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Single Observation Adaptive Search for Continuous Simulation Optimization

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Optimizing the performance of complex systems modeled by stochastic computer simulations is a challenging task, partly due to the lack of structural properties (e.g., convexity). This challenge is magnified by the presence of random error whereby an adaptive algorithm searching for better designs can at times mistakenly accept an inferior design. In contrast to performing multiple simulations at a design point to estimate the performance of the design, we propose a framework for adaptive search algorithms that executes a single simulation for each design point encountered. Here the estimation errors are reduced by averaging the performances from previously evaluated designs drawn from a shrinking ball around the current design point. We show under mild regularity conditions for continuous design spaces that the accumulated errors, although dependent, form a martingale process and hence, by the strong law of large numbers for martingales, the average errors converge to zero as the algorithm proceeds. This class of algorithms is shown to converge to a global optimum with probability one. By employing a shrinking ball approach with single observations, an adaptive search algorithm can simultaneously improve the estimates of performance while exploring new and potentially better design points. Numerical experiments offer empirical support for this paradigm of single observation simulation optimization.

Key words: Simulation optimization, adaptive search, martingale processes
1. Introduction

Stochastic optimization problems, where an objective function is noisy and must be estimated, are finding applications in diverse areas, spanning engineering, economics, computer science, business and biological science. Simulation optimization algorithms, that integrate search for the optimum with observations from a noisy objective function, have been proposed in both continuous and discrete domains (Fu 2015, Pasupathy and Ghosh 2013).

Striking a balance between exploration of new points and estimation of potentially good points is critical for computationally efficient algorithms. We present a class of adaptive search algorithms that performs exactly one simulation per design point, which we call single observation search algorithms (SOSA). This class of SOSA algorithms combines exploration with estimation by estimating the expectation of the objective function at a point with an average of observed values from previously visited nearby points. The nearby points are within a shrinking ball around the current point. The challenge is to ensure that the errors associated with the estimates of the expectation of the noisy objective function do not bias the adaptive search algorithm and potentially lead to mistakenly accepting inferior solutions. We prove convergence to a global optimum for this class of SOSA algorithms under some mild regularity conditions. Then any adaptive search algorithm that fits into this class can utilize single observations within shrinking balls in contrast to multiple repetitions at a point to successfully search for a global optimum.

The idea of simulating a single observation per design point was first used by Robbins and Monro for estimating gradients in their classic stochastic approximation algorithm (see, for instance, Chau and Fu (2015), Kushner and Yin (2003), Robbins and Monro (1951)). This class of stochastic approximation algorithms have proven to be very successful at optimizing noisy functions on continuous domains (see Chau and Fu (2015)). These classic stochastic approximation algorithms are based on steepest descent, and are shown to converge to a local optimum. We consider the question of whether there exist single observation per design point algorithms that converge to a global optimum.

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optimum. The insight of Robbins and Monro in their classic stochastic approximation algorithm with single observations, is “... if the step sizes in the parameter updates are allowed to go to zero in an appropriate way as (iterations go to infinity) then there is an implicit averaging that eliminates the effects of the noise in the long run.” (Kushner and Yin 2003, p. viii). The random error associated with the steepest descent type method can be shown to be a martingale difference. This martingale property can be used to prove the convergence to a local optimum of the class of the steepest descent type methods. This paper shows how to extend implicit averaging in a way that provides convergence to a global optimum for our class of SUSA algorithms on non-convex, multi-modal, noisy optimization problems.

A necessary characteristic of our class of SUSA algorithms is that the underlying adaptive algorithm converges to a global optimal value when the objective function is not noisy. There are many such adaptive search algorithms for global optimization, for example, simulated annealing, evolutionary algorithms, model based algorithms (Hu et al. 2007, 2014), nested partition method (Shi and Ólafsson 2000a), interacting particle algorithms (Molvašoglu et al. 2009, 2010, Mete and Zabinsky 2014). (See Zabinsky (2011) for an overview.) We characterize algorithms in SUSA by their sampling distributions for generating sequential points. Since many adaptive search algorithms for global optimization maintain a sampling distribution that is bounded away from zero on the feasible region, we prove that this condition is sufficient to satisfy the assumptions of the convergence analysis and consequently, such an algorithm converges to a true global optimum using single observations with the shrinking ball approach.

The challenge in proving convergence to a global optimum with the additional demand of estimation is due to the complex dependencies among the observational errors introduced by an adaptive algorithm. An adaptive algorithm is influenced by past errors as it seeks new candidates, and thus may be biased toward design points that were observed to be better than their true value. We show in this paper that underlying martingale properties exhibited by these adaptive algorithms can nonetheless be used to prove convergence to a global optimum.
Baumert and Smith (2002) introduced the idea of estimating the objective function at a specific design point $x$ by including other points within a shrinking ball around $x$, thus never repeating a simulation at a single design point. Their algorithm was based on pure random search, which generates points independently and thus is not adaptive thereby avoiding dependencies among subsequent errors.

Andradóttir and Prudius (2010) investigated three methods, namely Adaptive Search with Resampling (ASR), deterministic shrinking ball and stochastic shrinking ball. They proved that the three algorithms are strongly convergent to a global optimum. But again, the deterministic and the stochastic shrinking ball approaches are based on pure random search with independent sampling.

Closely related works in the area of simulation optimization include: Stochastic Approximation (Borkar 2008, 2013, Spall 2003), Sample Average Approximation (SAA)(Kim et al. 2015, Kleywegt et al. 2002), Gradient-based Adaptive Stochastic Search for Simulation Optimization (Zhou et al. 2014), Nested Partitions Method (Shi and Ólafsson 2000b) and Low-Dispersion Point Sets (Yakowitz et al. 2000). These works either employ a gradient based approach, exploit special structure of the feasible region or use multiple observations of sampled design points. Stochastic approximation (Borkar 2008, 2013, Chau and Fu 2015, Kim et al. 2015, Kushner and Yin 2003, Robbins and Monro 1951, Spall 2003), starting with the seminal work of Robbins and Monro (Robbins and Monro 1951), is widely used for continuous problems with many applications. It is a first-order method with single or multiple observations per point that typically converges to a first-order stationary point. In contrast, SOSA does not require gradient information, allows a general continuous feasible region and uses a single observation per sampled design point. Sample average approximation (Homem-De-Mello 2003, Kim et al. 2015, Kleywegt et al. 2002) has been used in both continuous and discrete stochastic optimization. SAA takes a collection of draws of the random vector $U$ (e.g., $u_1, \ldots, u_N$) that may be independent identically distributed draws or possibly quasi-Monte Carlo samples or Latin hypercube designs, and then solves an associated
problem, $\min_{x \in S} \hat{f}(x)$ where $\hat{f}(x) = (1/N) \sum_{i=1}^{N} g(x, u_i)$. Under certain conditions, SAA converges to the global optimum as $N \to \infty$. Retrospective approximation (Pasupathy 2010) and variable sample size methods (Homem-De-Mello 2003) are efficient variations of SAA. SAA is also related to scenario-based stochastic programming (Shapiro et al. 2009), where the random draws are performed before the optimization over $x$ is performed. Other simulation optimization methods (ours included) consider each $x$ and then draw from $U$. Both methods have complex correlations and dependencies.

Meta-models (Barton and Meckesheimer 2006, Pedrielli and Ng 2015) are also commonly used in simulation optimization. SOSA, in principle, can be viewed as a meta-model and resembles Kriging (Pedrielli and Ng 2015) in that the objective function at non-sampled design points are estimated based upon weighted observations of the function at other sampled design points. However, Kriging assumes conditions that are different from ours.

In this paper, we consider a broad class of adaptive random search algorithms in a continuous domain, and prove that the accumulated error of the search process looking forward from the current candidate point follows a martingale process. Since the errors are zero in expectation for each point, the average error converges to zero from the strong law of large numbers for martingales (see “the impossibility of systems” in Feller (1971)). This allows us to prove convergence to a global optimum in probability for a broad class of adaptive random search algorithms with mild assumptions and using a single observation per design point.

We provide numerical results using two algorithms; both algorithms are run with single observations (SOSA), as well as with multiple replications implemented with ASR as in Andradóttir and Prudius (2010). The two algorithms are: a) a sampler based on a modified version of Improving Hit-and-Run (IHR) (see Zabinsky et al. (1993), Zabinsky and Smith (2013)); and b) a uniform local/global sampler originally used in Andradóttir and Prudius (2010). Andradóttir and Prudius (2010) demonstrated that ASR performed better computationally than non-adaptive shrinking ball approaches in high dimensions. In this paper, we demonstrate on two test problems in ten dimensions that SOSA performs better computationally than ASR for both sampling methods tested.
This is an encouraging result, that implicit averaging over shrinking balls can be very effective and can actually improve computational performance of stochastic searches in general. The averaging effect of nearby points seems to have the effect of discovering local and global trends in the objective function, thus enabling an adaptive algorithm to sample effectively.

2. Single Observation Simulation Optimization

The stochastic optimization problem we consider is

\[
\min_{x \in S} f(x) \quad (1)
\]

where \(x \in S \subset \mathbb{R}^d\) and

\[
f(x) = \mathbb{E}[g(x,U)]. \quad (2)
\]

However, the objective function \(f(x)\) cannot be evaluated exactly. Instead, we have a noisy evaluation available, that is, the performance at a design point \(x \in S \subset \mathbb{R}^d\) is given by \(g : S \times \Omega \rightarrow \mathbb{R}\), where \(U\) is a random element over a probability space denoted \((\Omega, \mathcal{A}, \mathbb{P})\). In discrete-event simulation, randomness is typically generated from a sequence of pseudo-random numbers, therefore, we let \(U\) represent an independent and identically distributed (i.i.d.) sequence of uniform random variables. We assume that \(f\) is continuous and \(S\) is compact so that a minimum exists. Let \(X^* = \text{Arg min}_{x \in S} f(x)\) denote the set of optimal solutions, and let \(f^*\) be the optimal value.

A common approach in simulation optimization is to estimate \(f(x)\) by observing the output of a simulation run, \(g(x,u)\), where \(u\) is a realization of the random variable \(U\). The difference between the observed performance and mean performance, denoted

\[
Z(x) = g(x,U) - f(x) \quad (3)
\]

represents the random observational error.

When the random observational errors are independent across all iterations of the algorithm, and identically distributed, then the strong law of large numbers can be invoked to prove the error goes to zero as iterations increase to infinity. The challenge with establishing convergence to optimality
for an adaptive algorithm for global optimization is that the random errors are in general neither identically distributed nor independent. An adaptive algorithm that favors “better” design points introduces complex dependencies among the errors. Since points that appear “better” influence the adaptive algorithm, the optimal value estimates tend to be negatively biased.

Illustration of Dependent Estimation Errors

Suppose $S$ is the union of two non-overlapping balls we will call ball $L$ and ball $R$. Suppose moreover that the objective function values $f(x)$, for $x \in L$, are better (less) than those in $R$. Suppose in the initial step of an adaptive algorithm we begin by sampling a point from ball $L$ and we observe its objective function value. Next we sample a point from the other ball $R$ and compare its value with that of our point in $L$. The third point will be sampled from the ball with the smaller observed value.

Suppose that the error associated with the first noisy observation in $L$ is negative, i.e., the observed value is smaller than the true $f(x)$. Now suppose the third point is in $R$. In this case, a negative error at the first point sampled means the error at the second point must also be negative since its expected value is inferior to the first point. This example illustrates that there is a dependency between the errors from the first and the second observations. In general, there are subtle dependencies that can be induced by adaptive search algorithms.

In Section 3, we show that, while errors looking backward from the current iteration point are dependent (e.g., looking at the first and second points, having sampled the third), errors looking forward when conditioning on the identity of the current iteration point (e.g., looking at the fourth point, having sampled the third) are independent of past errors.

In this paper, we rigorously analyze the accumulated error associated with a class of adaptive random search algorithms. As suggested in the example above, while the accumulated error of the entire process does not form a martingale, the accumulated error of the process after a point has been evaluated does form a martingale. This insight allows us to prove convergence to a global optimum with probability one for a broad class of adaptive random search algorithms with mild assumptions and using a single observation per design point. The key is to “slow down” the
estimation process so that it does not cause the optimization search process to converge prematurely to a wrong solution.

We make the following three assumptions regarding the problem. These assumptions are relatively mild, and are typically satisfied in most continuous simulation optimization problems.

**Assumption 1:** The feasible set $S \subset \mathbb{R}^d$ is a closed and bounded convex set with non-empty interior.

**Assumption 2:** The objective function $f(x)$ is continuous on $S$.

We consider two versions of the next assumption, Assumption 3 and Assumption 3'. Assumption 3 requires that the random error be bounded, and is satisfied by many bounded distributions. However, it does not include distributions having infinite support, such as Normal or Gamma distributions. Assumption 3' requires that the random error has bounded variance, and the Normal and Gamma distributions with finite variance satisfy Assumption 3'. In the next section, we see that Assumption 3 leads to a stronger convergence result (convergence with probability one) than Assumption 3' (convergence in probability).

**Assumption 3:** The random error $(g(x,U) - f(x))$ is uniformly bounded over $x \in S$, that is, there exists $0 < \alpha < \infty$ such that, for all $x \in S$, with probability one,

$$|g(x,U) - f(x)| < \alpha.$$

**Assumption 3':** The random error $(g(x,U) - f(x))$ has bounded variance over $x \in S$.

### 3. A Class of Adaptive Random Search Algorithms and Convergence Analysis

We propose a class of adaptive stochastic search algorithms with single observations per design for the continuous simulation optimization problem in (1). We show that, under some regularity conditions, an algorithm in this class will generate a sequence of objective function estimates that converge to the true global optimum with probability one.

In the course of the algorithm, design points are sequentially sampled from the design space $S$, according to an adaptive sampling distribution, denoted by $q_n$ at iteration $n$. The objective function
at a point is estimated using the observed function values of nearby sample points within a certain radius. The use of nearby points allows for a strong law of large numbers type of convergence by asymptotically eliminating the random error associated with the sequence of observations. The radius shrinks as the algorithm progresses, hence the image of shrinking balls. This shrinking process asymptotically eliminates the systematic bias of using estimates of neighboring points since the objective function $f(x)$ is continuous in $x$.

For each $x \in S$, let $B(x, r)$ be the ball centered at $x$ with radius $r$. Let $X_n$ and $Y_n$ be, respectively, the set of sample points obtained in the course of the Single Observation Search Algorithms and their corresponding function evaluations up to iteration $n$. For $x_i \in X_n$, the objective function estimate $\hat{f}_n(x_i)$ of $x_i$ comes from the average of the function evaluations of the sample points that fall into the balls centered at $x_i$. Let $l_n(x_i)$ denote the number of sample points that fall into the balls centered at $x_i$, called contributions to the estimate of $x_i$.

Consider the following class of algorithms using the shrinking ball concept. Note that, to guarantee convergence to a true global optimum, regularity conditions for the sampling density $q_n$ and the parameter sequences $r_n$ and $i_n$ are required.

**Single Observation Search Algorithms (SOSA)**

We are given:

- A continuous initial sampling density for search on $S$, $q_1(x)$, and a family of continuous adaptive search sampling distributions on $S$ with density

  $$q_n(x \mid x_1, y_1, \ldots, x_{n-1}, y_{n-1}), \ n = 2, 3, \ldots,$$

  where $x_n$ is the sample point at iteration $n$ and $y_n$ is its observed function value.

- A sequence of radii $r_n > 0$.

- A sequence $i_n < n$.

**Step 0:** Sample $x_1$ from $q_1$, observe $y_1 = g(x_1, u_1)$ from the simulation, where $u_1$ is a sample value having the same distribution as $U$ and independent of $x_1$. Set $X_1 = \{x_1\}$ and $Y_1 = \{y_1\}$. Also, set $\hat{f}_1(x_1) = \hat{f}_1^*(x_1) = y_1$, $l_1(x_1) = 1$ and $x_1^* = x_1$. Set $n = 2$. 
Step 1: Given $x_1, y_1, \ldots, x_{n-1}, y_{n-1}$, sample the next point, $x_n$, from $q_n$. Independent of $x_1, y_1, \ldots, x_{n-1}, y_{n-1}, x_n$, obtain a sample value $u_n$ having the same distribution as $U$ and evaluate the objective function value $y_n = g(x_n, u_n)$.

Step 2: Update $X_n = X_{n-1} \cup \{x_n\}$ and $Y_n = Y_{n-1} \cup \{y_n\}$. For each $x \in X_n$, update the contribution and the estimate of the objective function value as

$$l_n(x) = |\{k \leq n: x_k \in B(x, r_k)\}| = \begin{cases} l_{n-1}(x) & \text{if } x_n \notin B(x, r_n) \\ l_{n-1}(x) + 1 & \text{if } x_n \in B(x, r_n) \end{cases}$$

(4)

and

$$\hat{f}_n(x) = \frac{\sum_{\{k \leq n: x_k \in B(x, r_k)\}} y_k}{|\{k \leq n: x_k \in B(x, r_k)\}|} = \begin{cases} \hat{f}_{n-1}(x), & \text{if } x_n \notin B(x, r_n) \\ ((l_n(x) - 1)\hat{f}_{n-1}(x) + y_n)/l_n(x), & \text{if } x_n \in B(x, r_n) \end{cases}$$

(5)

where $B(x, r_k)$ is a ball of radius $r_k$ centered at $x$ and $|A|$ for a set $A$ denotes the number of elements in $A$. Estimate the optimal value as

$$\hat{f}_n^* = \min_{x \in X_n} \hat{f}_n(x)$$

(6)

and estimate the optimal solution as

$$x_n^* \in \{x \in X_n : \hat{f}_n(x) = \hat{f}_n^*\}$$

(7)

where $X_n$ is a subset of $X_n$ determined by the index $i_n$.

Step 3: If a stopping criterion is met, stop. Otherwise, update $n \leftarrow n + 1$ and go to Step 1.

Observe that the objective function estimate $\hat{f}_n(x)$ is defined for all $x \in S$. However, in the course of the algorithm, we only compute the estimate for $x \in X_n$. To do this, the simplest way is to go through $X_n$ once in each iteration, and update the objective function estimates through the recursive formula in (5). By doing so, up to iteration $n$, the estimates at sample points $x_1, x_2, \ldots, x_n$ will be updated $n, n-1, \ldots, 1$ times, respectively, which will result in at most $n(n+1)/2$ updates. At each iteration, the function estimate of a sample point will get updated only when the new sample point falls relatively close to a previously sampled point (within its ball of radius $r_k$). Therefore, up to iteration $n$, the function estimates will be calculated less than $n(n+1)/2$ times.
Notice that the algorithm takes the estimate of the optimal value on the $n^{th}$ iteration, $\hat{f}_n^*(x)$, not from all $n$ function estimates, but from a subset of the function estimates up to $i_n$. By slowing the sequence of estimates using $i_n$, we are able to ensure convergence of the optimal value estimate $\hat{f}_n^*(x)$ to the true optimal value $f^*$. The idea is that the shrinking balls must shrink slowly enough to still allow for the number of points in the balls to grow to infinity.

The main result of the paper is stated in Theorem 3, where we prove that an adaptive random search algorithm with single observations per design, using a shrinking ball approach, converges to a global optimum with probability one. The following corollary with the relaxed Assumption 3' proves convergence to a global optimal value in probability. The convergence analyses rest on the martingale property of the random error, which we establish in Theorem 1.

To develop this martingale property, we investigate the random error at a design vector, as in (3), $Z(x) = g(x,U) - f(x)$, and since

$$\mathbb{E}[Z(x)] = \mathbb{E}[g(x,U)] - f(x) = f(x) - f(x) = 0, \text{ for all } x \in S,$$

(8)

$Z(x)$ is a random error with zero expectation. It is worth noting that, rewriting (3) as

$$g(x,U) = f(x) + Z(x)$$

(9)

now states that, given a design point $x$, a random performance can always be decomposed into a sum of its expected performance and a random error with zero expectation.

To establish the convergence result for SOSA, it is necessary that the sequence of random numbers used as input to the simulation are independent of the past information. To state this more formally, first let $X_n$ and $Y_n$ denote the sample point and its corresponding objective function evaluation at iteration $n$, for $n = 1, 2, \ldots$. Then

$$Y_n = g(X_n, U_n),$$

(10)

where $\{U_n, n = 1, 2, \ldots\}$ are random elements, i.i.d. and have the same distribution as $U$. Since $X_n$, $U_n$ and $Y_n$ are generated sequentially, we can construct a filtration, starting with $\mathcal{F}_0 = \sigma(X_1)$, the $\sigma$-field generated by $X_1$, and, for $n = 1, 2, \ldots$, $\mathcal{F}_n = \sigma(X_1, U_1, \ldots, X_n, U_n, X_{n+1})$, the $\sigma$-field generated
by $X_1, U_1, \ldots, X_n, U_n, X_{n+1}$. Observe that $X_n$ is $\mathcal{F}_{n-1}$ measurable. Since $Y_n$ is a function of $X_n$ and $U_n$, $Y_n$ is $\mathcal{F}_n$ measurable. The process of $(X_n, Y_n)$ is then adapted to the filtration $\{\mathcal{F}_n\}_{n=0}^{\infty}$. It is crucial to the convergence results that $U_n$ is generated so that it is independent of $\mathcal{F}_{n-1}$.

We next establish that the expected random error conditioned on the filtration is zero. This property will induce a martingale process of accumulated errors and, finally, enable the optimal value estimates generated by the algorithm, $\{\hat{f}_n^*\}$, to converge to the true optimal value $f^*$.

Define the random error at iteration $n$ by $Z_n$ where

$$Z_n = Y_n - f(X_n).$$

Note that $Z_n$, as well as $Y_n$, depend on $X_n$ and $U_n$, but we suppress the arguments in the notation to simplify the presentation.

We now establish a crucial martingale property, that $\mathbb{E}[Z_n \mid \mathcal{F}_{n-1}] = 0$. Since $X_n$ is $\mathcal{F}_{n-1}$ measurable and $U_n$ is independent of $\mathcal{F}_{n-1}$,

$$\mathbb{E}[Y_n \mid \mathcal{F}_{n-1}] = \mathbb{E}[g(X_n, U_n) \mid \mathcal{F}_{n-1}] = \mathbb{E}[g(X_n, U_n) \mid X_n]$$

and since $U_n$ is identically distributed as $U$,

$$= \mathbb{E}[g(X_n, U) \mid X_n] = f(X_n).$$

(12)

Again, since $X_n$ is $\mathcal{F}_{n-1}$ measurable,

$$\mathbb{E}[Z_n \mid \mathcal{F}_{n-1}] = \mathbb{E}[Y_n - f(X_n) \mid \mathcal{F}_{n-1}] = \mathbb{E}[Y_n \mid \mathcal{F}_{n-1}] - f(X_n) = 0.$$

(13)

The result in (13) provides the basis of the martingale property, since the error on iteration $n$, conditioned on the past information in $\mathcal{F}_{n-1}$, is zero.

Furthermore,

$$\mathbb{E}[Z_n] = \mathbb{E}[\mathbb{E}[Z_n \mid \mathcal{F}_{n-1}]] = \mathbb{E}[0] = 0,$$

(14)

which establishes that, according to (12) and (14), we can decompose $Y_n$ into its conditional expectation term and its error term,

$$Y_n = f(X_n) + Z_n.$$
where the random error $Z_n$ has zero expectation.

Now we express the accumulated error in estimating $f(X_i)$ associated with the sample point $X_i$ in terms of the error coming from the points in the balls around $X_i$.

At iteration $n$, and for a fixed sample point $X_i$, for $i \leq n$, let $M_n(X_i)$ be the accumulated error in estimating $f(X_i)$ using the function evaluations from the points $X_k$, $k = 1, \ldots, n$ that fall into balls around $X_i$. We define an indicator function to identify the sample points in balls around $X_i$,

$$I_k(X_i) = \begin{cases} 
1 & \text{if } X_k \in B(X_i, r_k) \\
0 & \text{if } X_k \notin B(X_i, r_k) 
\end{cases}$$

for $k = 1, \ldots, n$. Using the indicator function, we have

$$M_n(X_i) = \sum_{k=1}^{n} I_k(X_i) Z_k. \quad (16)$$

It is important to note that $\{M_n(X_i), n = 1, 2, \ldots\}$ for $i > 1$ is not a martingale, due to the dependencies on early sample points in the sequence. In fact, when $n < i$, $M_n(X_i)$ depends on unrealized information of $X_i$. To be exact, $M_n(X_i)$, where $n < i$, is not $F_n$ measurable.

At each iteration, the best candidate $X_n^*$ is chosen as the point whose function estimate is the smallest so far. Since the function estimate is formed by the function observations of previous points in the ball around $X_n^*$, these objective function observations tend to be negatively biased. The errors from these points tend to be concurrently negative and, hence, are correlated.

We have discovered that we can decompose the accumulated error into two parts: the error from the sample points that preceded $X_i$, and the error from the sample points that were sampled after $X_i$, and establish the martingale property for the second part of the error. We let

$$M_i^n(X_i) = \sum_{k=i}^{n} I_k(X_i) Z_k, \quad n = i, i+1, \ldots. \quad (17)$$

be the accumulated error from function evaluations taken from iteration $i$ onward to iteration $n$. Note that $M_i^n(X_i)$ is the sum of $(n-i+1)$ error terms. Define $M_i^{i-1}(X_i) = 0$. Now, using (16) and (17), we decompose $M_n(X_i)$ into two parts

$$M_n(X_i) = \sum_{k=0}^{i-1} I_k(X_i) Z_k + M_i^n(X_i), \quad (18)$$
where \( I_0(X_i)Z_0 = 0 \) as a convention.

In Theorem 1, we show that the accumulated error about a sample point \( X_i \) from function evaluations taken from iteration \( i \) onward to iteration \( n \) is a martingale.

**Theorem 1.** For any \( i, i = 1, 2, \ldots \), \( \{ M^i_n(X_i), n = i, i + 1, \ldots \} \) is a martingale with respect to the filtration \( \{ F_n, n = i, i + 1, \ldots \} \).

**Proof:** Fix \( i \). Define

\[
\tilde{M}^i_n = \sum_{k=i}^{n} Z_k
\]

as the accumulated error from \textit{all} points sampled on the iterations from iteration \( i \) through iteration \( n \). We first show that \( \{ \tilde{M}^i_n, n = i, i + 1, \ldots \} \) is a martingale with respect to the filtration \( \{ F_n, n = i, i + 1, \ldots \} \). This is equivalent to showing that \( \mathbb{E}[|\tilde{M}^i_n|] < \infty \), and \( \mathbb{E}[\tilde{M}^i_n | F_{n-1}] = \tilde{M}^i_{n-1} \), for all \( n \geq i \). By Assumption 3, \( \mathbb{E}[|Z_n|] < \alpha < \infty \). By the triangular inequality of the absolute value function, \( \mathbb{E}[|\tilde{M}^i_n|] \leq (n - i + 1)\alpha < \infty \). In addition,

\[
\mathbb{E}[\tilde{M}^i_n | F_{n-1}] = \mathbb{E}[Z_n + \tilde{M}^i_{n-1} | F_{n-1}]
\]

\[
= \mathbb{E}[Z_n | F_{n-1}] + \mathbb{E}[\tilde{M}^i_{n-1} | F_{n-1}]
\]

\[
= \tilde{M}^i_{n-1}.
\]

The last equation follows by (13) and that \( \tilde{M}^i_{n-1} \) is \( F_{n-1} \)-measurable, for all \( n \geq i \). Therefore, \( \{ \tilde{M}^i_n, n = i, i + 1, \ldots \} \) is a martingale.

Observe that \( I_k(X_i) \) is \( F_{k-1} \)-measurable, for \( k = i, i + 1, \ldots \). Therefore, \( I_k(X_i) \) is a decision function with respect to the filtration \( \{ F_n, n = i, i + 1, \ldots \} \). Now, for \( n = i, i + 1, \ldots, \)

\[
M^i_n(X_i) = \sum_{k=i}^{n} I_k(X_i)Z_k = M^i_{n-1}(X_i) + I_n(X_i)(\tilde{M}^i_n - \tilde{M}^i_{n-1} - I_n(X_i)).
\]

Therefore, by the impossibility of systems (Feller 1971, pg 213), \( \{ M^i_n(X_i), n = i, i + 1, \ldots \} \) is a martingale. \( \square \)
We now express the estimate of the function value at a sample point and the estimate of the optimal value in terms of \( X_i, f(X_i) \) and \( M_n(X_i) \). For a fixed \( i \), let \( L_n(X_i) \) be the number of sample points that fall into the balls \( B(X_i, r_k) \) around \( X_i \) where \( k = 1, \ldots, n \) and \( n \geq i \), that is,

\[
L_n(X_i) = \sum_{k=1}^{n} I_k(X_i).
\]

The estimate of the function value at a sample point \( X_i \) can be expressed as

\[
\hat{f}_n(X_i) = \frac{\sum_{k=1}^{n} I_k(X_i) Y_k}{L_n(X_i)} = \frac{\sum_{k=1}^{n} I_k(X_i) f(X_k)}{L_n(X_i)} + \frac{M_n(X_i)}{L_n(X_i)}
\]

where the first term includes the systematic bias and the second term is the accumulated error. The systematic bias is created by the fact that an estimate of the function value at a point includes function value estimates of points around it with different expectations. The accumulated error \( M_n(X_i) \) can be decomposed (see (18)) into a non-martingale accumulated error term, \( \sum_{k=0}^{i-1} I_k(X_i) Z_k \), and a martingale process of the accumulated errors, \( M'_n(X_i) \), as a result of Theorem 1.

The estimate of the optimal value \( \hat{f}^* \), is

\[
\hat{f}^*_n = \min_{i=1, \ldots, i_n} \{ \hat{f}_n(X_i) \} = \min_{i=1, \ldots, i_n} \left\{ \frac{\sum_{k=1}^{n} I_k(X_i) f(X_k)}{L_n(X_i)} + \frac{M_n(X_i)}{L_n(X_i)} \right\}.
\]

Note that \( \hat{f}^*_n \) is the minimum taken not from all \( n \) function estimates but from a subset of the function estimates up to \( i_n \), where \( i_n \leq n \). The size of the subset \( i_n \) is a control parameter required to ensure the convergence of the optimal value estimate \( \hat{f}^* \) to the true optimal value \( f^* \), by slowing the estimation.

Since \( \hat{f}^*_n \), the estimate of the optimal value generated by SOSA, is taken from the minimum of a growing number of estimated function values, in order to prove that \( \hat{f}^* \) in fact converges to \( f^* \), the true optimal value, we require some form of uniformity in the convergence of these estimates. To establish the requirements, we add one more assumption.

Recall that \( L_n(x) \) is the number of sample points that fall in the balls around \( x \). Given a function of natural numbers \( \tilde{L}(n) \), we define \( D(n) \) to be the event that each design vector \( x \) has at least \( \tilde{L}(n) \) sample points in the balls around \( x \), that is,

\[
D(n) = \bigcap_{x \in S} \left\{ L_n(x) \geq \tilde{L}(n) \right\}.
\]
The objective function evaluations of these sample points (one function evaluation per sample point) around a design point will form the estimate of the objective function of that design point.

The key idea is that the number of sample points in the balls around \( x \) grows at least as fast as \( \tilde{L}(n) \) even though the radii of the balls are shrinking. In other words, the balls cannot shrink too quickly; the shrinking balls must maintain a threshold of sample points in them.

**Definition 1.** A function \( h(n) \) is called \( O(n^p) \) where \( p \in \mathbb{R} \) if there is a \( 0 < \kappa_r < \infty \) such that for all \( n \in \mathbb{N}, 0 \leq h(n) \leq \kappa_r n^p \). A function \( h(n) \) is called \( \Omega(n^p) \) where \( p \in \mathbb{R} \) if there is a \( 0 < \kappa_L < \infty \) such that for all \( n \in \mathbb{N}, h(n) \geq \kappa_L n^p \). A function \( h(n) \) is called \( \Theta(n^p) \) if it is both \( O(n^p) \) and \( \Omega(n^p) \).

**Assumption 4:** Assume there exists \( 1/2 < \gamma < 1 \) and a function \( \tilde{L}(n) \) that is \( \Omega(n^\gamma) \), such that

\[ \sum_{n=1}^{\infty} P(D(n)^c) < \infty \]

where \( D(n)^c \) is the complement of event \( D(n) \), and \( \gamma \) is called an order of local sample density.

Assumption 4 ensures that there are on the order of \( n^\gamma \) function evaluations used in the estimate of every point in the design space.

The single observation search algorithms can satisfy Assumption 4 if the sampling densities satisfy some regularity conditions, and if the radii of the balls do not shrink too quickly. In particular, if the search sampling density \( q_n, n = 1, 2, \ldots \) is uniformly bounded away from zero on \( S \) and \( r_n \) is of \( \Omega(n^{-(1-\gamma)/d}) \), then Assumption 4 is satisfied. Many adaptive search algorithms for global optimization do have their search sampling density bounded away from zero, and in the next section we consider two such algorithms and prove that they satisfy Assumption 4, and hence converge to a global optimum.

In leading up to Theorem 3, we expand the estimate of the function value in (19) as,

\[
\hat{f}_n(X_i) = \frac{\sum_{k=1}^{n} I_k(X_i) f(X_i)}{L_n(X_i)} + \frac{\sum_{k=1}^{n} I_k(X_i) (f(X_k) - f(X_i))}{L_n(X_i)} + \frac{\sum_{k=1}^{n-1} I_k(X_i) Z_k}{L_n(X_i)} + \frac{\sum_{k=n}^{\infty} I_k(X_i) Z_k}{L_n(X_i)}
\]

\[
= f(X_i) + \left( \frac{\sum_{k=1}^{n} I_k(X_i) f(X_k)}{L_n(X_i)} - f(X_i) \right) + \frac{\sum_{k=1}^{n-1} I_k(X_i) Z_k}{L_n(X_i)} + \frac{\sum_{k=n}^{\infty} I_k(X_i) Z_k}{L_n(X_i)} \tag{21}
\]
to clearly identify the correct value in the first term, the bias due to nearby points in the second term, the non-martingale accumulated error in the third term and the martingale accumulated error in the fourth term. The second term, the bias, is created by the fact that an estimate of the function value at a point is formed by the function value of that point and other points around it but within the balls. Based on the continuity of the objective function in Assumption 2, we employ Cesaro’s Lemma (see Lemma EC.1 in the e-companion) with the shrinking ball mechanism to show that the bias term is washed away by averaging. The third term, the non-martingale random error of the estimated objective function value of a specific sample point, is formed by the errors corresponding to other points that are sampled prior to that sample point. These non-martingale random errors can be highly correlated and may not cancel each other out by averaging alone. However, it is considered a fixed term once a specific point has been sampled. Therefore, the slowing sequence, \( i_n \), is employed to slow the growth of this term, causing this non-martingale random error to diminish to zero when divided by the number of points in the associated balls. The fourth term, the martingale random error of the estimated objective function value of a specific sample point, is formed by the errors corresponding to points that are sampled from that specific sample point onward. The slowing sequence together with the martingale property through the Azuma-Hoeffding Inequality (see Lemma EC.2 in the e-companion) cause the martingale random error to disappear. Thus, in the limit, we are left with the correct value for the estimate of the function value.

We next prove Theorem 2 showing that the probability the estimate is incorrect by \( \varepsilon \) amount for the early portion of the estimates goes to zero as \( n \) goes to infinity, i.e.,

\[
\lim_{n \to \infty} P \left( \bigcup_{i=1}^{i_n} \left\{ |\hat{f}_n(X_i) - f(X_i)| \geq \varepsilon \right\} \right) = 0.
\]

For notational convenience, define \( A(n, \varepsilon) \) as the event that, when we consider only the early portion of the sequence up to \( i_n \), at least one objective function estimate is incorrect by more than the target error \( \varepsilon \) allowed, for \( \varepsilon > 0 \), that is,

\[
A(n, \varepsilon) = \bigcup_{i=1}^{i_n} \left\{ |\hat{f}_n(X_i) - f(X_i)| \geq \varepsilon \right\}.
\]
Theorem 2 gives conditions under which, the probability of missing this error target for the early portion of the estimates goes to zero as \( n \) goes to infinity. Theorem 2 makes use of the martingale property established in Theorem 1 and the slowing sequence \( i_n \) in Sosa.

**Theorem 2.** If Assumptions 1, 2, 3 and 4 are satisfied, and if \( i_n \uparrow \infty \) such that \( i_n \leq ns \) where \( 0 < s < \gamma \), then, for all \( \varepsilon > 0 \),
\[
\sum_{n=1}^{\infty} P(A(n, \varepsilon)) < \infty.
\]

*Proof:* See the e-companion. \( \square \)

By Assumption 4, the algorithm generates sample points that fill the feasible region. Once the estimated objective function errors of all sample points are controlled as described in Theorem 2 and the objective function is continuous according to Assumption 2, the optimal value estimates converge to the true optimal value. This convergence property is formalized in Theorem 3.

**Theorem 3.** If Assumptions 1, 2, 3 and 4 are satisfied, and if \( i_n \uparrow \infty \) such that \( i_n \leq ns \) where \( 0 < s < \gamma \), then \( \hat{f}_n^* \to f^* \) with probability one.

*Proof:* See the e-companion. \( \square \)

If Assumption 3 is relaxed to Assumption 3', we have a weaker convergence in probability result.

**Corollary 1.** If Assumptions 1, 2, 3' and 4 are satisfied, and if \( i_n \uparrow \infty \) such that \( i_n \leq ns \) where \( 0 < s < \gamma \), then, for all \( \varepsilon > 0 \),
\[
\lim_{n \to \infty} P(|\hat{f}_n^* - f^*| \geq \varepsilon) = 0,
\]
i.e., \( \hat{f}_n^* \to f^* \) in probability.

*Proof:* See the e-companion. \( \square \)

Note that Assumption 4 can also be relaxed, but with a weaker convergence result as in Corollary 1. A weaker Assumption 4, requiring only that \( \lim_{n \to \infty} P(D(n)^c) = 0 \), produces the same effect through the last term of (EC.4), and thus, as in Corollary 1, \( \hat{f}_n^* \) converges to \( f^* \) only in probability.

Now, in Corollary 2, we show that not only does Sosa converge to the optimal value, but, under appropriate conditions, it converges to an optimal solution. Let \( X_n^* \) represent the optimal solution
estimate at iteration $n$. In the case of multiple optima, it is possible that the estimates jump within the set of optima depending on the underlying sampling distributions $q_n$, however, the distance between the optimal solution estimate $X^*_n$ and the set of global optima $\mathcal{X}^*$ converges to zero with probability one. For a solution $x \in \mathbb{R}^d$ and a subset $E \subset \mathbb{R}^d$, we define the distance from $x$ to $E$ as $\rho(x, E) = \inf_{y \in E} \|x - y\|$ where $\|\cdot\|$ denotes the Euclidean norm on $\mathbb{R}^d$. When there is a unique optimum, then the optimal solution estimate converges to the unique global optimum.

**Corollary 2.** Suppose all the conditions in Theorem 3 are satisfied. Then, with probability one,

$$\rho(X^*_n, \mathcal{X}^*) \to 0.$$  

In addition, if there is a unique optimum, $\mathcal{X}^* = \{x^*\}$, then $X^*_n \to x^*$ with probability one.

**Proof:** See the e-companion. $\square$

### 4. An Application: Single Observation Algorithms Based on Hit-and-Run

We compare four algorithms on two global optimization test problems with noisy objective functions. The four algorithms comprise two from the Single Observation Search Algorithms (SOSA) framework and two from the Adaptive Search with Resampling (ASR) framework. Introduced by Andradóttir and Prudius (2010), ASR is an adaptive search framework that advocates performing repeated observations of objective function at sampled design points, which is in contrast to SOSA. Within each framework, we try two different samplers, one is the Improving Hit-and-Run (IHR) sampler and the other one is the sampler originally used with ASR in Andradóttir and Prudius (2010), which we call the AP sampler. The four algorithms to be tested in this study are:

1. SOSA with IHR sampler (IHR-SO),
2. SOSA with AP sampler (AP-SO),
3. ASR with IHR sampler (IHR-ASR),
4. ASR with AP sampler (AP-ASR).

The IHR-SO algorithm modifies the improving hit-and-run algorithm (IHR) (see Ghate and Smith (2008), Zabinsky (2003), Zabinsky et al. (1993), Zabinsky and Smith (2013)) by incorporating the shrinking ball approach. The class of hit-and-run algorithms has a desirable property
of efficiently converging to a target distribution with very mild assumptions (Smith 1984, Zabinsky and Smith 2013). IHR is a simplified version of Annealing Adaptive Search (AAS) algorithm (Romeijn and Smith 1994, Zabinsky 2003) with zero temperature. AAS is later shown to be approximated by Model-based Annealing Random Search (MARS) (Hu et al. 2007, 2014). Therefore, IHR can be considered a simple representative of a global optimization search engine in this class.

An early version of improving hit-and-run with single observation, but without shrinking balls, has been applied to a simulation optimization problem with encouraging computational results in Kiatsupaibul et al. (2015). A more extensive numerical study (Linz et al. 2017) showed a benefit of SOSA compared to multiple replications.

The AP sampler has been employed as the sampling strategy for ASR in the study of Andradóttir and Prudius (2010) with promising computational results. AP samples locally from the neighborhood of the current optimal solution estimate and also globally from the whole feasible region. In a sense, it is a simple version of the nested partitions methods (Shi and Ólafsson 2000a) that sample from a partition local to the current optimal solution estimate and its complement. Therefore, AP can be considered a simple representative of another class of global optimization search engine. The original combination of AP and ASR is called AP-ASR in this study. Here we also combine AP with SOSA, called AP-SO, and compare it against the original AP-ASR.

The IHR-SO and AP-SO algorithms share the property of having a positive probability of sampling anywhere in the space. Thus, as shown in Theorem 4 and Corollary 3, they satisfy the four assumptions for continuous optimization problems and converge to a global optimal value.

**SOSA with IHR Sampler (IHR-SO)**

Parameters: \( \kappa_r \), starting ball radius; \( \gamma \), order of local sample density; \( \beta \), order of radius shrinkage; and \( s \), order of slowing sequence. The parameters specify the following radii and slowing sequences:

\[
\{ r_n = \kappa_r n^{-\beta}, n = 1, 2, \ldots \} \text{ and } \{ i_n = [n^s], n = 1, 2, \ldots \}.
\]

Follow the steps of SOSA but replace Step 1 by the following IHR sampler.
Step 1: Given \(x_1, y_1, \ldots, x_{n-1}, y_{n-1}\), let \(\tilde{x}_{n-1} \in \text{Arg min}_{x \in \mathcal{X}_{n-1}} \hat{f}_{n-1}(x)\). The IHR sampler obtains a new design point \(x_n\) by first sampling a direction \(v\) from the uniform distribution on the surface of a \(d\)-dimensional hypersphere, and second, sampling \(x_n\) uniformly distributed on the line segment \(\Lambda\) where \(\Lambda = \{\tilde{x}_{n-1} + \lambda v : \lambda \in \mathbb{R}\} \cap S\). Given \(x_n\), sample \(u_n\) having the same distribution as \(U\) and evaluate \(y_n = g(x_n, u_n)\).

**SOSA with AP Sampler (AP-SO)**

The same as IHR-SO but replace Step 1 by the following AP sampler.  

Step 1: Given \(x_1, y_1, \ldots, x_{n-1}, y_{n-1}\), let \(\tilde{x}_{n-1} \in \text{Arg min}_{x \in \mathcal{X}_{n-1}} \hat{f}_{n-1}(x)\). Sample from the AP sampler by sampling \(x_n\) uniformly on \(S\) with probability 0.5, and otherwise, sample \(x_n\) uniformly on points that are within \(R\) of \(\tilde{x}_{n-1}\) on each dimension, i.e., \(\{x \in S : |x^i - \tilde{x}_{n-1}^i| \leq R\} \cap S\). For each \(x^i\) is the \(i\th\) component of \(x\) and the radius parameter \(R\) is the tuning parameter of the sampler. Given \(x_n\), sample \(u_n\) having the same distribution as \(U\) and evaluate \(y_n = g(x_n, u_n)\).

**ASR with IHR Sampler (IHR-ASR)**

Parameters: \(b\) and sequence \(\{M(i) = [i^b]\}\); \(c\) and sequence \(\{K(i) = [i^c]\}\); \(\delta\), \(L\) and \(T\). 

For each \(x \in S\), the algorithm keeps track of the following accumulators: \(N_k(x)\), the number of objective function observations collected at \(x\) at iteration \(k\); \(S_k(x)\), the sum of these \(N_k(x)\) objective function observations; and \(\hat{f}_k(x) = S_k(x)/N_k(x)\).  

*Step 0*: Let \(i = 1, k = 0\), and \(\mathcal{X}_0 = \emptyset\).  

*Step 1*: Let \(k \leftarrow k + 1\). If \(k = M(i)\), then go to *Step 2*. Otherwise, go to *Step 3*.  

*Step 2*: Sample a new design point by performing the following steps.  

2.1: If \(i = 1\), sample \(x_i\) uniformly on \(S\). Otherwise, sample \(x_i\) from the IHR sampler.  

2.2: If \(i = 1\), accept \(x_i\). Otherwise, accept \(x_i\) if \(f_L(x_i) \leq \hat{f}_{k-1}(x_{k-1}^*) + \delta\), where \(f_L(x_i)\) is an average of \(L\) observations of \(f(x_i)\). If \(x_i\) is accepted, then let \(\mathcal{X}_k = \mathcal{X}_{k-1} \cup \{x_i\}\). If \(x_i\) is rejected, \(\mathcal{X}_k = \mathcal{X}_{k-1}\). Update \(N(x_i)\) and \(S(x_i)\).  

2.3: For each \(x \in \mathcal{X}_k\), if \(N_k(x) < K(i)\), obtain \(K(i) - N_k(x)\) additional objective function observations of \(f(x)\) and update \(N(x_i)\) and \(S(x_i)\) accordingly.
2.4: Let $i \leftarrow i + 1$. Go to Step 4.

Step 3: Resample from the existing design points by performing the following steps. Let $k'$ be the last iteration that a new point is sampled.

3.1: Sample $x$ from $X_{k-1}$ according to probability $p_k(x)$ where $p_k(x) \propto \exp(\hat{f}_k'(x)/T_{k'})$ and $T_{k'} = T/\log(k' + 1)$.

3.2: Obtain an estimate of the objective function value at $x$. Update $N(x_i)$ and $S(x_i)$.

3.3: Go to Step 4.

Step 4: Select an estimate of the optimal solution $x_k^*$ from $X_k^* = \arg\min_{x \in X_k} \hat{f}_k(x)$.

Step 5: If stopping criterion satisfied, stop. Otherwise, go to Step 1.

ASR with AP Sampler (AP-ASR)

The same as IHR-ASR but replace Step 2.1 by the AP sampler.

Theorem 4 establishes that IHR-SO and AP-SO satisfy Assumption 4 when the radii of the shrinking balls are chosen appropriately. To satisfy Assumption 4, basically, the algorithm needs to generate enough sample points to fill all the balls and the balls cannot shrink too fast. Sufficient conditions that allow the algorithm to generate enough points are that the sampling densities are bounded away from zero and the feasible region is a convex set. Theorem 4 then specifies an appropriate shrinking rate of the balls.

**Theorem 4.** If $S$ is a bounded convex set, $r_n$ is of $\Omega(n^{-\beta})$, where $\beta = (1 - \gamma)/d$ and $1/2 < \gamma < 1$, and the algorithm based on SOSA employs sampling densities $q_n$ that are bounded away from zero on $S$ for all $n$, then the algorithm satisfies Assumption 4.

**Proof:** See the e-companion. □

**Corollary 3.** If Assumptions 1, 2, and 3 are satisfied, and we choose $r_n$ to be $\Omega(n^{-\beta})$, where $\beta = (1 - \gamma)/d$, $1/2 < \gamma < 1$ and $i_n \uparrow \infty$ such that $i_n \leq n^s$ where $0 < s < \gamma$, then any algorithm based on SOSA with sampling densities $q_n$ that are bounded away from zero on $S$ for all $n$ generates a sequence of optimal value estimates that converges to the optimal value $f^*$ with probability one.
Proof: By Theorem 3 and Theorem 4, Corollary 3 follows. □

Observe that the sampling densities $q_n$ implied by IHR-SO and AP-SO on a compact set are bounded away from zero. Therefore, IHR-SO and AP-SO generate sequences of optimal value estimates that converge to the global optimum with probability one.

From Theorem 3 and Theorem 4, the three parameters $\gamma, \beta,$ and $s$ of IHR-SO and AP-SO cannot be chosen independently if one would like to guarantee convergence because $\beta = (1 - \gamma)/d$ and $0 < s < \gamma$. A value of $s$, the order of slowing sequence, $i_n$, identifies how quickly we would like to adopt new optimal value estimates. From pilot experiments, a small value of $s$ will slow down the algorithm. Therefore, we adopt a large value of $s$ close to one, $s = 0.9$. This choice of $s$ dictates a large value of $\gamma$, $\gamma = 0.91$, and a small value of $\beta$, $\beta = 0.009$ for the ten-dimensional test problems. Fortunately, a small value of $\beta$ makes the ball shrink slowly and, hence, allows the algorithm to collect more observations of the objective value at each design point, enhancing the accuracy of the objective value estimates.

An appropriate value of the starting ball radius $\kappa_r$ for IHR-SO and AP-SO is problem dependent. It depends on the size of feasible region, the smoothness of the objective function and also the adaptive sampling strategy $q_n$. In this study, we experimented with several values for $\kappa_r$ for each test problem.

The two algorithms based on SOSA (IHR-SO and AP-SO) use the parameter values

\[
\gamma = 0.91, \beta = 0.009, s = 0.9, \text{ and } \kappa_r = \begin{cases} 
0.1 & \text{for Problem 1} \\
1 & \text{for Problem 2.}
\end{cases}
\]

With this set of parameters, the shrinking ball radii and the slowing sequence are set to

\[
r_n = \begin{cases} 
0.1n^{-0.009} & \text{for Problem 1} \\
n^{-0.009} & \text{for Problem 2}
\end{cases} \text{ and } i_n = \lfloor n^{0.9} \rfloor.
\]

The two algorithms based on ASR (IHR-ASR and AP-ASR) use the same parameter values used in Andradóttir and Prudius (2010),

\[
b = 1.1, c = 0.5, \delta = 0.01, L = 0.1, \text{ and } T = 0.01.
\]
The tuning parameter $R$ for AP sampler is set to 0.07 for Problem 1, and 0.4 for Problem 2, for both AP-SO and AP-ASR.

We apply the four algorithms to two problems. The first problem is the shifted sinusoidal problem, from Ali et al. (2005).

**Problem 1: Shifted Sinusoidal Problem**

$$\min \ E \ [f(x) + (1 + |f(x)|)U] \quad \text{s.t.} \quad 0 \leq x_i \leq \pi, \ i = 1, \ldots, 10$$

where $f(x) = -[2.5\Pi_{i=1}^{10} \sin(x_i - \pi/6) + \Pi_{i=1}^{10} \sin(5(x_i - \pi/6))] + 3.5$

and $x \in \mathbb{R}^{10}$ and $U \sim \text{Uniform}[-0.1, 0.1]$. According to Ali et al. (2005), this problem contains 4,882,813 local optima with a single global optimum at $x^* = (4\pi/6, \ldots, 4\pi/6)$ and $f(x^*) = 0$.

The second problem is the Rosenbrock problem. This problem is also employed as a test case for AP-ASR in Andradóttir and Prudius (2010).

**Problem 2: Rosenbrock Problem**

$$\min \ E \ [f(x) + (1 + |f(x)|)U] \quad \text{s.t.} \quad -10 \leq x_i \leq 10, \ i = 1, \ldots, 10$$

where $f(x) = 10^{-6} \times \sum_{i=1}^{d-1} ((1 - x_i)^2 + 100(x_{i+1} - x_i^2)^2)$

and $x \in \mathbb{R}^{10}$ and $U \sim \text{Uniform}[-0.1, 0.1]$. The global minimum is at $(1, \ldots, 1)$ and $f^* = 0$.

We apply each algorithm to solve each problem 100 times. The initial point for each of these 100 replications is generated according to a uniform distribution on the feasible region and used for each algorithm. Each time we run the algorithms for 12,000 function evaluations. Note that IHR-SO and AP-SO require only one objective function evaluation per iteration. Therefore, each optimization run requires exactly 12,000 iterations. For IHR-ASR and AP-ASR, we count the objective function evaluations as the algorithms progress and stop once 12,000 function evaluations are reached, which may require less than 12,000 iterations. We then record the following four measurements averaged over 100 runs, at each iteration, for $n = 1, 2, \ldots, 12,000$:

- the optimal value estimate:
  $$\hat{f}^*_n = \min_{x \in \mathbb{R}^{10}} \hat{f}_n(x),$$
the (true) objective function of the optimal solution estimate (the best candidate):

\[ f(x^*_n) \text{ where } x^*_n \in \arg \min_{x \in X_n} \hat{f}_n(x), \]

the contributions to the optimal solution estimate (the counts of the sample points that contribute to the objective function estimate of the optimal solution estimate):

\[ l^*_n = \left| \{ k \leq n : x_k \in B(x^*_n, r_n) \} \right| \]

the average noise of the optimal solution estimate:

\[ \hat{e}^*_n = \frac{\sum_{k \leq n : x_k \in B(x^*_n, r_n)} (1 + |f(x)|)U_k}{l^*_n}. \]

Figure 1 shows the four performance measurements of the four algorithms (IHR-SO, AP-SO, IHR-ASR and AP-ASR) applied to Problem 1, and averaged over 100 runs, at selected number of function evaluations. Panel (a) also shows the 95% confidence intervals of the optimal value estimates, represented by vertical bars. The confidence interval represents the variation across simulation runs. It is calculated as the mean estimate plus and minus 1.96 times the standard deviation divided by 10 (the square root of the number of simulation runs).

The two SOSA algorithms, IHR-SO and AP-SO, outperform the two ASR algorithms, IHR-ASR and AP-ASR. From panels (a) and (b) of Figure 1, we observe that the optimal value estimates and the objective function value of the optimal solution estimates of SOSA (IHR-SO and AP-SO) converge to the global optimum (target) more quickly than those of ASR (IHR-ASR and AP-SO). Panel (c) shows how objective function observations accumulate at the optimal solution estimates over the course of each of the algorithms. The errors of the optimal value estimates (panel (d)) decrease as more observations accumulate. The two ASR algorithms are designed to accumulate observations uniformly over the course of the algorithms, as seen from panel (c) of Figure 1. Panel (c) of Figure 1 also shows that IHR-SO accumulates more observations over the course of the run, although adaptively. From panel (c) of Figure 1, IHR-SO and AP-SO are more aggressive at the beginning stage of the algorithms when the quality of the optimal value estimates are poor, not
Figure 1  Performance diagnostics for IHR-SO, AP-SO, IHR-ASR and AP-ASR with respect to Problem 1. Panels (a) and (b) exhibit the optimal value estimate (with confidence intervals) and the true objective function value at the optimal solution estimate (the best candidate), respectively. Panels (c) and (d) show the contributions to the best candidates and the average noises of the optimal solution estimate as functions of objective function evaluations, respectively.
spending too many observations at the early sample points. The SOSA algorithms then adaptively accumulate more observations for the optimal value estimates in the later stage of the algorithms, when the quality of the estimates get higher. Observe that the ASR algorithms accumulate more observations for the optimal value estimates at the early stage than SOSA algorithms do, slowing the algorithms down. Consequently, as shown in panel (d) of Figure 1, the average noises of the optimal value estimates reduce more rapidly at the early stage in the case of ASR algorithms. Towards the end, the average noises of the four algorithms are not much different. Observe that the errors at the optimal solution estimates are negatively biased. This is a common behavior found in algorithms whose optimal estimates are chosen as the minimum among all the estimates, since the ones that underestimate the true objective function value will be more likely to be chosen, causing the average errors to be negative.

Table 1 shows the statistics of the optimal value estimates $\hat{f}_n^*$ of the four algorithms at termination, when applied to Problem 1. As seen from Figure 1, the mean estimates of IHR-SO and AP-SO are closer to the optimal value (zero) than those of IHR-ASR and AP-ASR. For Problem 1, the mean squared errors of IHR-SO and AP-SO are also significantly smaller than those of IHR-ASR and AP-ASR. Furthermore, for Problem 1, IHR-SO consistently outperforms IHR-ASR, and AP-SO outperforms AP-ASR at the 50 and 75 percentiles as well as the worst estimate encountered.

Figure 2 shows the four performance measurements of the four algorithms (IHR-SO, AP-SO, IHR-ASR and AP-ASR) applied to Problem 2, and averaged over 100 runs, at each number of function evaluations, up to 4,000 function evaluations. Panel (a) also shows the optimal value estimates with 95% confidence intervals, which are quite tight.

Again the two SOSA algorithms, IHR-SO and AP-SO, outperform the two ASR algorithms, IHR-ASR and AP-ASR. From panels (a) and (b) of Figure 2, we observe that the optimal value estimates and the objective function value of the optimal solution estimates of SOSA (IHR-SO and AP-SO) converge to the global optimal value more quickly than those of ASR (IHR-ASR and AP-ASR).
Figure 2  Performance diagnostics for IHR-SO, AP-SO, IHR-ASR and AP-ASR with respect to Problem 2. Panels (a) and (b) exhibit the optimal value estimate (with confidence intervals) and the true objective function value at the optimal solution estimate (the best candidate), respectively. Panels (c) and (d) show the contributions to the best candidates and the average noises of the optimal solution estimate as functions of objective function evaluations, respectively.
Table 1 shows the statistics of the optimal value estimates $\hat{f}_n^*$ of the four algorithms at termination, when applied to Problem 2. At termination ($n = 4,000$), the optimal value estimates of all four algorithms are very close to the true optimal value of zero. The mean squared errors of AP-SO and IHR-ASR are the smallest, followed by that of IHR-SO.

**Table 1** Statistics of the optimal value estimates $\hat{f}_n^*$ of the four algorithms at termination. The experiments for Problem 1 terminate with $n = 12,000$. The experiments for Problem 2 terminate with $n = 4,000$.

We also demonstrate the effectiveness and the efficiency of the framework by modifying two adaptive search algorithms, namely Improving Hit-and-Run (IHR) and Andradóttir-Prudius (AP), to fit this framework. We also show that any algorithm with sampling density bounded away from zero on the feasible region, in particular IHR-SO and AP-SO, satisfy the four assumptions and, hence, converges with probability one to a global optimum. The two algorithms under the SOSA framework outperform the same two algorithms under the adaptive search with resampling (ASR) alternative, as demonstrated on two noisy objective functions. The performance confirms the theory developed and illustrates the potential of the new continuous simulation optimization paradigm.

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