

# A Recursive Random Search Algorithm for Large-Scale Network Parameter Configuration

## Abstract

Parameter configuration is a common procedure used in large-scale network protocols to support multiple operational goals. This problem can be formulated as a black-box optimization problem and solved with an efficient search algorithm. This paper proposes a new heuristic search algorithm, Recursive Random Search(RRS), for large-scale network parameter optimization. The RRS algorithm is based on the initial high-efficiency property of random sampling and attempts to maintain this high-efficiency by constantly “restarting” random sampling with adjusted sample spaces. Due to its root in random sampling, the RRS algorithm is robust to the effect of random noises in the objective function and is advantageous in optimizing the objective function with negligible parameters. These features are very important for the efficient parameter optimization of network protocols. The performance of RRS is demonstrated with the tests on a suite of benchmark functions. The RRS algorithm has been applied to the adaptive configuration of several network protocols, such as RED, OSPF and BGP. One example application in OSPF routing algorithm is presented.

## 1 Introduction

Today’s network protocols like BGP and OSPF were designed for one primary service: “best effort reachability.” But now network operators want to deploy Virtual Private Networks(VPN), manage traffic within ASes to meet Service Level Agreements(SLA), and between ASes (at peering points) to optimize complex peering agreements. The designers of such protocols included “parametric hooks” to allow operators to “tweak” the protocols and achieve such traffic management goals. However, the parameter setting process today is manual and is widely considered a black art. The configuration of many protocols, such as BGP, is tough, error prone and is likely to get harder as the protocol is overloaded to serve more functions[1]. Though some tools are emerging to aid operators, a lot more needs to be done. The on-line simulation system proposed in[2] is one such contribution to this important space, which can be used as

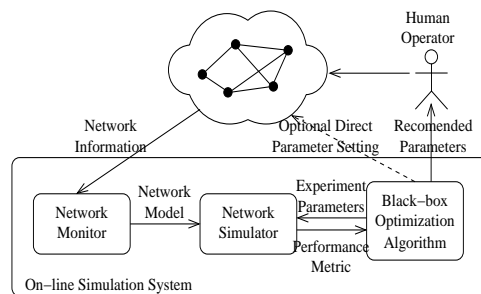


Figure 1: On-line simulation system for adaptive configuration of network protocols

a “recommendation service” to suggest a variety of “good” parameter settings and illustrate resulting flow patterns so that operators are better informed than their current manual procedures. As illustrated in Fig 1, the basic idea of this system is to formulate network protocol configuration as a black-box optimization problem. With the network

protocol considered as a black-box, network simulation can be used to evaluate its performance for various parameter settings. Based on this, an optimization algorithm can then be employed to find the good protocol configuration for the current network conditions. The “black-box” approach allows flexibility in terms of objectives and metrics of the desired optimization, and hence can be applied to a variety of configuration problems. And it can be used to manage any network protocol as long as the protocol has *tunable* parameters and their setting has substantial effect on network performance.

In this on-line protocol configuration framework, an efficient search algorithm is essential to its success. Although a large number of optimization algorithms have been proposed and successfully applied in practice[3, 4, 5], there is no single algorithm which can consistently outperform the others in every problem class[6]. For a specific type of problems, the performance of a search algorithm is dependent on whether its search strategy fit the features and requirements of the underlying problem. For network parameter optimization, the desired search algorithm should have *high efficiency*, *scalability* and *noise-robustness*. Traditional search algorithms, such as genetic algorithm and simulated annealing, could not provide such combination of properties. This paper therefore proposes a new optimization algorithm, Recursive Random Search(RRS), whose major feature is its basis on random sampling. The Recursive Random Search algorithm exploits the initial high efficiency of random sampling and remains in this high-efficiency phase by constantly restarting random sampling with adjusted sampling spaces. In addition, the RRS algorithm is also robust to noises in the objective function and is of great advantage when dealing with the objective function with negligible parameters.

The RRS algorithm has been tested on a suite of benchmark functions to examine its efficiency, noise-resistance and handling of negligible parameters. The results show that RRS outperforms the other algorithms. We have applied the RRS algorithm to the on-line configuration of several network protocols, such as, RED, OSPF and BGP. Simulation results demonstrate substantial improvement in network performance when the proposed on-line tuning is deployed.

The rest of this paper is organized as follows: In Section 2, we examine features of network parameter optimization problems and investigate their impact on the algorithm design. In Section 3, we discuss design concepts of the RRS algorithm and provide an overview of the algorithm. In Section 4, we describe the details of this algorithm. In Section 5, the tests of the algorithm on a suite of benchmark functions are presented. In Section 6, we summarize one example application of RRS to the adaptive configuration of OSPF routing algorithm. Finally, we conclude this paper in Section 7.

## 2 Network Parameter Optimization Problem

Like optimization problems arising in many engineering areas, network parameter optimization can be formulated as (assume minimization): given a real-valued objective function  $f : \mathbb{R}^p \rightarrow \mathbb{R}$ , find a global minimum,

$$\mathbf{x}^* = \arg \min_{\mathbf{x} \in D} f(\mathbf{x}) \quad (1)$$

where  $D$  is the predefined parameter space, usually a compact set in  $\mathbb{R}^p$ . In these problems, the objective function  $f(\mathbf{x})$  is often analytically unknown and the function evaluation can only be achieved through computer simulation or other indirect ways. This type of problems are also called “black-box” optimization problems where the objective function is modeled as a black-box. Since little *a priori* knowledge about the black-box is assumed, these problems are considered very hard to solve. In addition, the objective functions are often non-linear and multi-modal, and these problems are also called *global optimization* as opposed to local optimization where there is only one single extreme in  $f(\mathbf{x})$  and are much easier to solve.

Designing or selecting an efficient search algorithm requires examining the features of the target problem. For network parameter optimization problems, the following features are usually present.

**High efficiency** is required for the desired search algorithm. More specifically, the emphasis of the search algorithm should be on finding a better operating point within the limited time frame instead of seeking the strictly global

optimum. Network conditions vary with time and the search algorithm should *quickly find better network parameters* before significant changes in the network occur. Furthermore, network parameter optimization is based on network simulation which might be very time-consuming. This also requires a highly efficient search algorithm to obtain a desired solution with a minimum number of network simulations.

**High dimension** is another feature of these problems. For example, AT&T’s network has 1000s of routers and links. If all OSPF link weights of this network are to be tuned, there will be thousands of parameters present in the optimization. High-dimensional optimization problems are usually much more difficult to solve than low-dimensional problems because of “curse of dimensionality”[7].

**Noise** is often introduced into the evaluation of objective function since network simulation may be used for function evaluations. Due to inaccuracies in network modeling, simulation, etc., this empirical evaluation of objective function may be distorted from the original one, in other words, affected by small random noises. Fig 2 shows an example of 2-dimensional empirical objective function obtained with network simulation. It can be seen that there exist many irregular small random fluctuations imposed on the overall structure.

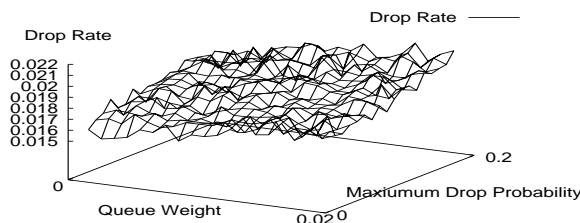


Figure 2: An empirical objective function obtained with network simulation (RED queueing management)

**Negligible parameters** may also be included in the objective function. These parameters contribute little to the objective function and should be ruled out from the optimization process. However, in practice, they are normally very difficult to be identified and eliminated effectively. If the search algorithm is able to automatically excluded these parameters from the optimization process, the efficiency of the optimization will significantly improved.

**“Globally convex”** or “big valley” structure[8, 9] may be present in the objective functions. That is, high-quality local optima tend to center around the global one and be close to each other, whereas low-quality local optima tend to distribute far way from the global one. “Globally convex” structure appears in many practical optimization problems, especially in the situations when the objective function is affected by random noises. Boese[10] has demonstrated the existence of this structure in complex Traveling Salesman Problem(TSP) and graph bisection problem, and presented an *intuitive* graph for this structure(Fig 3). Note that this figure is only an intuitive illustration of “global convex”. The structure should be considered as a global slowly-varying structure superimposed with many local irregularities. The same structure has been found in circuit/graph partitioning and job-shop scheduling, etc.[11]. Leary[12] also confirmed that there exist similar “funnel” structures in molecular conformation problems where the potential energy from the forces between atoms is minimized.

The issues described above are common in many practical optimization problems[13, 14]. For such class of problems, genetic algorithm[15] and simulated annealing[5] are the most common algorithms since they require little *a priori* information from the concerned problem and are generally applicable. However, these algorithms are mainly designed for full-optimization and often lacking in efficiency. Controlled random search[3], i.e., Price algorithm, is also recommended for this situation[13, 16] but still suffers from lack of efficiency. In practice, these optimization

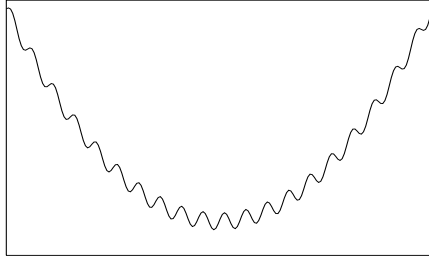


Figure 3: Big valley structure

algorithms are often combined with local search techniques, such as, deepest descent and pattern search, to improve their efficiency. However, since these local search techniques use fixed local structures to guide the search process, they are usually susceptible to the effect of noises[17]. For example, in pattern search, the wrong pattern may easily be derived if the samples for pattern exploration are corrupted by noises. Furthermore, for the objective function with “globally convex” structures, local methods also perform inefficiently since there exist a large number of low-quality local optima. Multistart local search algorithms will waste many efforts on examining these low-quality local optima and essentially work like an inefficient random sampling.

In contrast to most of the search algorithms, the RRS algorithm is mainly built upon random sampling. RRS performs the search process based on stochastic information on a certain sample area, therefore, its performance is less affected by noises. In addition, RRS is more efficient when dealing with the objective function with negligible parameters. This is because that random samples will still maintain its uniform distribution in the subspace composed of only those important parameters, and hence effectively removes negligible parameters from the optimization process. In this way, the efficiency of the search can be improved significantly. For the objective function with “globally convex” feature, RRS is able to detect the overall structure by its initial extensive sampling and approach global optima very quickly.

### 3 Design Ideas of Recursive Random Search

Random sampling is the simplest and most widely used search technique, which takes random samples from a uniform distribution over the parameter space. Despite its simplicity, random sampling is able to provides a strong probabilistic convergence guarantee, i.e., the optimization result converges to the global optimization with probability 1. Furthermore, random sampling has surprisingly proved to be more efficient than deterministic exploration methods, such as, grid covering, in terms of some probabilistic criteria and it is especially so for high-dimensional problems[18]. The disadvantage of random sampling is its apparent lack of efficiency. However, we will show that it is in fact *very efficient in its initial steps and its inefficiency is from the later sampling steps*. In the following, we first describe the initial high-efficiency property of random sampling, the basis of the Recursive Random Search algorithm, and then present the basic idea of RRS.

#### 3.1 Efficiency of Random Sampling

Given an measurable objective function  $f(\mathbf{x})$  on the parameter space  $D$  with a range of  $[y_{min}, y_{max}]$ , we can define the *distribution function* of objective function values as:

$$\phi_D(y) = \frac{m(\{\mathbf{x} \in D \mid f(\mathbf{x}) \leq y\})}{m(D)} \quad (2)$$

where  $y \in [y_{min}, y_{max}]$  and  $m(\cdot)$  denotes *Lebesgue measure*, a measure of the size of a set. For example, *Lebesgue measure* is area in a 2-dimensional space, and volume in a 3-dimensional space, and so on. Basically, the above equation represents the portion of the points in the parameter space whose function values are smaller than a certain level  $y$ .  $\phi_D(y)$  is a monotonously increasing function of  $y$  in  $[y_{min}, y_{max}]$ , its maximum value is 1 when  $y = y_{max}$  and its minimum value is  $m(x_*)/m(D)$  where  $x_*$  is the set of global optima. Without loss of generality, we assume that  $f(\mathbf{x})$  is a continuous function and  $m(\mathbf{x} \in D | f(\mathbf{x}) = y) = 0, \forall y \in [y_{min}, y_{max}]$ , then  $\phi(y)$  will be a monotonously increasing continuous function with a range of  $[0, 1]$ . Assuming a  $y_r \in [y_{min}, y_{max}]$  such that  $\phi_D(y_r) = r, r \in [0, 1]$ , a  $r$ -percentile set in the parameter space  $D$  can be defined:

$$A_D(r) = \{ \mathbf{x} \in D \mid f(\mathbf{x}) \leq y_r \} \quad (3)$$

Note that  $A_D(1)$  is just the whole parameter space  $D$  and  $\lim_{\epsilon \rightarrow 0} A_D(\epsilon)$  will converge to the global optima. Suppose the sample sequence generated by  $n$  steps of random sampling is  $\mathbf{x}_i, i = 1 \dots n$  and  $\mathbf{x}_{(1)}^n$  is the one with the minimum function value, then the probability of  $\mathbf{x}_{(1)}^n$  in  $A_D(r)$  is:

$$P(\mathbf{x}_{(1)}^n \in A_D(r)) = 1 - (1 - r)^n = p \quad (4)$$

Alternatively, the  $r$  value of the  $r$ -percentile set that  $\mathbf{x}_{(1)}^n$  will reach with probability  $p$  can be represented as:

$$r = 1 - (1 - p)^{1/n} \quad (5)$$

For any probability  $p < 1$ ,  $r$  will tend to 0 with increasing  $n$ , that means, random sampling will converge to the global optima with increasing number of samples. Fig 4 shows the  $r$ -percentile set that  $n$  steps of random sampling can reach with a probability of 99%. We can see that *random sampling is highly efficient at initial steps since  $r$  decreases exponentially with increasing  $n$ , and its inefficiency is from later samples*. As shown in Fig 4, it takes only 44 samples to reach a point in  $A_D(0.1)$  area, whereas all future samples can only improve  $r$  value of  $\mathbf{x}_{(1)}^n$  at most by 0.1.

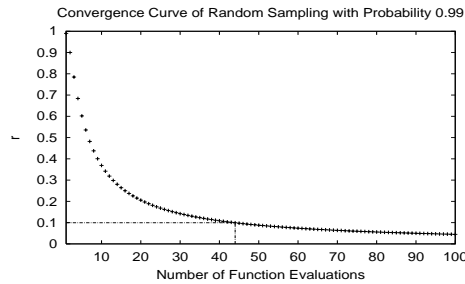


Figure 4:  $A_D(r)$  of  $\mathbf{x}_{(1)}^n$  in random sampling with probability 0.99

### 3.2 Overview of The RRS Algorithm

The basic idea of RRS is to maintain the initial efficiency of random sampling by “restarting” it before its efficiency becomes low. However, unlike the other methods, such as hillclimbing, random sampling cannot be restarted by simply selecting a new starting point. Instead we accomplish the “restart” of random sampling by *changing its sample space*. Basically, we perform random sampling for a number of times, then move or resize the sample space according to the previous samples and start another random sampling in the new sample space. Given a black-box objective function, a desired optimization process should start with inspecting macroscopic features of the objective function, and then look further into microscopic features in selected promising areas. The search process of RRS algorithm

is fully consistent with this idea. In the beginning of the search, RRS performs sampling from the whole parameter space and thus examines the overall structure of the objective function. With the search continuing and the sample space gradually shrinking, the search gets more and more details of the objective function until it finally converges to a local optimum.

A stochastic search algorithm usually comprises two elements: *exploration* and *exploitation*. Exploration examines the macroscopic features of the objective function and aims to identify promising areas in the parameter space, while exploitation focuses on the microscopic features and attempts to exploit local information to improve the solution quickly. Various search techniques can be used for these two purposes. Since macroscopic features are hard to be characterized, some unbiased search techniques, such as random search and random walk, are often used for exploration. Some algorithms also try to build a simple model to characterize the macroscopic features of the objective function and perform exploration based on this model. However, to choose an appropriate model for a certain problem is very difficult and requires extensive *a priori* knowledge. Local search methods are the most commonly used techniques for exploitation, and hence exploitation is also called *local phase* in many literature and accordingly exploration is also known as *global phase*. Derivative-based local search methods, such as quasi-Newton method[19] and deepest descent[20], are very efficient for differentiable objective functions, however, they are not suitable for many practical problem because of its sensitivity to noises and limitation for the differentiability of objective function[13]. Direct search, such as Nelder-Mead simplex method[21] and pattern search[22], do not exploit the derivative of the objective function and are more suitable for the concerned problems.

Basically, the RRS algorithm uses random sampling for exploration and recursive random sampling for exploitation. Ideally it should only execute the exploitation procedure in promising areas. However, it is difficult to determine which areas are more promising and should be exploited. Many algorithms, such as multistart type algorithms, do not differentiate areas and hence may waste much time in trivial areas. Our approach is to identify a certain *r*-percentile set  $A_D(r)$  and only start exploitation from this set. In this way, most of trivial areas will be excluded from exploitation and thus the overall efficiency of the search process can be improved. This can be illustrated by the example shown in Fig 5. The left graph shows a contour plot of a 2-dimensional multi-modal objective function and the right graph

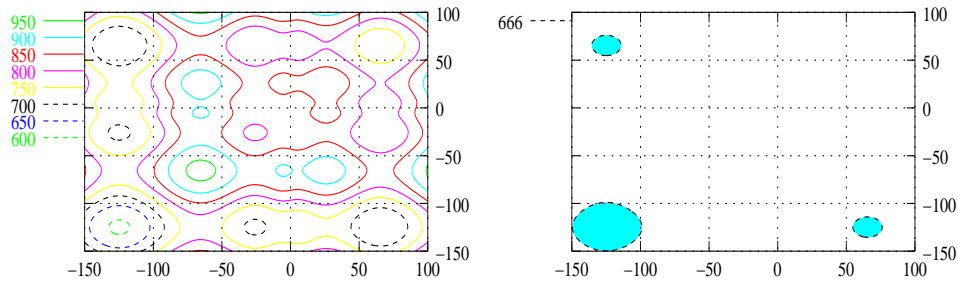


Figure 5: Contour plot of an objective function(left) and its region of  $A_D(0.05)$ (right)

shows the set of  $A_D(0.05)$ . As shown in the figure, the function has many local optima; however, only three regions of attraction remain in  $A_D(0.05)$  (shaded areas in the right plot). Each of these regions encloses a local optimum and the one with the biggest size happens to contain the global optimum. This is often true for many optimization problems since the region of attraction containing the global optimum usually has the largest size [7]. If we perform random sampling on the whole parameter space, the samples falling in  $A_D(r)$  are also uniformly distributed over  $A_D(r)$ , consequently, they are more likely to belong to the region containing the global optimum. That means, if exploitation is started from these points, the search will arrive at the global optimum with a larger probability than other non-global optima.

It is desirable that the size of  $A_D(r)$  region identified by exploration is as small as possible such that most of trivial areas are filtered out. On the other hand, its smallest size is limited by the efficiency of random sampling, i.e.,

it should be within the reach of initial high-efficiency steps of random sampling so that identifying a point in it will not take too long to lower the overall efficiency.

## 4 Algorithm Details

The basic idea of the RRS algorithm is to use random sampling to explore the whole parameter space and only start exploitation, i.e., recursive random sampling, for those points which fall in a certain  $A_D(r)$  region. The pseudo-code of the algorithm is shown in Algorithm 1 and we will explain its details in the following with reference to the lines of the pseudo-code.

### 4.1 Exploration

In the exploration phase, random sampling is used to identify a point in  $A_D(r)$  for exploitation. The value of  $r$  should be first chosen. Based on this value and a predefined confidence probability  $p$ , the number of samples required to make  $Pr(\mathbf{x}_{(1)}^n \in A_D(r)) = p$  can be calculated as (according to Equation 4):  $n = \frac{\ln(1-p)}{\ln(1-r)}$  (line 1 in pseudo-code). The algorithm uses the value of  $f(\mathbf{x}_{(1)}^n)$  in the first  $n$  samples as the threshold value  $y_r$  (line 4) and any future sample with a smaller function value than  $y_r$  is considered to belong to  $A_D(r)$ . In later exploration, a new  $\mathbf{x}_{(1)}^n$  is obtained every  $n$  samples and  $y_r$  is updated with the average of these  $\mathbf{x}_{(1)}^n$  (lines [21-23]). Note that this calculation of  $y_r$  is not intended to be an accurate estimation of the threshold for  $A_D(r)$ , instead it only function as the adjustment for the balance between exploration and exploitation. In other words, it is to ensure that on the average the exploration process will not continue for  $n$  samples and hence enter its low-efficiency phase.

In this exploration method, the confidence probability  $p$  should choose a value close to 1, for example, 0.99. The value of  $r$  decides the balance between exploration and exploitation and should be chosen carefully as discussed before. According to the current experience, we have used  $r = 0.1$  and  $p = 0.99$  in the algorithm, and with such values it only takes 44 samples to find a point for the estimation of  $y_r$ .

### 4.2 Exploitation

As soon as exploration finds a promising point  $\mathbf{x}_0$  whose function value is smaller than  $y_r$ , we start a recursive random sampling procedure in the neighborhood  $N(\mathbf{x}_0)$  of  $\mathbf{x}_0$ . The initial size of  $N(\mathbf{x}_0)$  is taken as the size of  $A(r)$ , i.e.,  $r \cdot m(D)$ , where  $D$  is the original parameter space since  $\mathbf{x}_0$  belongs to  $A(r)$  with a high probability. Currently a simple method is used to construct  $N(\mathbf{x}_0)$ : assume the parameter space  $D$  is defined by the upper and lower limits for its  $i$ th element,  $[l_i, u_i]$ , the neighborhood of  $\mathbf{x}_0$  with a size of  $r \cdot m(D)$  is the original parameter space scaled down by  $r$ , i.e.,  $N_{S,r}(\mathbf{x}_0) = \{z \in S \mid |z_i - x_{0,i}| < r^{1/n} \cdot (u_i - l_i)\}$  (line 10), where  $x_{0,i}$  is  $i$ th element of  $\mathbf{x}_0$  and  $z_i$   $i$ th element of  $\mathbf{z}$ . With this new sample space  $N_{S,r}(\mathbf{x}_0)$ , random sampling is continued. And then based on the obtained samples, the sample space is re-aligned or shrunk as exemplified in Fig 6 until its size falls below a predefined level  $\xi$ , which decides the resolution of the optimization.

**Re-align sub-phase** As described above, exploitation first starts in the neighborhood  $N(\mathbf{x}_0)$  of  $\mathbf{x}_0$ . If  $\phi_{N(\mathbf{x}_0)}(f(\mathbf{x}_0))$  (defined in Equation 5) is large, that means most points in  $N(\mathbf{x}_0)$  are better than  $\mathbf{x}_0$ . Therefore, if we do random sampling in  $N(\mathbf{x}_0)$ , it will be highly likely to find a point better than  $\mathbf{x}_0$  with a small number of samples. Let's define an expected value of  $\phi_{N(\mathbf{x}_0)}(f(\mathbf{x}_0))$ ,  $v$ , with a confidence probability  $q$ , random sampling should find a better point in  $N(\mathbf{x}_0)$  with  $l = \frac{\ln(1-q)}{\ln(1-v)}$  (line 2) samples. If a better point is found within  $l$  samples, we replace  $\mathbf{x}_0$  with this point, move the sample space to the new  $N(\mathbf{x}_0)$  and keep its size unchanged (lines [11-13]). This is called *re-align* operation. For example, in Fig 6, the exploration identifies a promising point  $C_1$  and then the exploitation (i.e., random sampling) start in the neighborhood  $R_1$  of  $C_1$ . After a few samples, a new point  $C_2$  is found to be better than  $C_1$ , hence the sample space is moved from  $R_1$  to the neighborhood  $R_2$  of  $C_2$ . In this

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**Algorithm 1: Recursive Random Search**

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1 Initialize exploration parameters  $p, r, n \leftarrow \ln(1-p)/\ln(1-r)$  ;
2 Initialize exploitation parameters  $q, v, c, s_t, l \leftarrow \ln(1-q)/\ln(1-v)$ ;
3 Take  $n$  random samples  $x_i, i = 1 \dots n$  from parameter space  $D$ ;
4  $\mathbf{x}_0 \leftarrow \arg \min_{1 \leq i \leq n} (f(\mathbf{x}_i)), y_r \leftarrow f(\mathbf{x}_0)$ , add  $f(\mathbf{x}_0)$  to the threshold set  $\mathbf{F}$ ;
5  $i \leftarrow 0, \text{exploit\_flag} \leftarrow 1, \mathbf{x}_{opt} \leftarrow \mathbf{x}_0$ ;
6 while stopping criterion is not satisfied do
7   if  $\text{exploit\_flag} = 1$  then
8     // Exploit flag is set, start exploitation process
9      $j \leftarrow 0, f_c \leftarrow f(\mathbf{x}_0), \mathbf{x}_l \leftarrow \mathbf{x}_0, \rho \leftarrow r$ ;
10    while  $\rho > s_t$  do
11      Take a random sample  $\mathbf{x}'$  from  $N_{D,\rho}(x_l)$ ;
12      if  $f(\mathbf{x}') < f_c$  then
13        // Find a better point, re-align the center of sample space to the new point
14         $\mathbf{x}_l \leftarrow \mathbf{x}', f_c \leftarrow f(\mathbf{x}')$ ;
15         $j \leftarrow 0$ ;
16      else
17         $j \leftarrow j + 1$ ;
18      endif
19      if  $j = l$  then
20        // Fail to find a better point, shrink the sample space
21         $\rho \leftarrow c \cdot \rho, j \leftarrow 0$ ;
22      endif
23    endw
24     $\text{exploit\_flag} \leftarrow 0$ , update  $\mathbf{x}_{opt}$  if  $f(\mathbf{x}_l) < f(\mathbf{x}_{opt})$ ;
25  endif
26  Take a random sample  $\mathbf{x}_0$  from  $S$ ;
27  if  $f(\mathbf{x}_0) < y_r$  then
28    // Find a promising point, set the flag to exploit
29     $\text{exploit\_flag} \leftarrow 1$ ;
30  endif
31  if  $i = n$  then
32    // Update the exploitation threshold every  $n$  samples in the parameter space
33    Add  $\min_{1 \leq i \leq n} (f(\mathbf{x}_i))$  to the threshold set  $\mathbf{F}$ ;
34     $y_r \leftarrow \text{mean}(\mathbf{F}), i \leftarrow 0$ ;
35  endif
36   $i \leftarrow i + 1$ ;
37 endw
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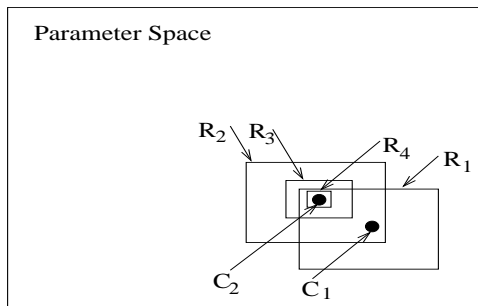


Figure 6: Shrink and Re-align Process

way, even if the initial  $N(\mathbf{x}_0)$  (i.e.,  $R_1$  in the example) might miss the local optimum, the later re-align moves will still lead the search to converge to the local optimum.

**Shrink sub-phase** If random sampling fails to find a better point in  $l$  samples, that suggests  $\phi_{N(\mathbf{x}_0)}(f(\mathbf{x}_0))$  is smaller than the expected level  $v$ . In this case, we reduce  $N(\mathbf{x}_0)$  by a certain ratio  $c \in [0, 1]$ , i.e., generate a new neighborhood  $N'(\mathbf{x}_0)$  whose size is  $c \cdot m(N(\mathbf{x}_0))$  (lines [15-16]). This is called *shrink* operation, which is performed only when we *fail* to find a better point in  $l$  samples. When the size of sample space is reduced to a value such that  $\phi_{N(\mathbf{x}_0)}(f(\mathbf{x}_0))$  is larger than  $v$ , then “re-align” will take over again to moves to the local optimum. With re-align and shrink alternately performed, the sample space will gradually converge to the local optimum. For example, in Fig 6, after  $l$  unsuccessful samples in  $R_2$ , the sample space is shrunk to  $R_3$ , then to  $R_4$  if sampling in  $R_3$  continue to fail. The exploitation process continues until the size of sample space falls below a certain threshold, whose value is dependent on the resolution requirement of the optimization problem.

## 5 Tests on Standard Benchmark Functions

As discussed before, the design objectives of RRS are: high efficiency, scalability to high-dimensional problems, robustness to function evaluation noises and capability of handling negligible parameters. This section will present the performance tests of RRS in these aspects. A suite of classical benchmark functions have been used in our performance tests, such as, Square Sum, Rastrigin[23] and Griewangk[7], most of which have a large number of local optima and are considered very difficult to optimize. For example, Fig 7 shows the 2-dimensional version of one benchmark function, Rastrigin function.

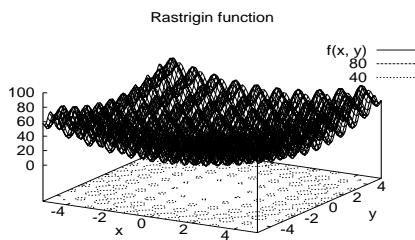


Figure 7: Two-dimensional Rastrigin function (Note: our tests use 20-2000 dimensional versions of this function.)

The performance of a search algorithm is usually measured by examining the number of function evaluations to obtain a point satisfying a certain goodness criterion. Since the emphasis of our design objective is not on full optimization but achieving high efficiency in the limited time frame, we have used the convergence curve of the

optimization, i.e., the optimization result as a function of the number of function evaluations, to compare the efficiency of algorithm especially in the initial part of the optimization process. Basically, we execute the search algorithm for a certain number of function evaluations and draw the convergence curve. The performance of the algorithms are compared based on these convergence curves.

## 5.1 Tests on Efficiency of RRS

In the efficiency tests, the performance of RRS is compared with two popular search algorithms: controlled random search and multistart pattern search. Controlled random search is recommended for black-box optimization problems in many literature[24, 13]. Multistart type algorithms are also one of the most popular methods in practice[14] and have been demonstrated to work very well and outperform some more sophisticated algorithms, such as genetic algorithm and simulated annealing, in many practical problems[25, 26, 27]. Pattern search[22] is one of local search techniques which are usually recommended for black-box optimization[28].

In the tests, the search algorithms are executed on each function with dimension varying from 20 to 2000. To eliminate randomness caused by stochastic elements in the search algorithms, each test is repeated for 50 times with random starting points and the average of the results is used. The following parameters for the RRS algorithm have been used in the tests:  $p = 0.99, r = 0.1, c = 0.5, v = 0.8, q = 0.99, s_t = 0.001$ . Fig 8 shows one set of test results for Rastrigin function. Complete results on all benchmark functions are presented in [29]. The results show that the RRS algorithm performs much more efficiently than the other two search algorithms. Controlled random search is more like pure random search in the beginning of the search. From the results, we can see that it does perform very efficiently at its initial few steps and is better than multistart pattern search. However, with the search continuing, its performance quickly degrades since random sampling loses its efficiency in later sampling. The results also demonstrated that multistart pattern search cannot perform well for the objective function with a large number of local optima.

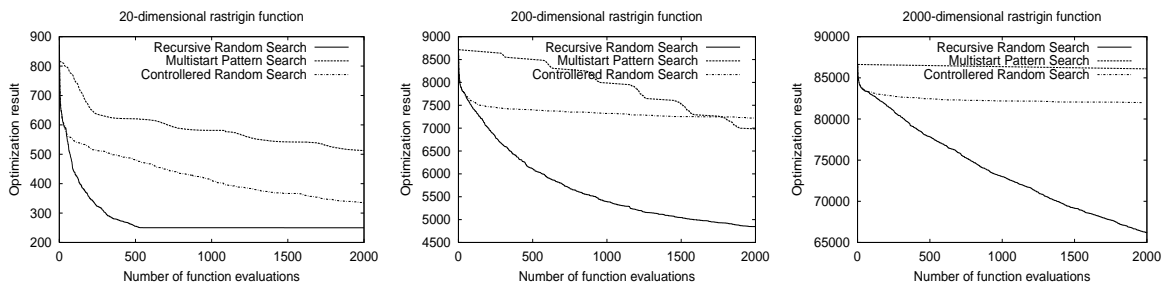


Figure 8: Performance tests on Rastrigin function

## 5.2 Tests on Noise-Resistance of RRS

Compared with local search methods, random sampling is more robust to the effect of noises. This section will compare the performance of RRS and multistart pattern search algorithm for noise-affected objective functions. Directly imposing random noises on the objective function may introduce the randomness into test results. Therefore, to obtain consistent results, Rastrigin function have been used to emulate the situations where the evaluation of the objective function is affected by small noises. Rastrigin function is defined as:

$$f(\mathbf{x}) = n \cdot A + \sum_{i=1}^n (x_i^2 - A \cdot \cos(2\pi x_i)) \quad (6)$$

It can be also considered as a simple sphere function  $\sum_{i=1}^n x_i^2$  superimposed with the noise term  $\sum_{i=1}^n A \cdot \cos(2\pi x_i)$ . The magnitude of noises is determined by the value of  $A$ . To test the noise-resistance of the search algorithms, we vary

the noise level in Rastrigin function, i.e., the value of  $A$ , and see how the search algorithms perform under different magnitudes of noises. Note that the noise magnitude should not be too large to distort the overall structure of the original function. Fig 9 shows the test results on Rastrigin functions with different noise level and different dimensions. The results demonstrate that increasing magnitude of noises seriously degrade the performance of multistart pattern search while the effect on RRS is slight.

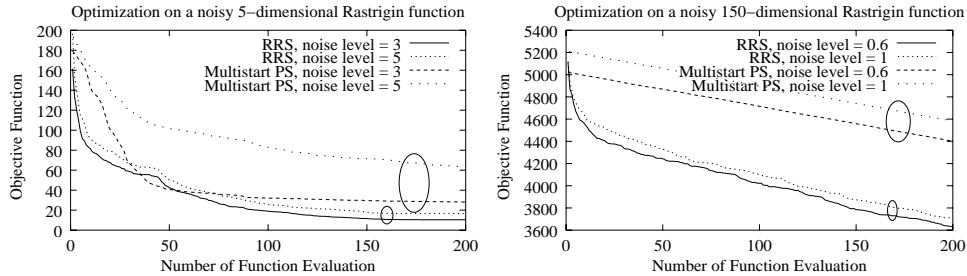


Figure 9: Noise-resistance tests of search algorithms

### 5.3 Tests on Objective Functions with Negligible Parameters

To simulate the situation of negligible parameters, an  $n$ -dimensional test function in Equation (7) is used:

$$f(\mathbf{x}) = \sum_{i=1}^5 x_i^2 + 10^{-12} \cdot \sum_{i=5}^n x_i^2 \quad (7)$$

where  $-500 < x_i < 500, i = 1 \dots n$ . In this function, the first five parameters essentially determine the function value while the others are trivial parameters with little effect. The tests are performed for the cases where there are 0, 5 and 10 negligible parameters, and the performances of RRS and multistart pattern search are compared. Fig 10 shows the test results. It can be seen that the introduction of trivial parameters has little effect on the performance of RRS while the performance of multistart pattern search degrades considerably with increasing number of trivial parameters.

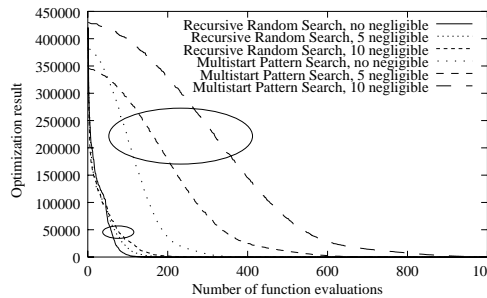


Figure 10: Performance tests on objective functions with negligible parameters

## 6 Application to Parameter Optimization of Network Protocols

The RRS algorithm has been applied to the adaptive configuration of several network protocols. For example, Random Early Detection (RED) algorithm is dynamically tuned to maintain its stability in varying network conditions and achieve high utilization as well as low queuing delay. And we have also tuned `local_pref` attributes of a BGP

domain to achieve the load balancing objective for outbound traffic. These applications are presented in detail in [30]. This section summarizes one of them as an example, i.e., traffic engineering by tuning OSPF routing metrics.

Traffic engineering is a very important aspect in networking research. The objective of traffic engineering is to distribute the offered traffic load evenly across the network such that network resources are optimally utilized. In current Internet, IP traffic is mapped onto the network by standard routing protocols. However, these routing protocols normally do not take into account current network conditions and Quality of Service(QoS) constraints. As a result, the routing generated by these algorithms tend to generate a highly uneven mapping of traffic. That is, some links may get very congested and the other may be consistently underutilized.

Suppose the offered traffic load in a network is defined by a demand matrix, where the row index represents the source, the column index is the destination and the element of the matrix is the offered load from the source to the destination. The demand matrix can be obtained through network measurement or estimation. A routing algorithm will decide the network path taken by the traffic from a source to a destination. In this way, the traffic load represented by the demand matrix is mapped to the network. Open Shortest Path First(OSPF) is the *de facto* standard routing protocol for the intra-domain traffic, i.e., the traffic transferred within the same network domain. In OSPF algorithm, each network link is assigned with a link weight and the routing decision is completely based on the values of the link weights i.e., it routes the traffic from source to destination through the path with the minimum total link weight. Traditionally, the link weights in OSPF are set heuristically without considering QoS requirements. With the knowledge of the demand matrix, the OSPF link weights can be tuned to achieve traffic engineering objectives. This tuning problem has been demonstrated to be NP-hard [31] can be tackled with a black-box optimization approach.

To formulate the black-box optimization problem for OSPF link weight setting, an optimization objective, i.e., the performance metric of the network, has to be defined first. Consider a network composed of  $n$  links. For one link  $l_i$ , we calculate its link cost  $\phi_i$  based on the following formula:

$$\phi_i(x_i) = \begin{cases} 1 & \text{for } 0 \leq x_i < 1/3 \\ 3 & \text{for } 1/3 \leq x_i < 2/3 \\ 10 & \text{for } 2/3 \leq x_i < 9/10 \\ 70 & \text{for } 9/10 \leq x_i < 1 \\ 500 & \text{for } 1 \leq x_i < 11/10 \\ 5000 & \text{for } 10 \leq x_i \end{cases} \quad (8)$$

where  $x_i$  is the normalized traffic load distributed on link  $l_i$ . Then, the objective of this optimization problem is to minimize the total link cost

$$\Phi = \sum_{i=1}^n \phi_i \quad (9)$$

This performance metric has been used for the design of AT&T WorldNet backbone network[31]. It heavily penalizes the routing causing congestion. Consequently the optimization of this metric will result in a good routing which evenly distributes traffic and voids link congestion.

Let  $\mathbf{w}$  denote the link weight vector, whose  $i$ th element is the link weight for link  $l_i$ . For each specific  $\mathbf{w}$ , network simulation is run to evaluate the objective function  $\Phi$ . Given a parameter space for  $\mathbf{w}$ , optimization can be performed to obtain the optimal or near-optimal solution of  $\mathbf{w}$ . Fig 11 shows the convergence curves of RRS on ARPANET and MCI network topologies which have 140 and 62 link weights, respectively. For comparison, we also show the convergence curves of a tabu-enhanced multistart hillclimbing search algorithm used in [31]. The straight lines in the figures indicates the performance metric when one of heuristic configuration methods, unit link weights, is used. The results show that the RRS algorithm performs very efficiently in the test. If we use the performance of the heuristic setting as the benchmark level, we can see that RRS may find a solution better than the heuristic setting with 62.4% fewer function evaluations than the other algorithm.

Besides the performance metric presented in this paper, we also tried other metrics, such as, overall packet loss rate, whose optimization is especially important for TCP traffic load. In practice, the operator can formulate arbitrarily

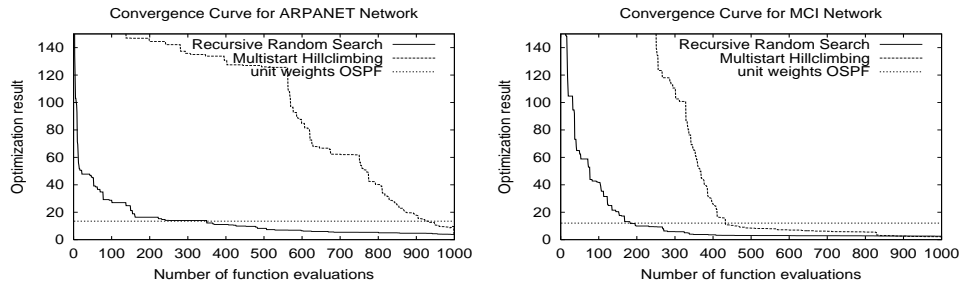


Figure 11: Convergence curves of optimization algorithms for OSPF link weight setting of ARPANET(140 links) and MCI network(62 links)

complex objectives according to their needs and request RRS to work on it to give the best-answer-so-far within a time bound.

## 7 Conclusion

This paper presented a new heuristic search algorithm, Recursive Random Search, which is designed for on-line network configuration optimization problems where the efficiency and noise-robustness of the algorithm are highly emphasized. In contrast to most other search algorithms, the new algorithm is mainly based on random sampling and does not include any traditional local search methods which are not scalable to high-dimensional problems and sensitive to the effect of noises. By constantly restarting random sampling with adjusted sample spaces, high efficiency at initial steps of random sampling can be maintained.

The test results on a suite of benchmark functions have shown that the RRS algorithm performs very efficiently, and is more robust to the effect of noises than the other algorithms. It is also demonstrated that RRS is of great advantage when dealing with the problems with negligible parameters. The RRS algorithm has been successfully used to the on-line configuration of several network protocols, such as RED, BGP and OSPF. However, its potential applications are not limited to these and the employment on other network protocols is under investigation.

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