markov stochastic Petri net (SM-SPN) formalism (outlined below) using an extension of the DNAmaca [7] Markov chain modelling language. From here, the semi-Markov state space is generated and we extract passage time densities, cumulative distribution functions and transient distributions.

5.1. Semi-Markov Stochastic Petri Nets

Semi-Markov stochastic Petri nets are extensions of GSPNs [3], which can handle arbitrary state-dependent

holding-time distributions and which generate an underlying semi-Markov process rather than a Markov process. Formally a SM-SPN consists of a 4-tuple, $(PN, \mathcal{P}, \mathcal{W}, \mathcal{D})$, where:

- $PN = (P, T, I^-, I^+, M_0)$ is the underlying Place-Transition net. P is the set of places, T , the set of transitions, $I^{+/-}$ are the forward and backward incidence functions describing the connections between places and transitions and M_0 is the initial marking.
- $\mathcal{P} : T \times \mathcal{M} \rightarrow Z^+$, denoted $p_t(m)$, is a state-dependent priority function for a transition.
- $\mathcal{W} : T \times \mathcal{M} \rightarrow \mathbb{R}^+$, denoted $w_t(m)$, is a marking-dependent weight function for a transition, to allow implementation of probabilistic choice.
- $\mathcal{D} : T \times \mathcal{M} \rightarrow (\mathbb{R}^+ \rightarrow [0, 1])$, denoted $d_t(m)$, is a marking-dependent cumulative distribution function for the firing-time of a transition.

In the above \mathcal{M} is the set of all reachable markings for a given net. Further, we define the following general net-enabling functions:

- $\mathcal{E}_N : \mathcal{M} \rightarrow P(T)$, a function that specifies net-enabled transitions from a given marking.
- $\mathcal{E}_P : \mathcal{M} \rightarrow P(T)$, a function that specifies priority-enabled transitions from a given marking.

The net-enabling function, \mathcal{E}_N , is defined in the usual way for standard Petri nets: if all preceding places have occupying tokens then a transition is net-enabled. Similarly, we define the more stringent priority-enabling function, \mathcal{E}_P . For a given marking, m , $\mathcal{E}_P(m)$ selects only those net-enabled transitions that have the highest priority, i.e. :

$$\mathcal{E}_P(m) = \{t \in \mathcal{E}_N(m) : p_t(m) = \max\{p_{t'}(m) : t' \in \mathcal{E}_N(m)\}\}$$

Now for a given priority-enabled transition, $t \in \mathcal{E}_P(m)$, there is a probability that it will actually fire after a delay sampled from its firing distribution, $d_t(m)$:

$$\mathbb{P}(t \in \mathcal{E}_P(m) \text{ fires}) = \frac{w_t(m)}{\sum_{t' \in \mathcal{E}_P(m)} w_{t'}(m)}$$

Note that the choice of which priority-enabled transition is fired in any given marking is made by a probabilistic selection based on transition weights, and is not a race condition based on finding the minimum of samples extracted from

firing time distributions. This mechanism enables the underlying reachability graph of the SM-SPN to be mapped directly onto a semi-Markov chain.

The marking-dependence of the weights and distributions does, in fact, allow us to translate SPNs and GSPNs into the SM-SPN paradigm in a straightforward manner, but that translation is not within the scope of this paper.

5.2. A Distributed Voting System

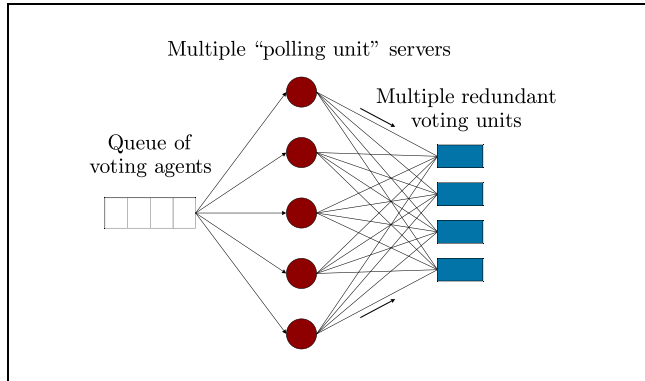


Fig. 1. A queuing model of a voting system

Fig. 1 shows the distributed components of a voting system with breakdowns and repairs, which we will use to generate a semi-Markov model. A voting agent queues to vote in the buffer; then, as a polling unit becomes free, it can receive the agent's vote and the agent can be marked as having voted. The polling unit contacts all the currently operational central voting units to register votes with all of them; this is done in order to prevent multiple vote fraud and to provide fault tolerance through redundancy. The polling unit then becomes available to receive another voting agent.

The semi-Markov stochastic Petri net for this system is shown in Fig. 2. Voting agents vote asynchronously, moving from place p_1 to p_2 as they do so. A restricted number of polling units which receive their votes transit t_1 from place p_3 to place p_4 . At t_2 , the vote is registered with as many central voting units as are currently operational in p_5 .

The system is considered to be in a failure mode if either all the polling units have failed and are in p_7 or all the central voting units have failed and are in p_6 . If either of these complete failures occur, then a high priority repair is performed, which resets the failed units to a fully operational state. If some but not all the polling or voting units fail, they attempt self-recovery. The system will continue to function as long as at least one polling unit and one voting unit remain operational.

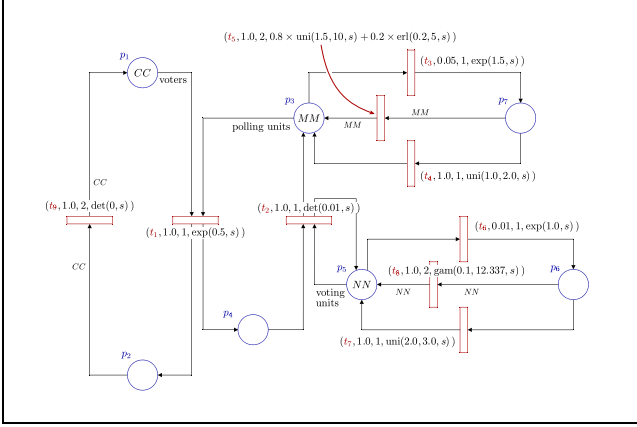


Fig. 2. A semi-Markov stochastic Petri net of the voting system

```

\transition{t5}{
  \condition{p7 > MM-1}
  \action{
    next->p3 = p3 + MM;
    next->p7 = p7 - MM;
  }
  \weight{1.0}
  \priority{2}
  \sojournTimeLT{
    return (0.8 * uniformLT(1.5,10, s)
      + 0.2 * erlangLT(0.001,5, s));
  }
}

```

Fig. 3. Excerpt from specification of voting example, showing definition of transition t_5 .

This example is defined in full as a DNAmaca specification [7], an excerpt of which is shown in Fig. 3. This defines transition t_5 , saying that it:

- is enabled when place p_7 has greater than $MM - 1$ tokens in it.
- removes MM tokens from place p_7 and adds MM tokens to place p_3 , when fired.
- has a weight 1.0 (used to define probabilistic choice between transitions when two or more are concurrently enabled).
- has a priority of 2, which will enable it above other transitions which would otherwise be structurally enabled but have a lower priority.
- is given a firing distribution which, with probability 0.8, is a uniform distribution or, with probability 0.2,

System	CC	MM	NN	States
0	18	6	3	2061
1	60	25	4	106,540
2	100	30	4	249,760
3	125	40	4	541,280
4	150	40	5	778,850
5	175	45	5	1,140,050

Tab. 1. Different configurations of the voting system as used to present results

is an Erlang distribution. The Laplace transform $g^*(s)$ for this firing time distribution is:

$$0.8 \times \text{uniformLT}(1.5, 10, s) + 0.2 \times \text{erlangLT}(0.001, 5, s)$$

where

$$\text{uniformLT}(a, b, s) = \frac{e^{-as} - e^{-bs}}{s(b-a)}$$

and

$$\text{erlangLT}(\lambda, n, s) = \left(\frac{\lambda}{\lambda + s} \right)^n.$$

In general, any arbitrary Laplace transform function can be specified as a firing distribution using the `\sojournTimeLT{...}` pragma.

5.3. Results

In this section, we compute passage time quantities for the time taken for a number of voters to pass from place p_1 to p_2 (a voter throughput quantity), as well as for the time taken for a fully operational system to enter a failure mode (i.e. when MM polling units fail in place p_7 or when NN central voting units fail in place p_6). We also extract simple reliability quantiles from cumulative distributions of the passage times, and transient measures for the voter throughput passage.

For the voting system described in Fig. 2, Table 1 shows how the size of the underlying SMP varies according to the configuration of the variables CC , MM , and NN , which are the number of voters, polling units and central voting units, respectively.

5.3.1 Example passage time distributions

Fig. 4 shows the density of the time taken for the passage of 175 voters from place p_1 to p_2 in system 5 as computed

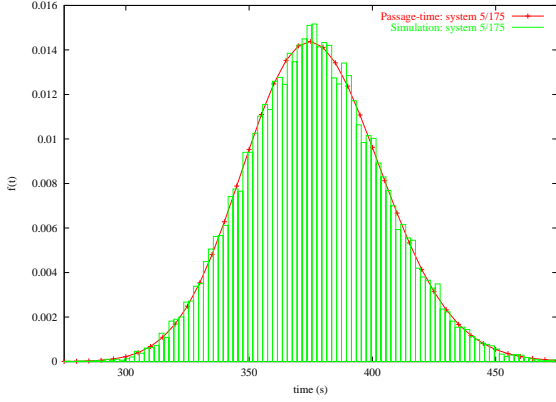


Fig. 4. Analytic and simulated density for the time taken to process 175 voters in system 5 (1.1 million states).

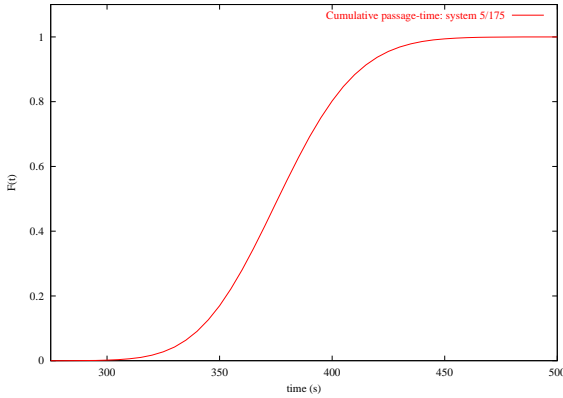


Fig. 5. Cumulative distribution function for the time taken to process 175 voters in system 5 (1.1 million states).

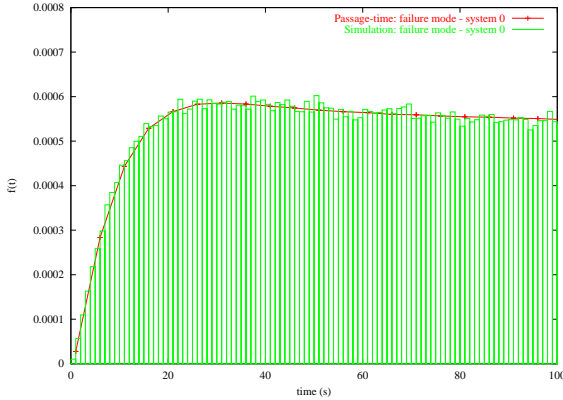


Fig. 6. Analytic and simulated density for failure mode passage in system 0 (2061 states).

by both our truncated iterative technique and by simulation. The close agreement provides mutual validation of the analytical method, with its numerical approximation, and the

simulation. It is interesting that, qualitatively, the density appears Normal. Certainly, the passage time random variable is a (weighted) sum of a large number of independent random variables, but these are, in general, not identically distributed.

Fig. 5 shows a cumulative distribution for the same passage as Fig. 4. This is easily obtained by inverting the Laplace transform $L_{\vec{t}_{ij}}(s)/s$; it allows us to extract response time quantiles, for instance:

$$\mathbb{P}(\text{system 5 processes 175 voters in under 440s}) = 0.9858$$

Fig. 6 shows analytic and simulated results for the time to complete failure in an initially fully operational voting system. It is produced for a much smaller system (2061 states) as the probabilities for the larger systems were so small that the simulator was not able to register any meaningful distribution for the quantity without using rare-event techniques. As we wanted to validate the passage time algorithm, we reduced the number of states so that the simulator would register a density. Examining very-low-probability events is an excellent example of where analytical techniques outperform simulations that would take many hours or even days to complete.

5.3.2 Example transient distribution

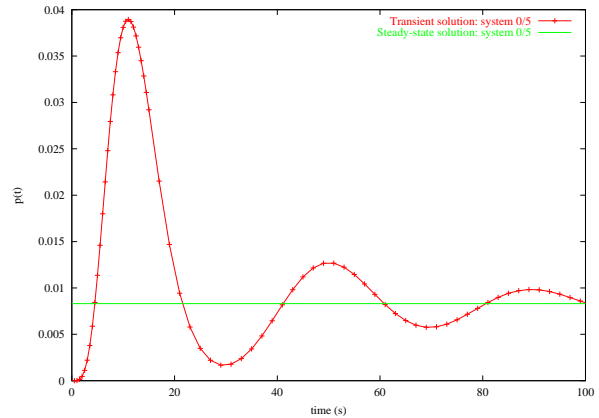


Fig. 7. Transient and steady-state values in system 0, for the transit of 5 voters from the initial marking to place p_2

We can use the transformation of Eq. (7) to generate transient distributions from passage time densities. Fig. 7 shows the transient distribution for the transit of five voters from place p_1 to p_2 . As expected, the transient distribution tends towards its steady-state value as $t \rightarrow \infty$.

Slave Processors	Time (s)	Speedup	Efficiency
1	549.08	1.00	1.000
8	71.11	7.72	0.965
16	39.16	14.02	0.876
32	24.10	22.79	0.712

Tab. 2. Time, speedup and efficiency for varying numbers of slave processors when calculating a passage time at 5 t -points for system 1, using Euler inversion (total of 165 s -point evaluations).

5.3.3 Tool scalability

Table 2 shows the time, speedups and efficiency for the analysis pipeline of Section 4 with varying numbers of slave processors when calculating 5 t -points for a passage time of system 1. The slave processors, each of which has a 2 GHz Intel Pentium 4 processor and 512 MB RAM, are part of a shared departmental network connected by 100MBps Ethernet. The master processor used was a dual 1 GHz Pentium III server with 2GB RAM (note, however, that a much lower spec machine would have been adequate as the master processor since it does not perform significant computation, nor does it require large amounts of memory). Even though exclusive access to the slave processors could not be guaranteed and the problem size in system 1 is relatively small, our distributed analysis pipeline still exhibits excellent scalability.

6. Conclusion

In this paper, we have derived passage time density, quantile and transient results for distributed systems with underlying semi-Markov state spaces of up to 10^6 states.

An iterative passage time generation algorithm was derived, implemented and compared against simulation. Our implementation optimises storage by relating the function to a set of s -points necessary for Laplace transform inversion. In this way, storage of an arbitrary distribution is kept constant and successive vector-matrix iterations do not suffer from the problem of representation-explosion.

Finally, we used a semi-Markov stochastic Petri net in conjunction with a semi-Markov extension to the DNAmaca language to specify a model of a distributed voting system, generate the corresponding semi-Markov state space and solve for a variety of passage time measures.

Our research efforts in the near future will include studying the convergence behaviour of our algorithm, with the goal of obtaining analytical bounds on the truncation error. In

addition, we will apply specialist techniques, e.g. using hypergraph partitioning of data structures, to achieve scalable algorithms for systems with up to $\sim 10^8$ states and beyond.

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