Fully Sequential Procedures for Large-Scale Ranking-and-Selection Problems in Parallel Computing Environments

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Fully sequential ranking-and-selection (R&S) procedures to find the best from a finite set of simulated alternatives are often designed to be implemented on a single processor. However, parallel computing environments, such as multi-core personal computers and many-core servers, are becoming ubiquitous and easily accessible for ordinary users. In this paper, we propose two types of fully sequential procedures that can be used in parallel computing environments. We call them vector-filling procedures and asymptotic parallel selection procedures, respectively. Extensive numerical experiments show that the proposed procedures can take advantage of multiple parallel processors and solve large-scale R&S problems.

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1. Introduction

Selecting the alternative with the largest or smallest mean performance from a finite number of alternatives is a common problem in many areas of operations research and management science. For instance, in designing a multistage manufacturing line, one may need to determine the best allocation of the buffer space to maximize the average throughput; in controlling an inventory system, one may need to identify the best reorder point to minimize the average cost; and in managing an ambulance service, one may need to select the optimal vehicle dispatching policy to minimize the average response time. In all of these examples the mean performances of the alternatives may be evaluated by running simulation experiments. This type of optimization problem is known as a ranking-and-selection (R&S) problem in the simulation literature.

Many R&S procedures have been developed (see, for instance, Kim and Nelson (2006b) for an introduction to the topic). These procedures typically allocate the simulation effort to all alternatives such that the best can be selected with certain statistical guarantees, e.g., a prespecified probability of correct selection (PCS). However, these procedures are often designed to handle a relatively small number of alternatives. As pointed out by Kim and Nelson (2006b), the two-stage procedure of Rinott (1978), hereinafter called Rinott’s procedure, is typically applied to fewer than 20 alternatives, and the fully sequential procedure of Kim and Nelson (2001), hereinafter called KN, is considered useful for fewer than 500 alternatives. The NSGS procedure of Nelson et al. (2001) is designed specifically to solve large-scale R&S problems. However, the largest test problem reported in their paper has only 500 alternatives.

In practice, however, there are many R&S problems that have thousands to tens of thousands of alternatives. Traditionally, these problems are solved using optimization-via-simulation (OvS) algorithms (see, for instance, Hong and Nelson 2009 for a recent review of OvS). Many of the OvS algorithms for this type of problem guarantee global convergence, i.e., they guarantee selecting the best alternative as the simulation effort goes to infinity. To achieve global convergence, however, these algorithms evaluate all alternatives as the simulation effort goes to infinity, and therefore become essentially R&S procedures. When they stop short...
of infinity, as they always do, there is often no statistical guarantee on the quality of the selected solution, and the solution may be significantly inferior to the optimal one. The goal of this paper is to provide R&S procedures, where the objective is to select the system with the largest mean response, that are valid and effective in parallel computing environments.

In the past few years there has been rapid adoption of parallel computing. Multiple-core processors are ubiquitous today; they are used not only in servers and personal computers but also in tablet computers and smart phones. Moreover, large quantities of computing (e.g., parallel processors) delivered as a service through the Internet, often called cloud computing, is becoming readily available and affordable to ordinary users. This motivates us to consider how to solve large-scale R&S problems in parallel computing environments. In particular, we are interested in whether current R&S procedures are statistically valid and efficient in parallel computing environments, and if they are not, how to design new procedures that are.

R&S problems can fit easily into parallel computing environments. If most of the computing time is used to generate independent simulation observations from various alternatives, then this can be done by executing the simulation programs in a parallel scheme without requiring any synchronization among different processors. This level of parallelization is called “embarrassingly parallel” (see, for instance, Foster 1995), and it makes parallel computing very attractive to solve R&S problems. This advantage of using parallel simulation technology for R&S problems has also been discussed by Chen (2005) and Yücesan et al. (2001).

The total computing effort required to solve an R&S problem typically increases only moderately as the problem size increases. Taking Rinott’s procedure (which samples from each alternative in two stages and only compares results after all sampling is completed) as an example, we plot the expected total number of samples as a function of the number of alternatives \( k \) in the solid line in Figure 1 (see Rinott 1978 for the procedure and Nelson et al. 2001 for a similar figure). To make this result more intuitive, suppose that we have 100 parallel processors and each processor can handle an R&S problem with 500 alternatives on its own as a single processor in the allowable amount of time using Rinott’s procedure. In the same amount of time, Rinott’s procedure on all processors can handle a similar problem with at least 30,000 alternatives, which significantly enlarges the size of R&S problems that may be solvable. In Figure 1, we also plot the maximum (or worst case) and the average expected total numbers of samples for the \( \mathcal{K} \mathcal{N} \) procedure (see Kim and Nelson 2001 for the details). Notice that when the number of alternatives increases, the proportion of clearly inferior alternatives often increases much faster than that of good alternatives. Therefore, fully sequential procedures, e.g., \( \mathcal{K} \mathcal{N} \), that allow early eliminations often require a much smaller expected sample size than the worst case, which makes fully sequential procedures more attractive for large-scale R&S problems than two-stage procedures such as Rinott’s.

From an implementation point of view, two-stage procedures are easier to parallelize than sequential procedures. For instance, a naive approach to implementing Rinott’s procedure is as follows: In the first stage, we distribute \( kn_0 \) replications equally among all processors and compute the first-stage sample variances and second-stage sample sizes of all alternatives after all processors finish their jobs. In the second stage, we again distribute all additional samples equally among all processors and select the alternative with best sample mean after all processors finish their jobs. Notice that the total time required to complete the procedure is determined by the processor that finishes its job last. When replication times of different alternatives are different or they are random, the total time of this approach may be quite long and cause many processors to be idle. To improve efficiency, one may estimate the replication time for each alternative after the first stage, formulate the sample allocation problem in the second stage as a stochastic parallel machine scheduling problem and minimize its makespan (i.e., total time to completion). Interested readers may refer to Pinedo (2008, Chapter 12) for more background on scheduling.

If the number of alternatives is less than or equal to the number of processors, then it makes sense to use multistage procedures, such as Rinott’s procedure, to reduce communication among processors. If the number of alternatives is much larger than the number of processors, however, it makes more sense to use fully sequential procedures with eliminations, such as \( \mathcal{K} \mathcal{N} \), to save simulation effort.

**Figure 1.** The expected total number of observations vs. the number of alternatives for both Rinott and \( \mathcal{K} \mathcal{N} \) when the initial sample size \( n_0 = 16 \), variances across all alternatives \( \sigma_i^2 = 1 \), and the indifference-zone parameter \( \delta = \sigma_i / \sqrt{n_0} \) in the slippage configuration where the difference between the means of the best and all other alternatives equals to \( \delta \).
by eliminating inferior alternatives early. Since large-scale R&S problems typically have a number of alternatives that is a few orders of magnitude larger than the number of available processors, in this paper we focus on designing fully sequential procedures to solve such problems.

There are many different configurations of parallel computing environments, ranging from multi-core personal computers to many-core servers to local computer farms to clouds on the Internet. We focus mainly on designing statistically valid fully sequential procedures for multi-core personal computers and many-core servers. Without communications via the Internet, the time for loading simulation programs to different processors and transmitting data among processors is almost negligible. When implementing these procedures in clouds on the Internet, however, there may be packet delays or even losses, which may affect the validity of the procedures. Therefore, we leave the design of fully sequential procedures for cloud implementations as a topic for future research.

When designing fully sequential procedures for a parallel computing environment, a critical question is what makes fully sequential procedures on multiple processors different from their counterparts on a single processor? A succinct answer to this question is that the input and output sequences of observations are different on multiple processors, whereas they are the same on a single processor. A single processor system works like a single-server queue, the departure (i.e., output) sequence is the same as the arrival (i.e., input) sequence. A multiple processor system works like a multiple-server queue; the departure sequence is in general different from the arrival sequence when the service time (i.e., replication time of an observation in our situation) is random. In a simulation study we may control the input sequence deterministically. For instance, in \( \mathbb{H}_N \) we simulate all alternatives one at a time according to a predetermined order. Therefore, the output sequence of a single processor system is also the same deterministic sequence. However, the same deterministic input sequence on a multiple-processor system may result in a random output sequence.

The randomness in the output sequence creates implementation issues as well as statistical issues when designing fully sequential R&S procedures. From an implementation point of view, randomness in the output sequence makes sample size synchronization difficult. For instance, when alternative 1 has 30 observations, alternative 2 may have 40 and alternative 3 may have only 20. Thus, procedures that require perfect synchronization of sample sizes from all alternatives are either difficult to implement or inefficient (i.e., using only a portion of the observations, such as setting the sample size to 20 in our three-alternative example). However, implementation issues may be easy to handle because there exist fully sequential procedures that allow unequal sample sizes from different alternatives (e.g., Hong 2006). The statistical issues caused by randomness in output sequence are more critical. First, when the performance of an alternative is correlated with its replication time, observations with shorter replication times tend to be available earlier and the sequence of output observations may not be independent even though they use independent random numbers. This problem also exists when simulating a single alternative using multiple processors; see Heidelberger (1988). Second, even when the performance of the alternatives is independent of their replication times (even when the replication times are constant), sample sizes of surviving alternatives depend on elimination decisions that in turn depend on sample-mean information of the alternatives. This type of dependence destroys the independence between sample means and sample sizes that are exploited in R&S procedures using a famous result from Stein (1945). More details on the statistical issues caused by random output sequences are discussed in §2.

In this paper we propose two solutions to deal with these issues. If one insists on making existing fully sequential R&S procedures suitable for parallel simulation schemes, implying that we may only perform comparisons based on the input sequence of samples, then we suggest creating a vector to record the observations exactly in the order of the input sequence and make comparisons based on a predetermined comparison rule. For instance, to implement \( \mathbb{H}_N \), one may perform a comparison after all surviving alternatives have their first \( r \) observations available for any \( r = n_0, n_0 + 1, \ldots \). Therefore, the procedure will have the same statistical validity as \( \mathbb{H}_N \). We call this type of procedure vector-filling (VF) as it fills the vector of observations based on the input sequence. Although the VF procedures have finite-sample statistical validity, they may not use all available observations at the time of comparison and may also add complexity in implementation as one needs to track the input order. Another issue with the VF procedures is that they may consume a large amount of memory to store the vector for R&S problems having a large number of alternatives. Even though the problem can be partly relieved by using a more effective memory management scheme, we may still encounter out-of-memory errors in some implementations. If one is content with asymptotic validity, we also design an asymptotic parallel selection (APS) procedure that allows unequal sample sizes for all alternatives and makes elimination decisions based on all available observations. The APS procedure can be shown to be asymptotically valid as the indifference-zone parameter goes to zero, an asymptotic regime also used by Kim and Nelson (2006b).

Our work is related to three streams of simulation literature. The first is the literature on R&S. In this paper we take a frequentist’s view and consider the indifference-zone (IZ) formulation of the problem. The IZ formulation was first proposed by Bechhofer (1954) and related procedures are summarized in Bechhofer et al. (1995) and Kim and Nelson (2006b). There are also many Bayesian formulations and procedures for R&S problems. For instance, Chen et al. (2000) and Chick and Inoue (2001a, b) allocate a
finite number of samples to different alternatives to maximize the posterior probability of correct selection. Instead of considering only the statistical measure of probability of correct selection, Chick and Gans (2009) and Chick and Frazier (2012) take sampling cost into account and formulate the R&S problems using dynamic programming techniques. A comprehensive comparison of the performance among different R&S procedures (designed under either a frequentist or a Bayesian formulation) has been conducted by Branke et al. (2007) in which they conclude that no R&S procedure can dominate in all situations.

The second stream of literature is on parallel and distributed simulation (PADS). According to Heidelberger (1988), PADS has two different approaches to parallelizing the simulation experiments: the first one is that each processor simulates multiple independent replications and the other one is that multiple processors cooperate on a single realization or replication. There is a vast literature on PADS from the 1980s and 1990s, where the focus was on the synchronization issues related to correct ordering of events in discrete-event simulations (see, for instance, Misra 1986 and Fujimoto 1990). Recently, cloud computing has also been applied to handle PADS (Fujimoto et al. 2010).

The third stream of literature is on simulation output analysis in a parallel and distributed simulation environment. Heidelberger (1988) discusses a variety of statistical properties for sample mean estimators calculated by observations from terminating simulations in a parallel simulation environment under three different stopping rules. Glynn and Heidelberger (1991) further study mean performance estimators for both terminating simulations and steady-state regenerative simulations under a completion-time constraint. Recently, Hsieh and Glynn (2009) proposed two new estimators for steady state simulations by weighting the sample average across replications on multiple processors according to some model selection criterion. To the best of our knowledge, there are only three papers using parallel and distributed simulation to solve R&S problems. The first is by Yücesan et al. (2001), who implement an optimal computing budget allocation (OCBA) algorithm in a web-based parallel environment to select the best alternative based on a Bayesian approach. The second is by Chen (2005), who applied a multistage R&S procedure with the simulation tasks of each stage distributed to multiple processors. Both papers test their procedures using only small-scale problems (both with only 10 alternatives), so it is not clear whether their procedures are capable of handling large-scale R&S problems. The third is by Ni et al. (2013), who proposed a “zipping” method to solve large-scale R&S problems in a high performance computing environment. The basic idea of their “zipping” method is to retrieve the independent and identically distributed (i.i.d.) sequence by controlling the seeds of the random number generator for each alternative, which is similar to the idea of our VF procedures.

In developing one of the pioneering works in handling R&S problems in parallel computing environments, we would like to highlight three main contributions of this paper. First, we demonstrate that large-scale R&S problems can be solved efficiently in parallel computing environments. Therefore, using parallel computing environments is a viable solution when there are a large number of alternatives. Second, we show that naive implementations of existing sequential R&S procedures in parallel computing environments may cause unexpected statistical issues that make these procedures inefficient or even invalid. To circumvent these issues, we propose the VF procedures that preserve the original statistical guarantees by carefully managing the output sequence of the simulation replications. Third, we propose the APS procedure that does not require active managing of the output sequence but is asymptotically valid. The APS procedure is simple to implement and the numerical study shows that it works well for the test problems.

The remainder of this paper is organized as follows: In §2, we use a queueing analogy to illustrate differences between using a single processor and using multiple processors to solve R&S problems. Based on the properties we identify in §2, we then propose two general approaches to designing R&S procedures, namely, the VF procedures and the APS procedures, in §§3 and 4. In §4, we also show the statistical validity of the APS procedure in a meaningful asymptotic regime. Numerical implementation of these two procedures as well as the numerical results are shown in §5, followed by some concluding remarks in §6.

2. The Randomness of Output Sequence

Suppose there are \( k \) alternatives whose mean performance can be evaluated by simulation on \( m \) processors. Let \( X_{ij} \) denote the \( l \)th observation from alternative \( i \), and we assume that \( X_{ij}, l = 1, 2, \ldots \), are i.i.d. random variables with a finite mean \( \mu_i \) for all \( i = 1, 2, \ldots, k \) and that \( X_{il} \) and \( X_{j\ell} \) are mutually independent for all \( l, d \), and \( i \neq j \). Under the IZ formulation, we further assume that \( \mu_1 - \delta \geq \mu_2 \geq \cdots \geq \mu_k \), where \( \delta \) is the IZ parameter; our goal is to design fully sequential procedures that are statistically valid, can select alternative 1 as the best with a probability at least \( 1 - \alpha \), and can be implemented on multi-core personal computers or many-core servers (with a total of \( m > 1 \) processors). For mathematical simplification, in this paper we assume that the \( m \) processors are identical in their processing speeds and that the time for loading simulation programs into processors and the time for transmitting data among the processors is negligible.

2.1. Queueing Analogy

To better understand the difference between implementation of fully sequential procedures on a single processor (i.e., \( m = 1 \)) and on multiple processors (i.e., \( m > 1 \)), we describe the simulation process using a queueing model.
analog. In this analogy, observations from alternative \( i \) are represented by class \( i \) customers, and \( m \) identical processors are represented by a server pool with \( m \) homogeneous servers. There is no arrival process in this queueing model; instead, all customers are waiting in the queue with a predetermined order at the beginning of the simulation process, and this predetermined order of customers is called the input sequence. When the simulation starts, the first \( m \) customers are assigned to the \( m \) servers. Once a server finishes the service of its current customer (i.e., generating the observation), the first customer waiting in queue will be immediately routed to that server. The departure process captures the order of customers who have finished service (i.e., the observations), and the order of departing customers is called the output sequence. When implementing fully sequential selection procedures, we perform comparisons and eliminations among the surviving alternatives based on the observations in the output sequence. When an alternative \( i \) is to be eliminated, the class \( i \) customers in the input sequence will abandon the queue and therefore will not be simulated.

For a fully sequential procedure implemented on a single processor, it is worthwhile noting that its input and output sequences are always the same. However, when the procedure is implemented on multiple processors, the output sequence may be different from the input sequence because the simulation times of different alternatives may be different, and the output sequence may even be nondeterministic because the simulation times of alternatives may be stochastic. See Figure 2 for an illustration. Because the output sequence may be different from the input sequence, we define \( Y_{ij} \) as the \( j \)th observation of alternative \( i \) in the output sequence, in addition to \( X_{ij} \), which represents the \( j \)th observation of alternative \( i \) in the input sequence. Notice that when simulation is conducted on a single processor, \( X_{ij} = Y_{ij} \). When simulation is conducted on multiple processors, however, it is possible that \( X_{ij} \neq Y_{ij} \).

Let \( \Gamma_{il} \) denote the (random) amount of time it takes to run \( X_{ij} \), the \( l \)th replication of alternative \( i \) in the input sequence. In the queueing analogy, \( \Gamma_{il} \) is the service time of the \( l \)th customer of class \( i \) in the queue. We assume that \( \Gamma_{il} > 0 \) almost surely (a.s.) and it has a finite mean \( \gamma_i > 0 \). Thus, \( \{(X_{ij}, \Gamma_{il}), l = 1, 2, \ldots\} \) is a sequence of i.i.d. bivariate random vectors. However, as the comparisons and elimination decisions for a sequential procedure are made based on the output sequence \( \{Y_{il}, l = 1, 2, \ldots\} \), it is critical to understand the statistical properties of \( \{Y_{il}, l = 1, 2, \ldots\} \). In the remainder of this section, we show that \( \{Y_{il}, l = 1, 2, \ldots\} \) may not be an i.i.d. sequence and it may compromise the statistical validity of existing fully sequential selection procedures.

2.2. Random Sample Sizes

When implementing a fully sequential selection procedure, one needs to specify the input sequence, which we call the sample allocation rule (SAR). SARs describe how observations from different alternatives are repeated in the input sequence. The most straightforward SAR is the round-robin rule that takes one observation from each surviving alternative in a predetermined order (say, alternatives 1, 2, \ldots, \( k \)); it is used in the \( \mathcal{AM} \) procedures of Kim and Nelson (2001, 2006a). For simplicity of presentation, we only consider the round-robin SAR in this paper.

Let \( t \geq 0 \) denote the run time from the start of the procedure and \( N_i(t) \) denote the number of completed observations of alternative \( i \) by time \( t \) for all \( i = 1, 2, \ldots, k \); then \( \{N_i(t), t \geq 0\} \) are continuous-time stochastic processes for all \( i = 1, 2, \ldots, k \) and \( \{N_i(t), t \geq 0\} \) are typically dependent. To better understand \( \{N_i(t), t \geq 0\} \) and its characteristics under single processor and multiple processors, we create a new phantom alternative and call it alternative \( p \). In the input sequence, alternative \( p \) is queued at the end of each round-robin cycle. For instance, when the alternatives are simulated in the order of alternative 1 to \( k \), alternative \( p \) is queued right after every alternative \( k \). Furthermore, alternative \( p \) has a simulation time of 0 and its observations are not compared to other alternatives.

**Figure 2.** An illustration using queueing models.
Therefore, it is clear that alternative \( p \) does not affect the implementation of the procedure, and this is why it is called a phantom alternative.

Let \( N_r(t) \) denote the number of observations of the phantom alternative in the output sequence by time \( t \). Let \( t_r = \inf\{t \geq 0: N_r(t) = r\} \) for all \( r = 1, 2, \ldots \), which is the time that the \( r \)th observation from the phantom alternative is obtained, and let \( N_r = N_r(t_r) \) be the number of completed observations of alternative \( i \) at time \( t_r \) for all \( i = 1, 2, \ldots, k \) and \( r = 1, 2, \ldots \). Next, we convert the continuous-time process \( \{N_r(t), t \geq 0\} \) into a discrete-time process \( \{N_r, r = 1, 2, \ldots\} \) for all \( r = 1, 2, \ldots \) and for any surviving alternative \( i \). Therefore, the discrete-time processes \( \{N_r, r = 1, 2, \ldots\} \) are deterministic for all surviving alternatives, which makes the statistical validity of \( \mathcal{R} \mathcal{N} \) easier to analyze.

When there are multiple processors, however, the discrete-time process \( \{N_r, r = 1, 2, \ldots\} \) becomes a stochastic process and \( N_r \) is typically different from \( N_r' \) when both alternatives \( i \) and \( j \) are surviving; i.e., \( N_r/N_{pr} \neq N_r'/N_{pr}' \). To illustrate this, we simulate four alternatives with eight processors using a round-robin rule, where the time to generate an observation of alternatives \( i \) follows an exponential distribution with mean \( i \) units of time, and we plot \( N_r/N_{pr} \) for all \( i = 1, 2, 3, 4 \) and \( r = 1, 2, \ldots, 100 \) in Figure 3. From the figure, we see clearly that \( N_r/N_{pr} \) are random and typically different from 1, but they appear to converge to 1 as \( r \) increases.

### 2.3. Loss of I.I.D. Property

For each alternative \( i \), even though \( \{X_{il}, l = 1, 2, \ldots\} \) is an input sequence of i.i.d. random variables, the output sequence \( \{Y_{il}, l = 1, 2, \ldots\} \) may no longer be i.i.d. observations of alternative \( i \) at time \( t_r \) for all \( i = 1, 2, \ldots, k \) and \( r = 1, 2, \ldots \). Let \( \mu_i \) denote the mean of \( Y_{il} \) for all surviving alternatives, which makes the statistical implementation of the procedure, and this is why it is called \( \mathcal{R} \mathcal{N} \) easier to analyze.

We use a simple example from Heidelberger (1988) as an illustration. Suppose there is only one alternative, alternative \( 1 \), to be simulated and \( X_1 = \Gamma_1 \) follows an exponential distribution with mean \( \mu_1 \). Then Heidelberger (1988) shows that the first observation from the output sequence, \( Y_{1l} \), is the shortest of \( m \) i.i.d. exponential random variables, that is, \( Y_{1l} = \min\{X_{11}, X_{12}, \ldots, X_{1m}\} \), which is exponentially distributed with mean \( \mu = \mu_1/m \). In the e-companion (available as supplemental material at http://dx.doi.org/10.1287/opre.2015.1413, EC.1.1), we derive closed-form expressions for \( Y_{il}, l = 1, 2, \ldots \). Based on these closed-form expressions, we can easily verify that \( Y_{il}, l = 1, 2, \ldots \), are not i.i.d. The mean of \( Y_{il} \) is

\[
\mathbb{E}[Y_{il}] = \mu_i \left[ 1 - \left( 1 - \frac{1}{m} \right)^n \right],
\]

and the expectation of the sample mean estimator is

\[
\mathbb{E}[\bar{Y}(n)] = \mu_i \left[ 1 - \frac{m-1}{n} \left( 1 - \left( 1 - \frac{1}{m} \right)^n \right) \right],
\]

which means \( \bar{Y}(n) \) has a downward bias. However, the bias goes to zero as the sample size \( n \to \infty \). Furthermore, the moment generating function (MGF) of \( Y_{il} \) is

\[
M_{Y_{il}}(t) = \frac{1}{1 - m \mu t} - \frac{m \mu t}{1 - m \mu t} \left( \frac{m-1}{m} \cdot \frac{1}{1 - \mu t} \right)^l,
\]

where \( t \) is in a sufficiently small neighborhood of 0. Notice that \( \lim_{n \to \infty} M_{Y_{il}}(t) = 1/(1 - m \mu t) \), which is exactly the MGF of an exponential random variable with mean \( m \mu = \mu_i \) (see the e-companion EC.1.1 for detailed derivations).

Furthermore, let \( M_{Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il}}(t) \) denote the MGFs of \( Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} \), respectively, for any \( l = 1, 2, \ldots \) and \( n = 1, 2, \ldots \). In EC.1.1., we also show that

\[
\lim_{n \to \infty} \left( M_{Y_{il}}(t) \cdot M_{Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il}}(t) - M_{Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il}}(t) \right) = 0.
\]

Therefore, the dependence between \( Y_{il} \) and \( Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} + Y_{il} \) also vanishes as \( n \to \infty \). In this sense, as \( n \to \infty \), the output sequence may be viewed as i.i.d. and statistically equivalent to the input sequence.

When there are multiple alternatives, however, the system dynamics are much more complicated, and we are not able to derive closed-form expressions for the distribution of \( Y_{il} \). Nevertheless, it is still quite clear that \( Y_{il}, l = 1, 2, \ldots \) are no longer i.i.d. observations and that \( \{Y_{il}, l = 1, 2, \ldots\} \) and \( \{Y_{il}, l = 1, 2, \ldots\} \) are likely dependent on each other. As an example, we simulate four alternatives on eight processors where \( X_{il} = \Gamma_{il} \) follows an exponential distribution with mean \( i \) time units. In Figure 4, we plot \( \mathbb{E}[\bar{Y}(n)] \), \( i = 1, 2, 3, 4 \), with the sample size \( n \) varying from 1 to 100, estimated from 1,000 macroreplications.
2.4. Dependence Caused by Eliminations

If \( X_i \) is independent of \( \Gamma_i \) (or even if \( \Gamma_i \) is constant) for all \( i = 1, 2, \ldots, k \), \( \bar{Y}(n) \) becomes an unbiased estimator of \( \mu_i \). However, the elimination decisions inherent to sequential selection procedures may still introduce dependence among the sample sizes of surviving alternatives and thus introduce dependence among their sample means. To illustrate this type of dependence, suppose there are three alternatives to be simulated on two processors and the replication times of alternatives 1, 2, 3 are fixed as 2, 1, 1 time units, respectively. Furthermore, suppose that the input sequence is in a round-robin order of 1, 2, and 3. Then the simulation process can be described as in Figure 5. At each time point \( t_i \) (which corresponds to the completion time of the \( r \)th phantom alternative), we conduct comparisons among all surviving alternatives. Notice that when all three alternatives are surviving, they all have an equal sample size \( N_i(t_r) = N_2(t_r) = N_3(t_r) = r \) at \( t_r \) for all \( r = 1, 2, \ldots \). Suppose at some time point \( t_n \), alternative 2 is eliminated; then at the next time point, \( t_{n+1} \), \( N_1(t_{n+1}) = n \) but \( N_1(t_{n+1}) = n + 1 \). Therefore, sample sizes of surviving alternatives depend on elimination decisions, which depend on sample means of all alternatives. This type of dependence may cause the sample means of the surviving alternatives to be dependent on each other.

When there is a large number of alternatives with random replication times simulated on many processors, the dynamics of elimination decisions can be more complicated, leading to more complicated dependence among the sample sizes and sample means of surviving alternatives. It is worthwhile noting that this problem is caused by the use of multiple processors. When a fully sequential procedure is implemented on a single processor, eliminations do not cause dependence because the output sequence of surviving alternatives remains the same as that without elimination.

In §§2.2–2.4, we have shown that the use of multiple processors may create various statistical issues when the observations in the output sequence are used to implement sequential selection procedures. To solve the problem, we take two different approaches and discuss them in the next two sections. In the first approach, we implement sequential procedures using the observations in the output sequence based on their order in the input sequence. Therefore, the finite-time statistical validity of these procedures may be guaranteed. However, this approach requires significant accounting and often a large amount of memory for storing the observations, and it may use only a portion of the observations in the output sequence (thus may not be efficient). Therefore, we propose another approach that designs sequential selection procedures that are asymptotically valid. This is possible because, as shown in §§2.2–2.4, the statistical properties of the sample-mean estimators \( \bar{Y}_i(n) \) tend to behave nicely as the sample sizes go to infinity.

3. Vector Filling Procedures

As mentioned above, if we restrict our attention to the use of observations exactly according to the predetermined order in the input sequence, then all existing fully sequential selection procedures are statistically valid when implemented in a parallel computing environment. To achieve this goal, we may create a vector to record the observations in the same order of the input sequence and place phantom alternative \( p \) in the positions where elimination decisions are scheduled. Then we can conduct comparison and elimination decisions when all observations from surviving alternatives ahead of every position for the phantom alternative have been collected in the vector. We call this type of procedures a vector-filling procedure. In this section, we provide a simple vector-filling procedure that extends the well-known \( \mathfrak{K}N \) to a parallel computing environment.

To simplify the presentation, we suppose there are \( m+1 \) parallel processors (or threads), which we call processors
0, 1, . . . , m. Processor 0 is used to manage the input sequence and to conduct comparisons and eliminations, while processors 1, 2, . . . , m are used to simulate the alternatives.

**Procedure 1 (Vector-Filling KN Procedure).**

Step 0. Setup. Select confidence level 1/k < 1 − α < 1, IZ parameter δ > 0, and first-stage sample size n0 ≥ 2. Let \( h^2 = (n_0 - 1) \left[ (2\alpha/(k-1))^{(2/(n_0-1))} - 1 \right] \).

Step 1. Initialization. Let \( I = \{1, 2, \ldots, k\} \) be the set of alternatives still in contention. Processor 0 manages the input sequence in which all alternatives are stored in a round-robin order from 1 to k. Processor 0 also performs the tasks listed in the following steps 2 through 4, including conducting pairwise comparisons and eliminations. The first \( m \) replications in the input sequence are assigned to the remaining \( m \) processors, processors 1, 2, . . . , m, to be simulated. Processors 1, 2, . . . , m work as follows: take the first alternative queued in the input sequence, generate an observation, and submit the result to processor 0.

The \( i \)th replication from alternative \( i \) in the input sequence is denoted as \( X_{ij} \). These available observations are stored in a vector in the same order as the input sequence. Let \( n_i = \max \{n \geq 0 : X_{ij} \text{ is available for all } l \leq n \text{ and for } i \in I\} \). Notice that \( n_i = 0 \) at the beginning of simulation. Start the simulation.

Step 2. Variance Estimation. When \( n_a \geq n_0 \), compute the sample variance of the difference between alternatives \( i \neq j \),

\[
S_{ij} = n_0 - 1 \sum_{l=1}^{n_0} (X_{il} - X_{jl}) - \left( \bar{X}_i(n_0) - \bar{X}_j(n_0) \right)^2,
\]

where \( \bar{X}_i(n_0) \) is the first-stage sample mean of alternative \( i \) with \( n_0 \) observations. Set \( r = n_0 \).

Step 3. Elimination. Set \( I^{r, \text{old}} = I \).

\[
I = I^{r, \text{old}} \setminus \{i \in I^{r, \text{old}} : \bar{X}_i(r) - \bar{X}_j(r) < \min(0, -h^2 S_{ij}^2 /(2r\delta) + \delta/2) \}
\]

for some \( j \in I^{r, \text{old}}, j \neq i \),

where \( A \setminus B = \{x : x \in A \text{ and } x \notin B\} \), and remove alternative \( i \) from the input sequence for all \( i \in I^{r, \text{old}} \).

Step 4. Stopping Rule. If \( |I| = 1 \), then stop all processors and select the alternative whose index is in \( I \) as the best. Otherwise, processor 0 checks whether it is ready for the next elimination. Let \( r = r + 1 \).

(a) If \( r \leq n_a \), go to step 3.

(b) Otherwise, wait for a new observation from any alternative \( j \) in \( I \), say \( X_{ij} \); record \( X_{ij} \) in the vector; update \( n_i \); and go to (a).

Remark 1. The statistical validity of the vector-filling \( \mathcal{KN} \) (VKN) procedure is the same as that of \( \mathcal{KN} \) because, statistically, the two procedures are identical in conducting comparisons and making elimination decisions.

Remark 2. The VKN procedure may require a large amount of memory to store simulation outputs exactly following the order in the input sequence, especially when the R&S problem has a large number of alternatives and the variances of replication times are high. For instance, when we apply the VKN procedure to solve a test problem with 10^4 alternatives on a personal computer with 4.00 GB RAM (see §5 for detailed settings), we encountered situations where the procedure has to be terminated after using up all of the memory. One may design careful accounting schemes that would dramatically reduce the memory requirements, e.g., by storing cumulative sums up to the point where all replications in the input sequence have returned.

**4. Asymptotic Parallel Selection Procedures**

The nice asymptotic properties of the sample-mean estimators \( \bar{Y}_i(n) \) in §2 motivate us to design fully sequential selection procedures that are asymptotically valid and are computationally more efficient than VF procedures. Our goal is to design a simple and easily executable, fully sequential procedure that uses all simulation observations in the output sequence, allows different surviving alternatives to have different sample sizes, and has a provable asymptotic validity in a meaningful asymptotic regime.

To design such a procedure the key is to decide when to compare surviving alternatives and make elimination decisions. For that we introduce the concept of a phantom alternative, which is an alternative that does not need to be simulated (i.e., simulation time is zero) and compared and is used only for counting purposes. We add the phantom alternative after each round-robin cycle in the input sequence and then start the simulation. Whenever the phantom alternative completes (i.e., appears in the output sequence), we compare all surviving alternatives using all their available observations (in the output sequence) and make elimination decisions. Notice that when there is only a single processor, the phantom alternative completes at the moment that the last surviving alternative in the round-robin cycle completes; then comparing and eliminating according to the phantom alternative are exactly the same as what the \( \mathcal{KN} \) procedure does. When there are multiple processors, different surviving alternatives may have different sample sizes at the moment that a phantom alternative completes. However, the difference between the sample size of a surviving alternative and that of the phantom alternative is always bounded by the number of processors. Therefore, the difference between the sample means of a surviving alternative computed based on input and output sequences may vanish as the sample size of the alternative goes to infinity. This provides a key to insuring the asymptotic validity of the procedure. Therefore, the completion times of the phantom alternative serves as a drumbeat process that synchronizes the comparisons and eliminations and insures the asymptotic validity.
**Procedure 2 (Asymptotic Parallel Selection (APS) Procedure).**

*Step 0. Setup.* Select confidence level $1/k < 1 - \alpha < 1$, IZ parameter $\delta > 0$, and first-stage sample size $n_0 \geq 2$. Let $a = -\log(2\alpha/(k-1))$.

*Step 1. Initialization.* Let $I = \{1, 2, \ldots, k\}$ be the set of alternatives still in contention. Processor 0 manages the input sequence in which all alternatives are stored in a round-robin order from 1 to $k$. Processor 0 also performs the tasks listed in the following steps 2 through 4, including conducting pairwise comparisons and eliminations. The first $m$ replications in the input sequence are assigned to the remaining $m$ processors, processors $1, 2, \ldots, m$, to be simulated. Processors $1, 2, \ldots, m$ work as follows: take the first alternative queued in the input sequence, generate an observation, and submit the result to processor 0.

Add a phantom alternative $p$ after each round-robin cycle in the input sequence (but not the set $I$). Let $r$ denote the stage that is the current sample size of the phantom alternative in the output sequence. Let $Y_{il}^{(r)}$ denote the $l$th completed observation from alternative $i$ in the output sequence, and let $N_{il}$ denote the number of completed observations from alternative $i$ in the output sequence at the time when the $r$th observation of the phantom alternative is added to the output sequence. Record the triple $(N_{il}, \sum_{l=1}^{N_{il}} Y_{il}^{(r)}, \sum_{l=1}^{N_{il}} Y_{il}^{(r)^2})$ for all $i \in I$.

*Step 2. Collecting Initial Observations.* Start simulations on processors $1, 2, \ldots, m$ and wait until $r = n_0$.

*Step 3. Elimination.* For all $i \in I$, let
\[
\bar{Y}(N_{il}) = \frac{1}{N_{il}} \sum_{l=1}^{N_{il}} Y_{il}^{(r)},
\]
\[
S_i^2(N_{il}) = \frac{1}{N_{il}-1} \left[ \sum_{l=1}^{N_{il}} Y_{il}^{(r)} - \frac{1}{N_{il}} \left( \sum_{l=1}^{N_{il}} Y_{il}^{(r)} \right)^2 \right].
\]
For all $i, j \in I$ and $i \neq j$, if $N_{il} \geq n_0$ and $N_{ij} \geq n_0$, let
\[
\tau_{ij,r} = \left[ \frac{S_i^2(N_{il})}{N_{il} - 1} + \frac{S_j^2(N_{ij})}{N_{ij} - 1} \right]^{-1};
\]
otherwise, let $\tau_{ij,r} = 0$. This ensures that the comparisons are done between alternatives that have at least $n_0$ observations. Let $I^{\text{old}} = I$ and let
\[
I = I \setminus \{i \in I^{\text{old}} : \tau_{ij,r} \approx (\bar{Y}(N_{il}) - \bar{Y}(N_{ij})) \]
\[
< \min \{0, -a/\delta + \delta/2\tau_{ij,r} \}
\]
for some $j \in I^{\text{old}}$ and $j \neq i$.

Remove alternative $i$ from the input sequence for all $i \in I^{\text{old}} \setminus I$.

*Step 4. Stopping Rule.* If $|I| = 1$, then stop all processors and select the alternative whose index is in $I$ as the best. Otherwise, wait for a new observation. If the new observation is from any alternative $i \in I$, then update $(N_{il}, \sum_{l=1}^{N_{il}} Y_{il}^{(r)}, \sum_{l=1}^{N_{il}} Y_{il}^{(r)^2})$ and wait for the next observation; if the observation is from alternative $p$, then update $r = r + 1$ and go to step 3.

**Remark 3.** In the APS procedure, we keep updating the sample variances for the surviving alternatives, which is similar to $\mathcal{N}^{++}$ of Kim and Nelson (2006a) designed for R&S problems in steady-state simulations. In order to show the asymptotic validity of sample-variance updating, we need some technical condition on the first-stage sample size $n_0$, which is stated in Theorem 1. Although the technical condition facilitates the asymptotic proof, it does not prescribe a specific choice of first-stage sample size in practice.

Notice that in the APS procedure, we only make elimination decisions when a phantom alternative completes, but we use all available observations at that time. Therefore, the APS procedure has several advantages when compared to the VKN procedure. First, it makes use of all available observations, which leads to a higher efficiency, especially when the simulation effort for generating each observation is substantial (e.g., the replication time is relatively long). Second, it requires significantly less memory to store the observations, which makes it feasible to solve large-scale R&S problems.

The following theorem establishes the asymptotic validity of the APS procedure.

**Theorem 1.** Let $X_{il}$ denote the $l$th replication from alternative $i$ in the input sequence and $\Gamma_{il}$ denote the replication time to generate the observation $X_{il}$, with unknown means $\mu_i = E[X_{il}]$ and unknown (but finite) variance $\sigma_i^2 = \text{Var}[X_{il}]$, for all $i = 1, 2, \ldots, k$ and $l = 1, 2, \ldots$. Assume that $X_{il}$ and $X_{im}$ are independent of each other when $i \neq j$ or $l \neq n_l$; that $\Gamma_{il} > 0$ a.s. for all $i$ and $l$; and that $\mu_i - \delta \geq \mu_j$, where $\delta$ is the IZ parameter. Moreover, let the first-stage sample size $n_0 = n_0(\delta)$ be a function of $\delta$ such that $n_0 \to \infty$ and $\delta n_0 \to 0$ as $\delta \to 0$. Then, as $\delta \to 0$, the APS procedure selects alternative 1 as the best with a probability at least $1 - \alpha$.

Notice that the asymptotic regime of $\delta \to 0$ (as well as the true difference between the best and the second-best alternatives $\mu_1 - \mu_2 \to 0$) is also used by Kim and Nelson (2006a) in analyzing the $\mathcal{N}^{++}$ procedure, where the observations are taken from steady-state simulations that are stationary but not independent. In Theorem 1, however, we assume that the observations ordered by the input sequence $\{X_{il}, l = 1, 2, \ldots\}$ are i.i.d. but, as shown in §2.3, the observations ordered based on the output sequence $\{Y_{il}, l = 1, 2, \ldots\}$ may not be.

We prove Theorem 1 in the next two subsections.

### 4.1. Brownian Motion Construction

For a clear presentation, in the remainder of this section we consider only the situation where the difference between the mean of the best and all other alternatives equals the IZ parameter $\delta$, i.e., $\mu_2 = \mu_3 = \cdots = \mu_\ell = \mu_1 - \delta$, which is called the slippage configuration (SC) in the R&S literature.
Consider any pair of alternatives, $i$ and $j$. Let $N^\delta_{ij} = \lfloor 2a(\sigma_i^2 + \sigma_j^2)/\delta^2 \rfloor$, where $\delta = -\log(2\alpha/(k-1))$ is defined in step 0 of the APS procedure and $[x]$ denotes the smallest integer not less than $x$. In $\S 4.2$, it will be clear that $N^\delta_{ij}$ is the maximum number of observations needed from either alternative $i$ or $j$ when comparing alternatives $i$ and $j$. Let $s$ be any number in $[0, 1]$ and let $r = \lfloor sN^\delta_{ij} \rfloor$, where $[x]$ denotes the largest integer not greater than $x$. Define

$$Z_{ij}(s) = \begin{cases} \frac{\sigma_i^2}{r} + \frac{\sigma_j^2}{r} \sqrt{N^\delta_{ij}} \left( \sqrt{\frac{N^\delta_{ij}}{r}} \cdot [\bar{Y}(N_{ir}) - \bar{Y}(N_{jr})], \quad s \in \left[ \frac{n_0}{N^\delta_{ij}} \right]; \\ 0, \quad s \in \left[ 0, \frac{n_0}{N^\delta_{ij}} \right); \end{cases}$$

(1)

where $N_{ir}$ is the sample size of alternative $l$, $l = 1, \ldots, k$, when the sample size of the phantom alternative $p$ is $N_{ip} = r$. To make $Z_{ij}(\cdot)$ well-defined, we set $Z_{ij}(\cdot) = 0$ when either $N_{ir} = 0$ or $N_{jr} = 0$. In fact, we are only interested in the case that $r \geq n_0$, i.e., $s \in \left[ \frac{n_0}{N^\delta_{ij}} \right]$, because the APS procedure starts eliminating systems after at least $n_0$ observations. For mathematical completeness in the neighborhood of $s = 0$, we can artificially set $N_{ir} = r$, $S^2(N_{ir}) = \sigma_i^2$ and $\bar{Y}(N_{ir}) = 0$, $l = i$ or $j$, in order to make sure that $Z_{ij}(s) = 0$ for $s \in \left[ 0, \frac{n_0}{N^\delta_{ij}} \right)$. However, without additional specifications, all statements about $Z_{ij}(s)$ will refer to the process for $s \in \left[ \frac{n_0}{N^\delta_{ij}} \right]$. 

Hong (2006) shows that $Z_{ij}(s)$ with the random sample size $N_{ir}$ replaced by the deterministic $n_i$, the sample variance $S^2(N_{ir})$ replaced by $\sigma_i^2$, and $\bar{Y}(N_{ir})$ replaced by $\bar{X}_{in}$, $l = i$ or $j$, has the same distribution as a Brownian motion when the $X_{in}$’s are normally distributed. This result motivates the definition of $Z_{ij}(s)$ in Equation (1). The following lemma shows that $Z_{ij}(\cdot)$ converges to a Brownian motion process for all $j = 2, 3, \ldots, k$.

**Lemma 1 (Convergence to a Brownian Motion Process).** Let $\Omega[0, 1]$ be the Skorohod space of all right-continuous real-valued functions on $[0, 1]$ with limits from the left everywhere, endowed with the Skorohod $J_1$ topology (see the e-companion EC.1.3 for the definition of the standard $J_1$ metric and Section 3.3 in Whitt 2002 for more background on the space $\Omega$). Thus, $Z_{ij}(\cdot)$ defined by Equation (1) with $j = 2, 3, \ldots, k$ is an element of the Skorohod space $\Omega[0, 1]$. Suppose that the conditions in Theorem 1 are all satisfied. Then, under the SC, i.e., $\mu_2 = \mu_3 = \cdots = \mu_k = \mu_1 = \delta$, we have

$$Z_{ij}(\cdot) \Rightarrow \mathcal{B}_\Delta(\cdot), \quad \text{as } \delta \to 0,$$

where $\mathcal{B}_\Delta(t) = \mathcal{B}(t) + \Delta t$, a standard Brownian motion process with a constant drift $\Delta = \sqrt{2a}$. 

**Proof of Lemma 1.** Let $\varepsilon^\delta = n_0/N^\delta_{ir}$, where $N^\delta_{ij} = \lfloor 2a(\sigma_i^2 + \sigma_j^2)/\delta^2 \rfloor$. Recall that $n_0 \to \infty$ and $\delta^2 n_0 \to 0$ as $\delta \to 0$. Then, $N^\delta_{ij} \to \infty$ and $\varepsilon^\delta \to 0$ as $\delta \to 0$. Notice that $Z_{ij}(s) = 0$ for $0 \leq s < \varepsilon^\delta$, which implies that

$$Z_{ij}(0) = \mathcal{B}_\Delta(0) = 0 \quad \text{and} \quad Z_{ij}(\cdot) \text{ is right-continuous at } s = 0 \text{ for all } \delta.$$

(2)

We next focus only on $s \in [\varepsilon^\delta, 1]$, that is, $r \in [n_0, N^\delta_{ij}]$. 

We start by analyzing the first term on the right-hand side (RHS) of Equation (1). Recall that, at stage $r$, for alternative $l$ ($l = 1$ or $j$), totally $r$ replications have been sent to the $m$ processors with $N_{ir}$ replications completely and $r - N_{ir}$ still in simulation. Then, we have $r - m \leq r - N_{ir}$, which implies that, w.p.1,

$$\sup_{r \in [n_0, N^\delta_{ij}]} \frac{N_{ir} - r}{r} \leq \frac{m}{n_0} \to 0,$$

as $\delta \to 0$. In other words, as $\delta \to 0$, $n_0 \to \infty$, so that $r \to \infty$ and $N_{ir}/r \to 1$ w.p.1 as functions of $s$ on $[0, 1]$. Notice that this result can be easily extended to the closed space $[0, 1]$ given the definition that $N_{ir} = r$ for $s \in [0, \varepsilon^\delta]$. In fact, the pointwise convergence guarantees the uniform convergence on $[0, 1]$ under the definition that $N_{ir} = r$ on $[0, \varepsilon^\delta]$ and the condition that $\varepsilon^\delta \to 0$ and $\varepsilon^\delta N^\delta_{ij} \to \infty$ as $\delta \to 0$. Similar arguments can also be applied to $S^2(N_{ir})$ and $\bar{Y}(N_{ir})$, $l = 1$ or $j$. Therefore, we will establish uniform convergence by showing pointwise convergence in the following proof.

Let $\Omega_{ir} \subset \{1, 2, \ldots, r\}$ be the set of the indices of incomplete replications from alternative $l$; i.e., if $n \in \Omega_{ir}$, $X_{in}$ is still in simulation at stage $r$. Thus, $|\Omega_{ir}| = r - N_{ir}$. Notice that, for $l = 1$ or $j$,

$$S^2(N_{ir}) = \frac{1}{N_{ir} - 1} \left[ \sum_{n = 1}^{N_{ir}} X_{in}^2 - \frac{N_0}{N_{ir}} \left( \sum_{n = 1}^{N_{ir}} X_{in} \right)^2 \right],$$

which can be further written as

$$S^2(N_{ir}) = \frac{1}{N_{ir} - 1} \left[ \sum_{n = 1}^{N_{ir}} X_{in}^2 - \sum_{n = \Omega_{ir}} X_{in}^2 \right] \cdot \frac{N_0}{N_{ir}} \left( \sum_{n = 1}^{N_{ir}} X_{in} \right)^2$$

$$= \frac{1}{N_{ir} - 1} \left[ \sum_{n = 1}^{N_{ir}} X_{in}^2 - \frac{1}{r} \left( \sum_{n = 1}^{N_{ir}} X_{in} \right)^2 \right]$$

$$+ \frac{1}{N_{ir} - 1} \left[ \sum_{n \in \Omega_{ir}} X_{in}^2 + \frac{(N_{ir} - r- r) \bar{X}_{in}}{N_{ir}} \right]$$

$$+ \frac{2r}{N_{ir}} \bar{X}_{in} \sum_{n \in \Omega_{ir}} X_{in} - \frac{1}{N_{ir}} \left( \sum_{n \in \Omega_{ir}} X_{in} \right)^2 \right].$$

(3)

Similarly, as $\delta \to 0$ (which implies that $n_0 \to \infty$ and $r \to \infty$; for simplicity, we may state only one of them hereafter), then $r/(N_{ir} - 1) \to 1$ w.p.1. By the strong law of large numbers, we know that $r^{-1} \sum_{n = 1}^{N_{ir}} X_{in} \to E\left[ X_{in}^2 \right] = \sigma^2_i + \mu_2^2$, and $\bar{X}_{in} \to r^{-1} \sum_{n = 1}^{N_{ir}} X_{in} \to \mu_i$ w.p.1. By the
continuous mapping theorem (Durrett 2004), the first term of $S^2_t(N_i)$ in Equation (3) converges to $\sigma^2_1$ w.p.1 as $r \to \infty$.

Notice that, for both $l = 1$ and $j$ and all $n = 1, 2, \ldots, r$,

$$|X_{ln}| \leq \max_{l = 1, \ldots, r} |X_{ln}|.$$

Furthermore, we have $N_i \geq r - m$ and $|\Omega_i| \leq m$. Then, when $n_0 > m + 1$, implying that $r > m + 1$, the second term of $S^2_t(N_i)$ in Equation (3) can be bounded as follows,

$$\frac{1}{N_{lr} - 1} \left[ - \sum_{n \in \Omega_{lr}} X^2_n + \frac{(N_{lr} - r)r}{N_{lr}} \tilde{X}(r)^2 \right] + \frac{2r}{N_{lr}} \tilde{X}(r) \sum_{n \in \Omega_{lr}} X_n - \frac{1}{N_{lr}} \left( \sum_{n \in \Omega_{lr}} X_n \right)^2$$

$$\leq \frac{m r}{r - m} \left[ \frac{1}{r} \max_{r - m + 1} X^2_n + \frac{r}{r - m} \tilde{X}(r)^2 + \frac{r}{r - m} \tilde{X}(r) \sum_{n \in \Omega_{lr}} X_n \right] \leq \frac{1}{r - m} \max_{r - m + 1} X^2_n + \frac{r}{r - m} \tilde{X}(r)^2 \left( \frac{1}{r - m} \max_{r - m + 1} |X_n| \right)^2.$$

Recall that $r^{-1} \sum_{n=1}^r X^2_n \to \sigma^2_1 + \mu^2_j$ and $r^{-1} \sum_{n=1}^r X_n \to \mu_j$ w.p.1. Then, by Lemma EC.1 in the e-companion, we have

$$\frac{1}{r - m} \max_{r - m + 1} X^2_n \to 0 \text{ and } \frac{r}{r - m} \max_{r - m + 1} |X_n| \to 0 \text{ w.p.1.}$$

By the continuous mapping theorem, the second term of $S^2_t(N_i)$ converges to 0 w.p.1 as $r \to \infty$. Therefore, as $\delta \to 0$, $S^2_t(N_i) \to \sigma^2_1$ w.p.1. By continuous mapping theorem again, we obtain that

$$\frac{\sigma^2_1}{r} + \frac{\sigma^2_j}{r} \to 1 \text{ w.p.1 as } \delta \to 0. \quad (4)$$

We next analyze the second term on the RHS of Equation (1). Similarly to what we did above, we can write

$$\tilde{Y}(N_{lr}) = \frac{1}{N_{lr}} \sum_{n=1}^{N_{lr}} Y_n = \frac{1}{N_{lr}} \left( \sum_{n=1}^{N_{lr}} X_n - \sum_{n \in \Omega_{lr}} X_n \right)$$

$$= \frac{1}{N_{lr}} \left( \sum_{n=1}^{N_{lr}} (X_n - \mu_j) + \frac{r}{N_{lr}} \mu_j - \frac{1}{N_{lr}} \sum_{n \in \Omega_{lr}} X_n \right)$$

$$= \sigma_i \sqrt{N^{\delta}_{lj}} \cdot H_i(s) + \frac{r}{N_{lr}} \mu_j - \frac{1}{N_{lr}} \sum_{n \in \Omega_{lr}} X_n,$$

where

$$H_i(s) = \sum_{n=1}^{N_{lr}} (X_n - \mu_j) \sigma_i \sqrt{N^{\delta}_{lj}} \cdot \frac{\sigma_i}{\sigma_i \sqrt{N^{\delta}_{lj}}} = \sum_{n=1}^{N_{lr}} \frac{(X_n - \mu_j)}{\sigma_i \sqrt{N^{\delta}_{lj}}}, \quad s \in [\epsilon^\delta, 1],$$

and the last equality holds because $r = \lfloor N_{lj} \delta \rfloor$. Then we have

$$\sqrt{s} \sqrt{\frac{N^{\delta}_{lj}}{N_{lr}}} \left[ \tilde{Y}(N_{lr}) - \tilde{Y}(N_{lj}) \right]$$

$$= \left( \frac{N^{\delta}_{lj}}{N_{lr}} \cdot \frac{\sigma_i}{\sqrt{\sigma^2_i + \sigma^2_j}} \cdot H_i(s) - \frac{N^{\delta}_{lj}}{N_{lr}} \cdot \frac{\sigma_j}{\sqrt{\sigma^2_i + \sigma^2_j}} \cdot H_j(s) \right)$$

$$+ \sqrt{s} \frac{N^{\delta}_{lj}}{\sigma^2_i + \sigma^2_j} \left( \frac{r}{N_{lr}} \mu_1 - \frac{r}{N_{lr}} \mu_j \right)$$

$$- \sqrt{s} \frac{N^{\delta}_{lj}}{\sigma^2_i + \sigma^2_j} \left( \frac{1}{N_{lr}} \sum_{n \in \Omega_{lr}} X_n - \frac{1}{N_{lr}} \sum_{n \in \Omega_{lr}} X_n \right). \quad (5)$$

By Donsker’s Theorem (Whitt 2002, Theorem 4.3.2), as $\delta \to 0$,

$$H_i(s) \Rightarrow \mathcal{B}^i(s),$$

where $\mathcal{B}^i(s)$ is a standard Brownian motion process. Furthermore, because $H_i(s)$ and $H_j(s)$ are independent of each other, we have $\mathcal{B}^i(s)$ and $\mathcal{B}^j(s)$ are also independent of each other. Recall that $r = \lfloor N_{lj} \delta \rfloor$ and $r/N_{lr} \to 1$ w.p.1 as $\delta \to 0$. Then $N_{lj} \delta / N_{lr}$, as a function of $s$, converges to the function that is identically equal to 1 w.p.1, for $l = 1$ or $j$. By Whitt (2002, Theorem 11.4.5), we have

$$\sqrt{s} \sqrt{\frac{N^{\delta}_{lj}}{N_{lr}}} \cdot \frac{\sigma_i}{\sqrt{\sigma^2_i + \sigma^2_j}} \cdot H_i(s) - \sqrt{s} \sqrt{\frac{N^{\delta}_{lj}}{N_{lr}}} \cdot \frac{\sigma_j}{\sqrt{\sigma^2_i + \sigma^2_j}} \cdot H_j(s) \Rightarrow \left( \frac{\sigma_i}{\sqrt{\sigma^2_i + \sigma^2_j}} \cdot \mathcal{B}^i(s) + \frac{\sigma_j}{\sqrt{\sigma^2_i + \sigma^2_j}} \cdot \mathcal{B}^j(s) \right) \text{ w.p.1 as } s \in [0, 1].$$

(6)

where the last equation follows from the independence of $\mathcal{B}^i(s)$ and $\mathcal{B}^j(s)$ and $\mathcal{B}$ means “equal in distribution.”

Because $r/N_{lr} \to 1$ and $r/N_{lj} \to 1$ w.p.1 as $\delta \to 0$, $\mu_j - \mu_j = \delta$ and $N_{lj} \delta = [2\alpha(\sigma^2_i + \sigma^2_j) / \delta^2]$, we have as $\delta \to 0$,

$$s \sqrt{\frac{N^{\delta}_{lj}}{\sigma^2_i + \sigma^2_j}} \cdot \frac{r}{N_{lr}} \mu_1 - \frac{r}{N_{lr}} \mu_j \Rightarrow \sqrt{2\alpha} = s \Delta \quad (7)$$

w.p.1 for all $s \in (0, 1)$, where $\Delta = \sqrt{2\alpha}$ by definition.

Because $r = \lfloor N_{lj} \delta \rfloor$, we have $N_{lj} \delta \leq r + 1$. Recall that $N_i \geq r - m$ and $|\Omega_i| \leq m$. Then, when $n_0 \geq m + 1$ implying that $r \geq m + 1$,

$$s \sqrt{\frac{N^{\delta}_{lj}}{\sigma^2_i + \sigma^2_j}} \cdot \frac{1}{N_{lr}} \sum_{n \in \Omega_{lr}} X_n$$

$$\leq \sqrt{\frac{(r + 1)s}{\sigma^2_i + \sigma^2_j}} \cdot \frac{m}{r - m} \max_{n = 1, \ldots, r} |X_n|$$

$$= \sqrt{s \frac{m^2}{\sigma^2_i + \sigma^2_j}} \cdot \frac{\sqrt{(r + 1)s}}{r - m} \cdot \frac{1}{\sqrt{r}} \max_{n = 1, \ldots, r} |X_n|.$$
We have already shown that
\[ \frac{1}{r} \sum_{n=1}^{r} X_{in}^2 \rightarrow 0 \text{ w.p.1} \]
as \( r \rightarrow \infty \), which implies that
\[ \frac{1}{\sqrt{r}} \max_{n=1, \ldots, r} |X_{in}| \rightarrow 0 \text{ w.p.1} \]
as \( r \rightarrow \infty \). Notice also that \( \sqrt{r(r+1)/r} \rightarrow 1 \) as \( r \rightarrow \infty \). Therefore, as \( \delta \rightarrow 0 \),
\[ s \left( \frac{N_{ij}^0}{\sigma_i^2 + \sigma_j^2} \right)^{1/2} \frac{1}{N_{\ell r}^0} \sum X_{in} \rightarrow 0 \]w.p.1 for both \( l = 1 \) and \( j \).

Notice that (6), (7), and (8) correspond to the limits of the three terms on the RHS of Equation (5), respectively. Therefore, by Whitt (2002, Theorem 11.4.5), we have as \( \delta \rightarrow 0 \),
\[ \left( \sqrt{\frac{N_{ij}^0}{\sigma_i^2 + \sigma_j^2}} \left[ \tilde{Y}_i(N_{\ell r}) - \tilde{Y}_j(N_{\ell r}) \right] : 0 < s \leq 1 \right) \]
⇒ \( \mathcal{B}(s) + s\Delta = (\mathcal{B}_\Delta(s); 0 < s \leq 1) \).

Then by (4) and (9), and by in Whitt (2002, Theorem 11.4.5) again, we have \( Z_{ij}(\cdot) \Rightarrow \mathcal{B}_\Delta(\cdot) \) on \( [0, 1] \) as \( \delta \rightarrow 0 \). Combined with the result in Equation (2), we conclude the proof of the lemma.

**Remark 4.** Lemma 1 establishes the foundation for showing the statistical validity of the APS procedure. Notice that the proof of Lemma 1 does not require the condition that \( \Gamma_i \) and \( \Gamma_n \) are independent for \( i \neq j \) or \( l \neq n \). This indicates that the APS procedure can be implemented in parallel computing environments where the multiple processors are not identical so that the replication times may be dependent on each other.

### 4.2. The Asymptotic Validity

Before proving the asymptotic validity of the APS procedure, we first define the continuation region that determines the elimination decisions in the procedure. Let
\[ U_{ij}^0(s) = \max \left\{ 0, \frac{a}{\Delta} - \frac{\Delta}{2} \cdot s \right\}, \text{ w.p.1 as } \delta \rightarrow 0, \]
where \( \Delta = \sqrt{2}a \). Then the asymptotic region \( C \), formed by \( U(s) \) and \( -U(s) \), is a symmetric triangular region.

Let \( T_{ij}^\delta \) denote the stopping time at which \( Z_{ij}(\cdot) \) first exits the continuation region \( C_{ij}^\delta \), i.e.,
\[ T_{ij}^\delta = \inf \{ s : |Z_{ij}(s)| \geq U_{ij}^\delta(s) \}, \]
and let \( T_{ij} \) denote the stopping time at which \( \mathcal{B}_\Delta(\cdot) \) first exits the triangular region \( C \), i.e.,
\[ T_{ij} = \inf \{ s : |\mathcal{B}_\Delta(s)| \geq U(s) \}. \]

Lemma 1 establishes the weak convergence of \( Z_{ij}(\cdot) \) to \( \mathcal{B}_\Delta(\cdot) \) on \([0, 1]\). However, elimination decisions are only made at these stopping times. To bound the probability of incorrect selection, we need a stronger result that ensures the value at the stopping time \( Z_{ij}(T_{ij}^\delta) \) can be approximated by \( \mathcal{B}_\Delta(T_{ij}) \), which can be guaranteed by the following lemma.

**Lemma 2** (Convergence to a Brownian Motion Process at the Stopping Time). Suppose that the conditions in Theorem 1 are all satisfied. Then
\[ Z_{ij}(T_{ij}^\delta) \Rightarrow \mathcal{B}_\Delta(T_{ij}) \]
as \( \delta \rightarrow 0 \).

**Remark 5.** The key idea for proving Lemma 2 is exactly the same as that of proving Kim et al. (2005, Proposition 3.2). We summarize the proof in the e-companion EC.1.3 for completeness.

To prove the validity of the APS procedure, we also need the lemma of Fabian (1974), i.e., Lemma EC.2 in the e-companion, on the probability of \( \mathcal{B}_\Delta(\cdot) \) exiting the triangular continuation region \( C \). This lemma is the foundation of many sequential R&S procedures, including those of Kim and Nelson (2001, 2006a) and Hong and Nelson (2005, 2007). Now we are ready to prove Theorem 1.

**Proof of Theorem 1:** We start from the slippage configuration where \( \mu_1 - \delta = \mu_2 = \cdots = \mu_k \). Then we have
\begin{align*}
\lim_{\delta \rightarrow 0} \mathbb{P} \{ \text{select alternative 1} \} \\
= \lim_{\delta \rightarrow 0} \left[ 1 - \mathbb{P} \left( \bigcup_{j=1}^{k-1} \{ \text{alternative } j \text{ eliminates 1} \} \right) \right] \\
\geq 1 - \lim_{\delta \rightarrow 0} \sum_{j=1}^{k-1} \mathbb{P} \{ \text{alternative } j \text{ eliminates 1} \},
\end{align*}
where (12) is due to Bonferroni inequality. Notice that
\begin{align*}
\lim_{\delta \rightarrow 0} \mathbb{P} \{ \text{alternative } j \text{ eliminates 1} \} \\
= \lim_{\delta \rightarrow 0} \mathbb{P} \{ Z_{ij}(T_{ij}^\delta) \leq 0 \} \\
= \mathbb{P} \{ \mathcal{B}_\Delta(T_{ij}) \leq 0 \} \\
= \frac{1}{2} e^{-\left(\frac{\alpha}{k-1}\right)} \leq \frac{\alpha}{k-1},
\end{align*}
where (13) denotes the probability that alternative \( j \) eliminates alternative 1 since \( Z_{1j}(\cdot) \) exits the continuation region through the lower boundary, (14) follows from Lemma 2, and (15) follows from Lemma EC.2 in the e-companion. Plugging (15) into (12) yields

\[
\lim \inf_{\delta \to 0} \mathbb{P} \{ \text{select alternative 1} \} \geq 1 - \sum_{j=1}^{k-1} \frac{\alpha}{k-1} = 1 - \alpha.
\]

For general cases under the IZ formulation, i.e., \( \mu_1 - \delta \geq \mu_2 \geq \cdots \geq \mu_k \), \( Z_{ij}(\cdot) \) defined in (1) no longer converges in distribution to \( \mathcal{B}_\alpha(\cdot) \). However, we can define

\[
V_{ij}(s) = \frac{\sigma^2_j / r + \sigma^2_r / r}{S'_j(N_{ir})/N_{ir} + S'_j(N_{jr})/N_{jr}} \cdot s \sqrt{N^b_{ij}} \cdot \left( \tilde{Y}_i(N_{ir}) - \tilde{Y}_j(N_{jr}) - (\mu_i - \mu_j - \delta) \right).
\]

Then

\[
V_{ij}(s) \leq Z_{ij}(s), \quad \text{a.s.} \quad (16)
\]

By Lemma 1, we know that \( V_{ij}(\cdot) \Rightarrow \mathcal{B}_\alpha(\cdot) \) as \( \delta \to 0 \). Let

\[
T^{V}_{ij} = \inf \{ s : |V_{ij}(s)| \geq U^b_{ij}(s) \}.
\]

Then

\[
\lim \sup_{\delta \to 0} \mathbb{P} \{ \text{alternative j eliminates 1} \} = \lim \sup_{\delta \to 0} \mathbb{P} \{ Z_{ij}(T^V_{ij}) \leq 0 \} = \lim \sup_{\delta \to 0} \mathbb{P} \{ V_{ij}(T^V_{ij}) \leq 0 \} = \mathbb{P} \{ \mathcal{B}_\alpha(T_{ij}) \leq 0 \} = \frac{\alpha}{k-1}, \quad (18)
\]

where (17) follows from (16). Plugging (18) into (12) concludes the proof of the theorem.

5. Numerical Implementation

In this section, we report on an extensive numerical study to test the effectiveness and efficiency of both the VKN procedure and the APS procedure and their applicability to solve large-scale R&S problems in parallel computing environments.

5.1. Master/Slave Structure and a Parallel Computing Simulator

We design a parallel computing environment using the master/slave structure, a widely used structure for parallel computing, which contains two functions: a single master and multiple slaves. The master maintains data information for all alternatives and manipulates two daemon threads (daemon threads can be viewed as service providers for other threads running in the same program. When the only remaining threads are daemon threads, the program will exit automatically), called “to-do” and “compare.” The to-do thread manages the input sequence of all surviving alternatives and the compare thread conducts pairwise comparisons and elimination decisions based on the simulation observations collected from the slaves. Each slave, created as a daemon thread, works in a very simple cycle: taking an alternative from the to-do thread, generating an observation, and submitting the observation to the compare thread for comparison. In the procedures, we denote the master as processor 0 and the slaves as processors 1, 2, \ldots, \( m \).

This parallel structure is programmed in Java and can be easily implemented on various computer configurations, e.g., Windows operating systems on personal computers or Linux operating systems on local servers. Moreover, with a communication protocol (e.g., HTTP), it can be extended to computer farms or clouds. For more introduction about the master/alive structure, we refer to Silvay and Buyya (1999) for general details and Fujimoto et al. (2010) for an implementation in the cloud. We implement our procedures using the master/slave structure on a local server with 48 working cores and 64 GB memory. The server runs CentOS 6.2, a Linux-based operating system.

With respect to the simulation experiments that have been used to evaluate existing R&S procedures in the literature, the observations are often generated in a very simple way; e.g., Kim and Nelson (2001) simulate one observation by generating a normal random variable, so the replication time for generating one observation tends to be extremely short. This is a common and reasonable approach for testing procedures in a single-processor computing environment, where replication times do not affect the output sequence and therefore do not affect the properties of the procedure, as evaluated by the probability of correct selection, or the efficiency of the procedure, as evaluated by the expected total number of observations. In a parallel computing environment, however, replication times affect the output sequence, affecting both the properties and efficiency of the procedure. Therefore, we need to take them into consideration in experimental designs.

There are two approaches to evaluating the impact of (random) replication times. The first one is to make the slaves sleep artificially for a certain amount of time. However, this approach may be very time consuming in testing large-scale R&S problems with a large number of processors because replication times cannot be set too small for two reasons. One is due to the limitation of Java, which may lose accuracy when the elapsed time is less than one millisecond. In addition, it takes time to wake up a sleeping thread and it is difficult to fully control the frequent sleep-active cycles in a punctual manner. The other reason is the potential overhead on the master caused by comparison work. No matter how observations are generated, either in parallel or sequentially, they must be recorded one by one in the master for pairwise comparison. Even
if time for processing one observation is relatively small (within 0.1 milliseconds by a rough estimation), it could cause many observations to be queuing in front of the master when the unit comparison time exceeds the ratio of the replication time to the number of slaves. Such cases are not representative of real situations.

These concerns motivate us to consider a second approach to test parallel computing R&S procedures: we build a simulator on a single processor to simulate the situations in a parallel computing environment. As discussed in §2.1, experiments in a parallel computing environment can be considered as a multiserver queue. Thus, they may be simulated using a typical discrete-event simulation mechanism, where replication times are simulated under the simulation clock instead of the real clock. By managing the simulation clock properly, we can guarantee correct logic of events happening on the simulator.

In §5.2 we conduct an extensive numerical study on the simulator to evaluate the effectiveness and sampling efficiency of both the VKN and the APS procedures. In §5.3 we apply the APS procedure to solve a practical R&S problem with more than 20,000 alternatives in an actual parallel computing environment and analyze its performance.

5.2. The Effectiveness and Sampling Efficiency Tests

We assume that $X_i$ follows a normal distribution with mean $\mu_i$ and variance $\sigma_i^2$, and $\Gamma_i$ follows an exponential distribution with mean $E[\Gamma_i] = \gamma$ for all $i = 1, 2, \ldots, k$ and $l = 1, 2, \ldots$. To study how dependence between $X_i$ and $\Gamma_i$ affects the performance of the procedure, we consider three scenarios in which $X_i$ and $\Gamma_i$ are independent, positively correlated, and negatively correlated, respectively. For these three scenarios, we can use the NORTA method of Cario and Nelson (1998) to generate $\Gamma_i$ and $X_i$ as follows,

$$\Gamma_i = -\gamma \log(1 - \Phi(W_{i1}))$$

$$X_i = \mu_i + \sigma_i \left( \rho W_{i1} + \sqrt{1 - \rho^2} W_{i2}^2 \right)$$

where $W_{i1} = (W_{i1}^1, W_{i2}^2)$ is a bivariate standard normal vector with correlation zero. When $\rho = 0$, $X_i$ and $\Gamma_i$ are independent, and when $\rho > 0$ (or $< 0$), $X_i$ and $\Gamma_i$ are positively (or negatively) correlated.

We first consider the slippage configuration (SC) of means where $\mu_1 = \delta, \mu_2 = \mu_3 = \cdots = \mu_k = 0$ and the equal-variance configuration where $\sigma_i = 1$ for all $i$. The main goal of the experiment is to demonstrate that our procedures can solve large-scale problems using multiple processors, so we vary the number of alternatives from $k = 10^3$ to $k = 10^5$ and the number of processors as $m = 4, 48, 96$. The first-stage sample size is fixed to $n_0 = 16$, and the IZ parameter is specified as $\delta = 1/\sqrt{n_0}$. The expected replication time is $\gamma = 100$ units (in simulation clock time), and the correlation is $\rho = 0, \rho = 0.8$, and $\rho = -0.8$ for independent, positively correlated, and negatively correlated cases, respectively. The targeted probability of correct selection (PCS) is set to 0.95; i.e., $1 - \alpha = 0.95$. To achieve two-digit precision of the estimated PCS, we made 1,000 macroreplications in all configurations.

The SC is often considered as a difficult configuration since all inferior alternatives are close to the best. For many practical large-scale problems, a substantial number of the inferior alternatives may be significantly different from the best. Therefore, we consider another configuration of means, called grouped-decreasing-means (GDM) configuration, in which 10%, 20%, 30%, and 40% of the alternatives are $\delta$, $2\delta$, $3\delta$, and $4\delta$ different from the best. The means of GDM are defined as follows,

$$\mu_i = \begin{cases} 
\delta, & i = 1, \\
0, & i = 2, \ldots, [0.1k] + 1, \\
-\delta, & i = [0.1k] + 2, \ldots, [0.3k] + 1, \\
-2\delta, & i = [0.3k] + 2, \ldots, [0.6k] + 1, \\
-3\delta, & i = [0.6k] + 2, \ldots, k.
\end{cases}$$

To make the test problem more difficult to solve, we consider an increasing-variance configuration where $\sigma^2_i = |\mu_i - \delta| + 2\delta$, and the IZ parameter is $\delta = 0.5$. The first-stage sample size for the GDM is set to $n_0 = 10$.

In Table 1 we summarize the simulation results for both the VKN and APS procedures in all scenarios under the SC settings when the number of alternatives is $k = 10^4$. We report the average total number of observations generated (Total Samples) with 95% confidence interval, the average simulation time for completing one macroreplication of either procedure (Makespan) with 95% confidence interval, and the estimated PCS, across 1,000 macroreplications.

From the table we have several findings. First, both the VKN and APS procedures can deliver the desired PCS, but the VKN procedure tends to be more conservative than the APS procedure, which may be because the APS procedure uses variance updating and it is valid only asymptotically. Second, it seems that the correlations between performance outputs and replication times do not play an important role. Third, the total sample sizes needed for different numbers of processors $m$ are almost the same and the makespan reduces linearly as $m$ increases, which implies multiple processors are attractive for R&S problems. However, it is worthwhile pointing out that the makespan is computed without considering the time for processing elimination decisions on the simulator, which results a linear speedup. A linear speedup is seldom achieved in an actual parallel computing environment because the processing capacity of the master and the overhead of I/O (input/output) between the master and the slaves could affect the speedup as the number of processors increases. Intuitively, if there are too many slaves and the replication times are too short, then it is inevitable that many observations are ready to be sent back to the master for elimination while the master is not able to finish pairwise comparisons immediately; as a consequence, this may accumulate a queue in front of the master (see the e-companion EC.2 for an example).
For the GDM configuration, to avoid reporting similar results, we consider only the independent case (correlation $\rho = 0$) using the APS procedure when the number of processors is $m = 48$ and the number of alternatives varies from $k = 1 \times 10^3, 2 \times 10^3, \ldots, 10^4$. The estimated PCS for each $k$ is always greater than the desired level 0.95. Figure 6 plots the average total sample size for different $k$'s, from which we see that the total number of samples appears to increase almost linearly.

5.3. The Three-Stage Buffer Allocation Problem

We consider a three-stage flowline with a finite number of buffer storage locations in front of stations 2 and 3 (including the one in service at each station, denoted as $x_2$ and $x_3$) and an infinite number of jobs in front of station 1 (see Buzacott and Shanthikumar 1993, Pichitlamken et al. 2006, Xu et al. 2010). There is a single server at each station, and the service time at station $i$ is exponentially distributed with service rate $x_i$, $i = 1, 2, 3$. If the buffer of station $i$ is full, then station $i - 1$ is blocked (i.e., production blocking) and a finished job cannot be released from station $i - 1$. The total number of buffer locations and the total service rates are limited. The goal is to find an allocation of buffer locations and service rates such that the steady-state throughput of the flowline is maximized. The constraints of this problem are $x_1 + x_2 + x_3 \leq 20$, $x_4 + x_5 = 20$, $1 \leq x_i \leq 20$, and $x_i \in \mathbb{Z}_+$ for $i = 1, 2, \ldots, 5$. The problem has totally $k = 21,660$ feasible solutions. For any feasible solution, the throughput is estimated from running a simulation experiment with total simulation time being 1,000 units and the

<table>
<thead>
<tr>
<th>Configuration</th>
<th>Independent</th>
<th>Positive corr.</th>
<th>Negative corr.</th>
</tr>
</thead>
<tbody>
<tr>
<td>VKN Total samples</td>
<td>$3.528 \times 10^5$</td>
<td>$3.554 \times 10^5$</td>
<td>$3.469 \times 10^5$</td>
</tr>
<tr>
<td>M</td>
<td>$\pm 0.032 \times 10^5$</td>
<td>$\pm 0.027 \times 10^5$</td>
<td>$\pm 0.030 \times 10^5$</td>
</tr>
<tr>
<td>Makespan</td>
<td>$8.821 \times 10^6$</td>
<td>$8.884 \times 10^6$</td>
<td>$8.673 \times 10^6$</td>
</tr>
<tr>
<td>PCS</td>
<td>0.999</td>
<td>0.998</td>
<td>0.999</td>
</tr>
<tr>
<td>APS Total samples</td>
<td>$1.788 \times 10^5$</td>
<td>$1.788 \times 10^5$</td>
<td>$1.791 \times 10^5$</td>
</tr>
<tr>
<td>M</td>
<td>$\pm 0.013 \times 10^5$</td>
<td>$\pm 0.013 \times 10^5$</td>
<td>$\pm 0.013 \times 10^5$</td>
</tr>
<tr>
<td>Makespan</td>
<td>$4.470 \times 10^6$</td>
<td>$4.469 \times 10^6$</td>
<td>$4.478 \times 10^6$</td>
</tr>
<tr>
<td>PCS</td>
<td>0.986</td>
<td>0.987</td>
<td>0.984</td>
</tr>
</tbody>
</table>

For $m = 48$,

| VKN Total samples | $3.529 \times 10^5$ | $3.553 \times 10^5$ | $3.545 \times 10^5$ |
| M | $\pm 0.032 \times 10^5$ | $\pm 0.032 \times 10^5$ | $\pm 0.032 \times 10^5$ |
| Makespan | $7.358 \times 10^5$ | $7.401 \times 10^5$ | $7.385 \times 10^5$ |
| PCS | 1.000 | 0.999 | 0.996 |
| APS Total samples | $1.792 \times 10^5$ | $1.787 \times 10^5$ | $1.792 \times 10^5$ |
| M | $\pm 0.013 \times 10^5$ | $\pm 0.013 \times 10^5$ | $\pm 0.013 \times 10^5$ |
| Makespan | $3.733 \times 10^5$ | $3.722 \times 10^5$ | $3.733 \times 10^5$ |
| PCS | 0.981 | 0.988 | 0.986 |

For $m = 96$,

| VKN Total samples | $3.578 \times 10^5$ | $3.596 \times 10^5$ | $3.512 \times 10^5$ |
| M | $\pm 0.031 \times 10^5$ | $\pm 0.029 \times 10^5$ | $\pm 0.034 \times 10^5$ |
| Makespan | $3.732 \times 10^5$ | $3.760 \times 10^5$ | $3.671 \times 10^5$ |
| PCS | 0.998 | 0.998 | 0.997 |
| APS Total samples | $1.792 \times 10^5$ | $1.792 \times 10^5$ | $1.786 \times 10^5$ |
| M | $\pm 0.013 \times 10^5$ | $\pm 0.013 \times 10^5$ | $\pm 0.013 \times 10^5$ |
| Makespan | $1.867 \times 10^5$ | $1.867 \times 10^5$ | $1.861 \times 10^5$ |
| PCS | 0.982 | 0.984 | 0.978 |
Figure 6. Total sample size vs. number of systems when \( m = 48 \).

![Graph showing total sample size vs. number of systems when \( m = 48 \).](image)

By solving the balance equations for the underlying Markov chain from Buzacott and Shanthikumar (1993), we obtain that the optimal solutions are \((6, 7, 7, 12, 8)\) and \((7, 7, 6, 8, 12)\) (denoted as best alternatives) with steady-state throughput 5.776. We set the IZ parameter as \( SL_\delta = 0.01 \) and define the feasible solutions with steady-state throughput within \( SL_\delta \) from the best as good alternatives. The event of selecting one from either the best or the good alternatives is defined as a “correct selection.” Table 2 provides the information for all best and good alternatives.

48 processors and 64 GB memory and running CentOS 6.2, a Linux-based operating system. To understand how the number of processors (i.e., slaves) affects the performances of the APS procedure, we test this problem with different numbers of processors, \( m = 1, 4, 8, 16, 32, 48 \). In these experiments, we set the first-stage sample size \( n_0 = 10 \) and the desired PCS as 0.95.

Figure 7 captures a snapshot of the status of the threads for the master/slave structure after starting the APS procedure with 4 slaves. From the figure, we observe that all slaves (denoted as slaves 0 to 3) are working in parallel to generate samples, whereas the two threads in the master, named “consume sample” thread (i.e., the compare thread) and “produce alt” thread (i.e., the to-do thread), are idling at that time since the elimination has not been conducted and the input sequence has already been prepared.

In Table 3 we report the average total sample size with 95% confidence interval, the average makespan with 95% confidence interval, and the estimated PCS, based on 100 macroreplications. We find that the total sample sizes are almost the same for various numbers of slaves and the APS procedure can always deliver a correct selection. However, we also notice that the makespan (i.e., total time to complete the procedure) seems not to reduce in proportion to the number of slaves. This is because the R&S procedure

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### Table 2. Best and good alternatives for the buffer allocation problem.

<table>
<thead>
<tr>
<th>Alternative</th>
<th>Throughput</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>((6, 7, 7, 12, 8))</td>
<td>5.776</td>
<td>Best</td>
</tr>
<tr>
<td>((7, 7, 6, 8, 12))</td>
<td>5.776</td>
<td>Best</td>
</tr>
<tr>
<td>((6, 7, 7, 13, 7))</td>
<td>5.772</td>
<td>Good</td>
</tr>
<tr>
<td>((7, 7, 6, 13))</td>
<td>5.772</td>
<td>Good</td>
</tr>
<tr>
<td>((6, 7, 7, 11, 9))</td>
<td>5.771</td>
<td>Good</td>
</tr>
<tr>
<td>((7, 7, 6, 9, 11))</td>
<td>5.771</td>
<td>Good</td>
</tr>
</tbody>
</table>

---

**Figure 7.** A screenshot of the master/slave with the number of processors \( m = 4 \).
is not completely parallel. To estimate what percentage of the procedure is executed in parallel, we fit the average makespan based on Amdahl’s law (Amdahl 1967), which states that the speedup of parallelism can be defined as \(1/(1 - P + P/m)\), where \(P\) the proportion of a program that can be made parallel, \(1 - P\) is the remaining proportion that cannot be parallelized, and \(m\) is the number of processors. Suppose the makespan, \(T\), can be modeled as follows,

\[
T = \beta \left(1 - P\right) + P \cdot \frac{1}{m} + \epsilon = c_0 + c_1 \frac{1}{m} + \epsilon
\]

where \(\beta\) is the constant coefficient and \(\epsilon\) is random noise. By a linear regression, we obtain that \(c_0 = 40.4\) and \(c_1 = 332.9\) (with \(R^2 \geq 0.994\)), which implies that \(P \approx 0.892\), indicating that 89.2% of the program can be made parallel, a very high compatibility. Notice that this result is what we expected because the vast majority of the tasks are independent simulation runs that can be easily parallelized, and it suggests that large-scale R&S problems may be effectively solved using a parallel computing environment when it is available.

6. Conclusions and Future Work

In this paper, we show that it is very attractive to solve large-scale R&S problems using parallel computing environments, which may reduce total computational time by an order of magnitude and greatly enlarge the set of R&S problems that are considered solvable. However, we also find that a direct implementation of sequential R&S procedures in a parallel computing environment may lead to unexpected statistical issues and affect the statistical validity and efficiency of procedures. In this paper, we design two different approaches to solve R&S problems in parallel computing environments.

To further improve the efficiency of the procedures, there are a few issues that are worth future investigation. First, we adopted a straightforward round-robin rule in the input sequence in this paper. However, this may not be necessary. Indeed, a higher level of efficiency may be achievable if we use more carefully chosen input sequences. Second, the current master/slave structure requires a large amount of communication between the master and slaves. When simulation experiments are computationally fast or there are a very large number of slaves, the master may become a bottleneck. More effective ways of handling the operations on the master are also worth studying.

### Supplemental Material

Supplemental material to this paper is available at http://dx.doi.org/10.1287/opre.2015.1413.

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### References


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#### Table 3. Summary of three-stage-buffer-allocation example with different \(m\)’s.

<table>
<thead>
<tr>
<th>Number of slaves</th>
<th>(m = 1)</th>
<th>(m = 4)</th>
<th>(m = 8)</th>
<th>(m = 16)</th>
<th>(m = 32)</th>
<th>(m = 48)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total samples (\times 10^3)</td>
<td>2.426</td>
<td>2.434</td>
<td>2.442</td>
<td>2.442</td>
<td>2.433</td>
<td>2.436</td>
</tr>
<tr>
<td>±0.004</td>
<td>±0.004</td>
<td>±0.004</td>
<td>±0.004</td>
<td>±0.003</td>
<td>±0.004</td>
<td></td>
</tr>
<tr>
<td>Makespan (minutes)</td>
<td>370.5</td>
<td>129.4</td>
<td>94.4</td>
<td>68.3</td>
<td>41.7</td>
<td>34.2</td>
</tr>
<tr>
<td>±1.9</td>
<td>±2.3</td>
<td>±3.1</td>
<td>±4.1</td>
<td>±3.2</td>
<td>±1.9</td>
<td></td>
</tr>
<tr>
<td>PCS</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>

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