

# Population Training Approach to Unconstrained Numerical Optimization

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## Abstract.

This paper describes the preliminary results of an evolutionary approach based on population training in heuristics for minimization of unconstrained functions. The population-training algorithm has dynamic size population ranked by an adaptation coefficient. Computational tests are presented and its performance is compared with a steady-state genetic algorithm that combines some well-succeeded features of other genetic algorithms.

## 1. Introduction

The basic idea of genetic algorithms is to store a population of individuals (or chromosomes), representing candidate solutions for concrete problems, that evolves along the time (or generations) through a competition process, where the most adapted (better fitness) have better survival and reproduction possibilities. The evolutionary process is based on individuals' selection and modification of the solutions that they represent through genetic operators as crossover and mutation.

In function parameter optimization, real-coded genetic algorithms have been successively applied and authors have reported the easiness and flexibility of its implementation [1]. On the other hand, with fixed-length binary coding, there exists a fixed amount of precision that an algorithm can be hoped to achieve, and some precision matters have to be considered before an application.

Several test functions can be found in literature and, frequently, many researchers have used them to study the performance of optimization algorithms. This work uses some of well-known test functions, such as michalewicz, rosenbrock, schwefel, langerman, griewangk, and rastrigin [1][2][3][4]. There exist many applications related to function parameter optimization well suitable to evolutionary algorithm application, such as neural network training, fuzzy set optimization, inverse problems, and others.

This work describes the preliminary results of an evolutionary approach based on population training in heuristics applied to minimization of unconstrained functions. Some experiments are presented, aiming to compare this new approach with a standard real-coded

genetic algorithm with static size population ranked by individual fitness.

Population training approach consists in training a dynamic size population with respect to a specific-problem heuristic. In this case, the evolutionary process privileges well-adapted individuals in selection and improves its surviving time.

This paper is organized as follows. Section 2 presents a standard genetic algorithm that employs a steady-state population updating and others well-known features. Section 3 presents the foundations of the population-training algorithm. Section 4 shows computational results using test functions taken from the literature. The main conclusions are presents at the end of this paper.

## 2. Steady-State Genetic Algorithm

A standard genetic algorithm was built, incorporating well-known operators found in the literature, such as roulette wheel selection [5], blend crossover [6], and non-uniform mutation [1]. In this work, the standard genetic algorithm will be called Steady-State Genetic Algorithm (SSGA).

SSGA employs a steady-state method to update the population [7]. In this method, each descendant replaces one of the low-fitness individuals found in the population. Explaining better, each crossover generates one descendant that replaces a lower-fitness individual in the population. Then, a new lower-fitness individual is found and marked to be replaced after the next crossover, and so on.

This updating method was chosen because it does not work with temporal gaps, where a population is enough explored before be replaced by another entire one (generation updating). Instead, the population contains individuals generated in all epochs, as well as the updating method employed by the population-training algorithm, presented in following. In both approaches, the offspring directly compete with their parents for selections and crossover.

The SSGA codes the function variables directly as real numbers in individuals of a static size population that evolves along the generations always preserving the best individual.

In order to compare the difference between the behaviors of both approaches, some experiments were made. The first experiment with SSGA employs the 2-dimension Langerman's function. An initial population was generated and evolved along the generations. In the Figure 1, it is showed the population over the function landscape through four snap images: after 100, 500, 1000 and 5000 generations. Notice that a reasonable diversity is maintained in the first 500 evaluations, but quickly individuals are grouped around the three promising regions (about 1000 generations). At the end, the population abandons one of these regions, staying around two local minimums, one of the which is the

global minimum (about  $F(9.76,0.68) = -1.081$ ). One can see the best niche pointed out by lower arrow.

This experiment was repeated with the 2-dimension Schwefel's function that has several local minimums. Once again, the SSGA population converges quickly to lower value surfaces on search space. A snap image is showed in Figure 2, after 1000 generations. One can concludes that, in spite its simplicity, SSGA is able to achieve solutions in 2-dimension fashion and has a wanted behavior in these cases.

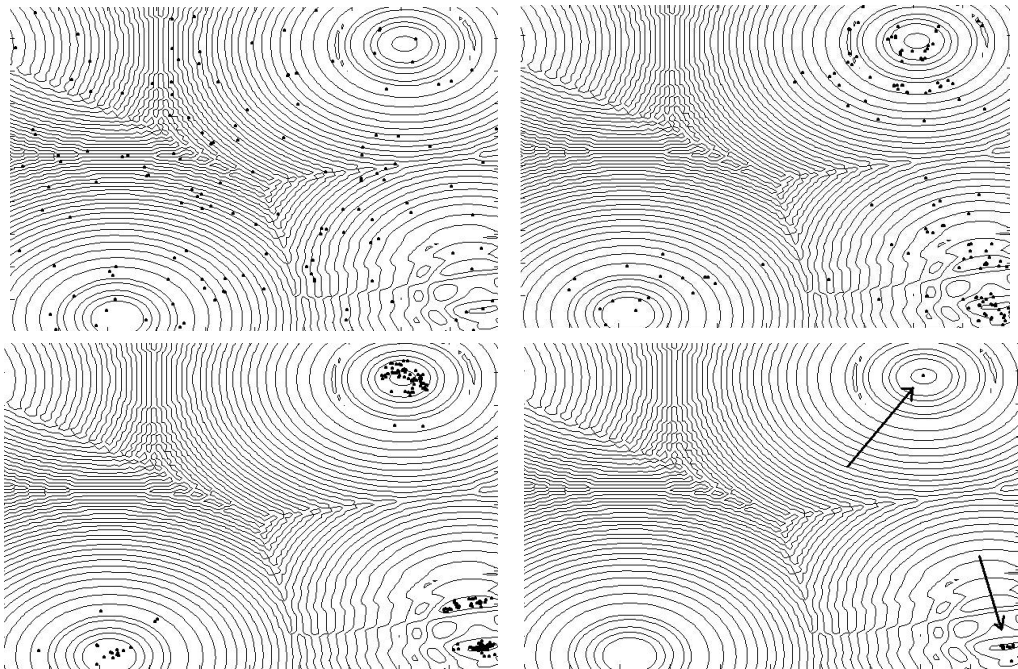


Figure 1. SSGA on 2-dimension contour map of Langerman's function after 100, 500, 1000 and 5000 generations.

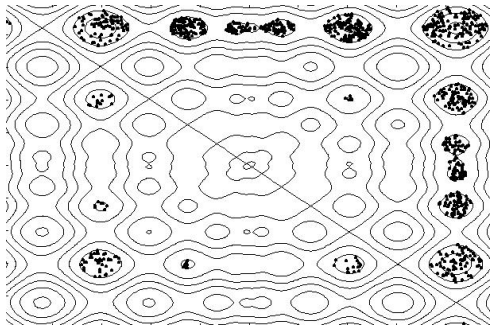


Figure 2. SSGA on 2-dimension contour map of Schwefel's function after 1000 generations.

### 3. Population Training Algorithm

The Population Training Algorithm (PTA) ranks a dynamic size population by a so-called adaptation coefficient. This new attribute considers both the objective function value and the error with respect to a specific training heuristic. The training heuristic is used to extract some information about the problem and guides the evolution process aiming performance improvement. A similar idea have been employed in Constructive Genetic Algorithm (CGA) proposed by Lorena and Furtado (2000) to clustering problems [8], and recently applied to permutation problems [9].

In CGA, the adaptation coefficient is also used to penalize schemata (individuals with incomplete information). In this work, the adaptation coefficient of an individual  $s_k$  is calculated by:

$$\delta(s_k) = d \cdot (G_{\max} - g(s_k)) - (g(s_k) - f(s_k)) \quad (1)$$

where  $G_{\max}$  is a constant upper bound to the objective function values;  $d$  is a proportionality constant with values in  $[0,1]$ ;  $g(s_k)$  is the objective function value for  $s_k$ ; and  $f(s_k)$  is the objective function value for a better neighbor of  $s_k$ , obtained by the training heuristic. In (1) there are two components:

- Interval 1 (I1):  $d \cdot (G_{\max} - g(s_k))$  measures the maximization of the difference between the upper bound  $G_{\max}$  and objective function value;
- Interval 2 (I2):  $g(s_k) - f(s_k)$  measures the error obtained with respect a training heuristic

Before the initial population is generated, the upper bound  $G_{\max}$  is set, by the selection of an individual with higher objective function value, inside a small group of them, which was generated at random. The constant  $d$  is a parameter to be tuned.

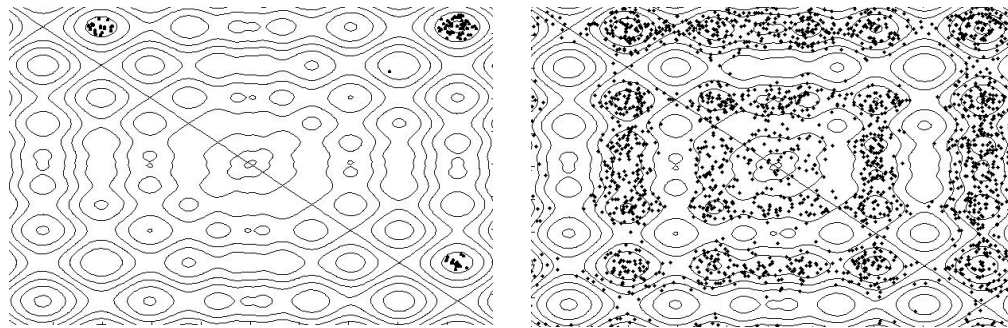
Each individual that is generated receives a ranking value  $\delta$ . Firstly, the function objective value  $g(s_k)$  is calculated. Then, a training heuristic is applied to  $s_k$ , trying improve it. If it fails,  $s_k$  is well-adapted

with respect to that heuristic. Otherwise, the amount I2 is non-zero and is subtracted from I1, penalizing the individual. Thus, once  $G_{\max}$  is not changed along the generations, the higher rankings in minimization problems are obtained by individuals with lower values of  $g(s_k)$  and  $g(s_k) - f(s_k)$ . In other words, the original problem is modeled as a new one: maximizing the interval I1 and minimizing the interval I2.

In the Figure 3, the effect of each one of the two intervals, I1 and I2, is showed separately. The individuals with maximum I1 are positioned in most promising regions (Figure 3a), whilst the individual with minimum I2 are positioned in plateaus or regions from where they are likely to be a local minimum (Figure 3b). The entire population fills a meaningful piece of search space. The unfilled pieces are that where  $g(s_k)$  closes to  $G_{\max}$  (high values) and the individuals initially generated in there are progressively eliminated by an adaptive rejection threshold, to be explained later.

#### 3.1. The training heuristic

For better understanding the I2 interval is indispensable to explain the heuristic employed to train the population and the evolution process. In the performed experiments, the following well-known local optimizers (acting as heuristics) to numerical optimization are tested: the down-hill simplex [10], and the popular finite-difference gradient method. These methods are computationally expensive, because their complexity is a function of the dimension of the problem (number of variables on objective function). Besides, the training method shall to be applied to each individual when it is generated. For this reason a third method was built and tested, achieving good results: the in-line search method.



**Figure 3. Interval minimization effect on 2-dimension contour map of Schwefel's function after 1000 generations: a) individuals with  $d \cdot (G_{\max} - g)$  values about 80% of the best  $\delta$ ; b) individuals with  $(g - f)$  values equal to zero (well-trained by a heuristic)**

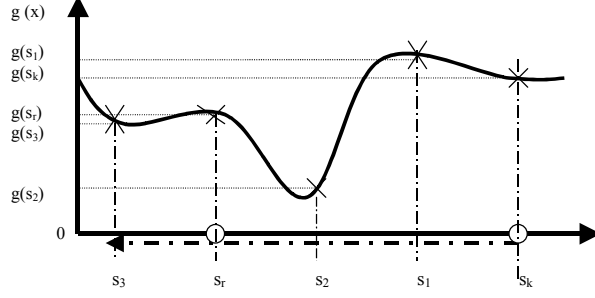


Figure 4. One-dimension example of the in-line search method

The in-line search method is based on the idea that it can be better to walk to well-adapted individuals. Thus, an individual  $s_r$  is chosen at random from the set of well-adapted ones. This set is defined through a percentage of population. For example, the 20% better ranked individuals form the set of individuals from where one will be selected as reference point  $s_r$  to in-line search method. Then, a fixed number of individuals ( $s_1, s_2, \dots, s_n$ ) are sampled along the line between  $s_k$  and  $s_r$  and the best objective function evaluation is held on as  $f(s_k)$  value. In order to extrapolate the region between  $s_k$  and  $s_r$ , the method take other samples after  $s_r$ .

The Figure 4 shows a one-dimension example of the in-line search method being performed. In this case, two individuals are selected between  $s_k$  and  $s_r$  and a third one is selected after  $s_r$ , extrapolating the interval. Notice that the second sample,  $s_2$ , had got the best objective function value. Then,  $g(s_2)$  is assigned to  $f(s_k)$ .

### 3.2. The recombination process

The ranked individuals in expression (1) are maintained in descendent order, i.e., the first individuals are better adapted than the last ones. In this work, the selection operator was built to privileges the well-adapted individuals. Two individual are selected: one come from the population elite and the other come from the entire population. The elite group is the same used to choose the reference point in the in-line search method. A similar selection operator has been employed on Constructive Genetic Algorithms and is called base-guide selection [8]. After the selection of two individuals, the blend crossover (BLX- $\alpha$ , with  $\alpha=0.25$ ) is also applied to generate another individual, and this new individual can undergo a non-uniform mutation [1] with low probability (about 1%). Then, the new individual is evaluated ( $\delta$ -calculation) and updated in the dynamic size population, according to its adaptation coefficient. The updating process, as well as the entire evolution process, is explained at following.

### 3.3. The evolution process

The updating process of PTA privileges the well-adapted individuals that are placed in the top of the ranked population. An adaptive rejection threshold,  $\alpha$ , provides a progressive elimination of the ill-adapted individuals. It will be related to evolution time (generation).

The population at the evolution time  $\alpha$ , denoted by  $P_\alpha$ , is dynamic and its size varies accordingly the value of the adaptive parameter  $\alpha$ , and can be emptied during the process. The parameter  $\alpha$  is now compared to the ranks in expression (1), thus yielding the following expression:

$$\alpha \geq \delta(s_k) = d \cdot (G_{max} - g(s_k)) - (g(s_k) - f(s_k)) \quad (2)$$

At the time they are created, individuals receive their corresponding rank  $\delta(s_k)$ . At each generation, these  $\delta$  are compared with the current evolution parameter  $\alpha$ . If the condition in (2) is satisfied, the individual  $s_k$  is eliminated.

Considering that the well-adapted individuals need to be preserved for recombination, the evolution parameter  $\alpha$  is started from the lowest  $\delta$  value (taken from the bottom of the ranked population), and then increases with step proportional to actual population size  $|P|$ .

$$\alpha = \alpha + k \cdot |P| \cdot \frac{\delta_{top} - \delta_{bot}}{Gr} + l \quad (3)$$

where  $k$  is a proportionality constant,  $l$  is the minimum increment allowed,  $Gr$  is the remaining number of generations, and  $(\delta_{top} - \delta_{bot})$  is the actual range of values of  $\delta$ .

One can observe that the adaptive increment of  $\alpha$  is affected by the own environment (population size, best and worst  $\delta$ 's, etc). Thus, once the PTA achieves very good regions and does not get to improve the best rank, the parameter  $\alpha$  goes eliminating the individuals until the population is emptied. The Figure 5 shows four snap images after 100, 500, 1000 and 5000 generations for 2-dimension Langerman's function.

Notice that dynamic population fills great part of the search space and, along the generations, and the offspring are attracted to promising regions, like SSGA, but the density of individuals is reasonably greater. Perhaps this fact can contribute for an intensification of pressure over the local minimums as well as over the global one. At the end, about generation 5000, the population was practically

emptied and few individuals are left in two minimums, including the same global minimum found by SSGA. The overall numerical results of all experiments that were made with both SSGA and PTA approaches are summarized and commented in next section.

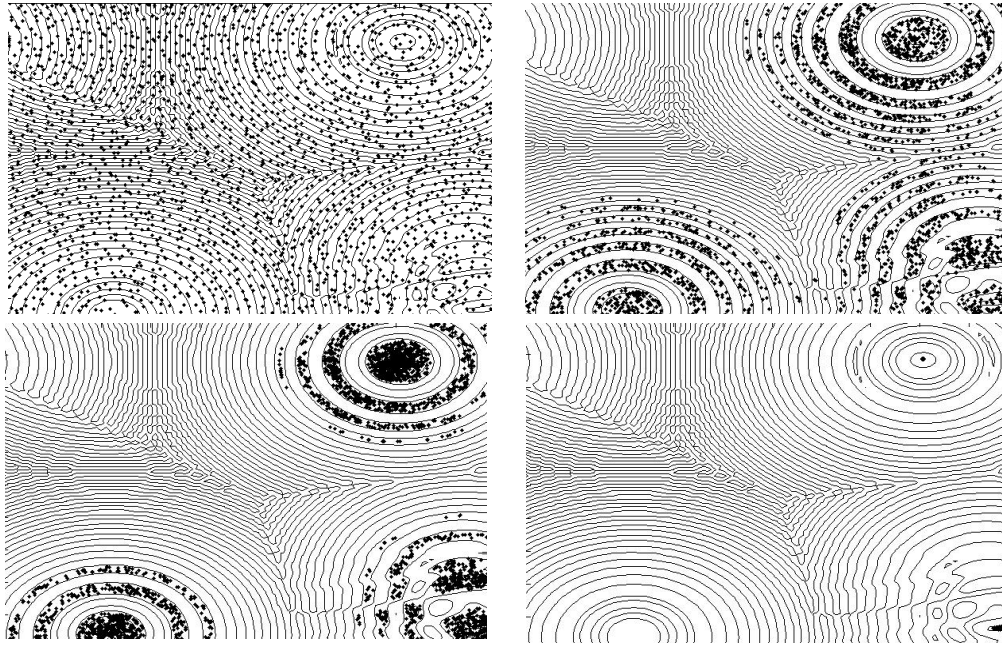


Figure 5. PTA on 2-dimension contour map of Langerman's function after 100, 500, 1000 and 5000 generations.

#### 4. Computational tests

The SSGA and PTA were coded in ANSI C and run on Intel Pentium III (450Mhz) hardware. For the computational tests, some parameters were adjusted in both approaches. Once adjusted, 20 trials were performed with 6 test function in the 5-dimensions and 10-dimension cases, giving 240 trials with each approach.

The best results found with SSGA were obtained with non-uniform mutation and blend crossover probabilities equal to 5% and 100% respectively and an initial population of 300 up to 1000 individuals. The blend crossover parameter  $\alpha$  is equal to 0.25, as well as in PTA. The best results found in PTA were obtained with  $d$  parameter set to 0.10. This configures the proportionality between intervals  $I_1$  and  $I_2$ . The  $k$  and  $l$  constants used at the step size of the adaptive rejection threshold  $\alpha$  were set to  $1 \cdot 10^{-2}$  and  $1 \cdot 10^{-5}$  in most of the test functions. But in 10-dimension Langerman's function, it was not possible to find the right values to them. The results for this test function were under the expectation. In PTA, the non-uniform mutation probability was set to 1%.

The stop criterion is: a) the expected best solution is found; or b) after 100 generations the best individual is not changed; or c) the limit of 751000 evaluations of the objective function is reached. The expected best solutions used on stop criteria are known in the literature and shown in Table I. Some executions of SSGA did not reach best solutions and had an excessive number of functions evaluations.

For each one of the 20 trials, the average ( $M$ ) of the solutions found is calculated and the percentage of the error related to the expected solution can be calculated by  $[M-F(s^*)]$ , where  $F(s^*)$  is the expected best solution.

Table I also shows the comparison between PTA and SSGA for 5-dimension and 10-dimension test functions. In 5-dimension, PTA approach presents a best overall performance, even without considering the poor performance of SSGA in rosenbrock's function. PTA obtains 7.66% of average error against 15,88% obtained by SSGA. In 10-dimension, once again SSGA runs poorly in rosenbrock's function (the best solution found was 6.10). Excluding the rosenbrock's result of both, there exist an equilibrium of performance: PTA with 23.95% of error and SSGA with 26.75%. In 5-dimension, PTA had better

performance in rosenbrock's, and langerman's functions, although in this last, both approaches did not have any success. In 10-dimension, PTA was better than SSGA in rosenbrock's, langerman's, and michalewicz 's functions.

The running times and the number of objective function evaluations are showed in Table I (averages). One can see that SSGA seems to be faster than PTA, although it perform more calls to the objective function.

**Table 1. Comparison of results**

	AVG Solution		Expected Solution	Error		AVG Evaluation / AVG Time (s)			
	PTA	SSGA		PTA	SSGA	PTA	SSGA		
Rosembrock(5)	0.012	0.351	0.001	0.011	0.350	486024.50	13.0	534000.00	5.7
Rastrigin(5)	0.001	0.001	0.001	0.000	0.000	178334.30	4.8	98992.70	1.1
Griewangk(5)	0.009	0.006	0.001	0.008	0.005	304628.80	8.1	540362.45	5.8
Langerman(5)	-0.963	-0.358	-1.400	0.437	1.042	103839.00	2.8	751024.00	8.0
Michalewicz(5)	-4.682	-4.687	-4.687	0.005	0.000	83140.80	2.2	92533.80	1.0
Schwefel(5)	-2094.672	-2094.908	-2094.914	0.243	0.006	359142.70	9.6	404562.20	4.3
Rosembrock(10)	0.102	6.640	0.001	0.101	6.639	703015.50	18.7	751000.00	8.0
Rastrigin(10)	0.007	0.001	0.001	0.006	0.000	142098.40	3.8	243424.20	2.6
Griewangk(10)	0.009	0.009	0.001	0.008	0.008	353668.30	9.4	574628.80	6.1
Langerman(10)	-0.004	-0.003	-1.400	1.396	1.397	422314.40	11.3	751000.00	8.0
Michalewicz(10)	-9.555	-7.693	-9.660	0.105	1.967	691504.10	18.4	751000.00	8.0
Schwefel(10)	-3942.296	-3952.952	-4189.829	247.533	236.877	368686.50	9.8	544419.20	5.8
						<b>349699.77</b>	<b>9.3</b>	<b>503078.94</b>	<b>5.4</b>

## 5. Conclusion

This work describes the preliminary results of an evolutionary approach based on population training in heuristics applied to minimization of unconstrained functions. The Population Training Algorithm (PTA) proposes a new concept of population training, working with a dynamic population, ranked by an adaptation coefficient. To compare its performance, it was built a steady-state genetic algorithm (SSGA) that combines some well-succeeded features of other genetic algorithms. Both algorithms are tested with some well-known test instances (functions) obtaining comparable good solutions. Some experiments confirm their good overall behavior seeking for local optimum. The PTA also has the reinforced effect of move away from unpromising solutions. Some further work is allowed on new heuristics for PTA and its adaptation to constrained problems.

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