Differential Evolution - A simple and efficient adaptive scheme for global optimization over continuous spaces

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Abstract

A new heuristic approach for minimizing possibly nonlinear and non differentiable continuous space functions is presented. By means of an extensive testbed, which includes the De Jong functions, it will be demonstrated that the new method converges faster and with more certainty than Adaptive Simulated Annealing as well as the Annealed Nelder&Mead approach, both of which have a reputation for being very powerful. The new method requires few control variables, is robust, easy to use and lends itself very well to parallel computation.

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Introduction

Problems which involve global optimization over continuous spaces are ubiquitous throughout the scientific community. In general, the task is to optimize certain properties of a system by pertinently choosing the system parameters. For convenience, a system's parameters are usually represented as a vector. The standard approach to an optimization problem begins by designing an objective function that can model the problem's objectives while incorporating any constraints. Especially in the circuit design community, methods are in use which do not need an objective function [1], [2], [3]. Although these methods can make formulating a problem simpler, they are usually inferior to techniques which make full use of an objective function. Consequently, we restrict ourselves to optimization methods which fully use the objective function. In most cases, the objective function is designed to transform the optimization problem into a minimization task. To this end, we will limit our investigation in the following to minimization problems.

When the objective function is nonlinear and non differentiable, direct search approaches are the methods of choice. The best known of these are the algorithms by Nelder&Mead [4], by Hooke&Jeeves [4], genetic algorithms [5], and evolutionary algorithms [6], [7] with the latter being truly continuous counterparts of genetic algorithms. At the heart of every direct search method is a strategy that generates variations of the parameter vectors. Once a variation is generated, a decision must be made whether or not to accept the newly derived parameters. All basic direct search methods use the greedy criterion to make this decision. Under the greedy criterion, a new parameter vector is accepted if and only if it reduces the value of the objective function. Although the greedy decision process converges fairly fast, it runs the risk of becoming trapped by a local minimum. Inherently parallel search techniques like genetic and evolutionary algorithms have some built-in safeguards to forestall misconvergence. By running several vectors simultaneously, superior parameter configurations can help other vectors escape local minima. Another method which can extricate a parameter vector from a local minimum is Simulated Annealing [8], [9], [10]. Annealing relaxes the greedy criterion by occasionally permitting an uphill move. Such moves potentially allow a parameter vector to climb out of a local minimum. As the number of iterations increases, the probability of accepting an uphill move decreases. In the long run, this leads to the greedy criterion. While all direct search methods lend themselves to annealing, it has mostly been used just for the Random Walk, which itself is the simplest case of an evolutionary algorithm [6]. Nevertheless, attempts have been made to anneal other direct searches like the method of Nelder&Mead [10] and genetic algorithms [8], [11].

Users generally demand that a practical optimization technique should fulfill three requirements. First, the method should find the true global minimum, regardless of the initial system parameter values. Second, convergence should be fast. Third, the program should have a minimum of control parameters so that it will be easy to use. In our search for a fast and easy to use "sure fire" technique, we developed a method which is not only astonishingly simple, but also performs extremely well on a wide variety of test problems. It is inherently parallel and hence lends itself to computation via a network of computers or processors. The basic strategy employs the difference of two randomly selected parameter vectors as the source of random variations for a third parameter vector. In the following, we present a more rigorous description of the new optimization method which we call Differential Evolution.



Problem Formulation

Consider a system with the real valued properties

$$g_{m}; m = 0, 1, 2, ..., P-1$$
 (1)

which constitute the objectives of the system to be optimized.

Additionally, there may be real valued constraints

$$g_{m}; m = P, P+1, ..., P+C-1$$
 (2)

which describe properties of the system that need not be optimized but neither shall be degraded. For example, one may wish to design a mobile phone with the dual objectives of maximizing the transmission power g_1 and minimizing the noise g_2 of the audio amplifier while simultaneously keeping the battery life g_3 above a certain threshold. The properties g_1 and g_2 represent objectives to be optimized whereas g_3 is a constraint. Let all properties of the system be dependent on the real valued parameters

$$x_{j}; j = 0, 1, 2, ..., D-1.$$
 (3)

In the case of the mobile phone the parameters could be resistor and capacitor values. For most technical systems realizability requires

$$x_j \in [x_{jl}, x_{jh}] \quad . \tag{4}$$

Usually, restrictions on the x_j will be incorporated into the collection g_m , $m \ge P$, of constraints. Optimization of the system means to vary the D-dimensional parameter vector

$$\underline{\mathbf{x}} = (\mathbf{x}_0, \, \mathbf{x}_1, \, \dots, \, \mathbf{x}_{D-1})^{\mathsf{I}} \tag{5}$$

until the properties g_m are optimized and the constraints g_m , m≥P, are met. An optimization task can always be reformulated as the minimization problem

min f_m(
$$\underline{x}$$
) (6)

where $f_m(\underline{x})$ represents the function by which the property g_m is calculated and its optimization or constraint preservation is represented as the minimization of $f_m(\underline{x})$. All functions $f_m(\underline{x})$ can be combined into a single objective function $z(\underline{x})$ [2], [12], which usually is computed either via the weighted sum

$$z(\underline{x}) = \sum_{m=0}^{P+C-1} w_m \cdot f_m(\underline{x})$$
(7)

or via with

$$z(\underline{x}) = \max(w_{m} \cdot f_{m}(\underline{x}))$$

$$w_{m} > 0.$$
(8)

The weighting factors w_m are used to define the importance associated with the different objectives and constraints as well as to normalize different physical units. The optimization task can now be restated as

The min-max formulation (8) and (10) guarantees that all local minima, the Pareto critical points, including the possibly multiple global minima, the Pareto points, can at least theoretically be found [2], [12]. For the objective function (7) and (10) this is true only if the region of realizability of \underline{x} is convex [1], [2], which in general does not apply in most technical problems.



(9)

The Method of Differential Evolution

Differential Evolution (DE) is a novel parallel direct search method which utilizes NP parameter vectors

$$\underline{x}_{i,G}, i = 0, 1, 2, ..., NP-1.$$
 (11)

as a population for each generation G. NP doesn't change during the minimization process. The initial population is chosen randomly if nothing is known about the system. As a rule, we will assume a uniform probability distribution for all random decisions unless otherwise stated. In case a preliminary solution is available, the initial population is often generated by adding normally distributed random deviations to the nominal solution $\underline{x}_{nom,0}$. The crucial idea behind DE is a new scheme for generating trial parameter vectors. DE generates new parameter vectors by adding the weighted difference vector between two population members to a third member. If the resulting vector yields a lower objective function value than a predetermined population member, the newly generated vector replaces the vector with which it was compared. The comparison vector can, but need not be part of the generation process mentioned above. In addition the best parameter vector $\underline{X}_{best,G}$ is evaluated for every generation G in order to keep track of the progress that is made during the minimization process.

Extracting distance and direction information from the population to generate random deviations results in an adaptive scheme with excellent convergence properties. We tried several variants of DE, the two most promising of which we subsequently present in greater detail.

Scheme DE1

Our first variant of DE works as follows: for each vector $\underline{x}_{i,G}$, i = 0, 1, 2, ..., NP-1, a trial vector \underline{v} is generated according to

$$\underline{\mathbf{v}} = \underline{\mathbf{x}}_{\mathbf{r}_1,\mathbf{G}} + \mathbf{F} \cdot (\underline{\mathbf{x}}_{\mathbf{r}_2,\mathbf{G}} - \underline{\mathbf{x}}_{\mathbf{r}_3,\mathbf{G}}),\tag{12}$$

with

$$r_1, r_2, r_3 \in [0, NP-1]$$
, integer and mutually different, and F > 0. (13)

The integers r_1 , r_2 and r_3 are chosen randomly from the interval [0, NP-1] and are different from the running index i. F is a real and constant factor which controls the amplification of the differential variation $(\underline{x}_{r_2,G} - \underline{x}_{r_3,G})$. Fig. 1 shows a two dimensional example that illustrates the different vectors which play a part in DE1.





Fig.1: Two dimensional example of an objective function showing its contour lines and the process for generating v in scheme DE1.

In order to increase the diversity of the parameter vectors, the vector

u_i

$$\underline{\mathbf{u}} = (\mathbf{u}_1, \mathbf{u}_2, \dots, \mathbf{u}_D)^{\mathrm{T}}$$
(14)

with

$$=\begin{cases} v_{j} & \text{for } j = \langle n \rangle_{D}, \langle n+1 \rangle_{D}, \dots, \langle n+L-1 \rangle_{D} \\ (x_{i,G})_{j} & \text{otherwise} \end{cases}$$
(15)

is formed where the acute brackets $\left< ~\right>_{\rm D}$ denote the modulo function with modulus D.

I.e. a certain seuence of the vector elements of \underline{u} are identical to the elements of \underline{v} , the other elements of \underline{u} acquire the original values of $\underline{x}_{i,G}$. Choosing a subgroup of parameters for mutation is similiar to a process known as crossover in evolution theory. This idea is illustrated in Fig. 2 for D=7, n=2 and L=3. The starting index n in (15) is a randomly chosen integer from the interval [0, D-1]. The integer L is drawn from the interval [0, D-1] with the probability $Pr(L=v) = (CR)^{V}$. $CR \in [0,1]$ is the crossover probability and constitutes a control variable for the DE1-scheme. The random decisions for both n and L are made anew for each trial vector \underline{v} .



Parameter vector containing the parameters x_i , j=0,1, ... , D-1

Fig. 2: Illustration of the crossover process for D=7, n=2 and L=3.

In order to decide whether the new vector \underline{u} shall become a population member of generation G+1, it will be compared to $\underline{x}_{i,G}$. If vector \underline{u} yields a smaller objective function value than $\underline{x}_{i,G}$, $\underline{x}_{i,G+1}$ is set to \underline{u} , otherwise the old value $\underline{x}_{i,G}$ is retained.

Scheme DE2

Basically, scheme DE2 works the same way as DE1 but generates the vector v according to

$$\underline{\mathbf{v}} = \underline{\mathbf{x}}_{i,G} + \lambda \cdot (\underline{\mathbf{x}}_{best,G} - \underline{\mathbf{x}}_{i,G}) + F \cdot (\underline{\mathbf{x}}_{r_2,G} - \underline{\mathbf{x}}_{r_3,G}), \qquad (16)$$

introducing an additional control variable λ . The idea behind λ is to provide a means to enhance the greediness of the scheme by incorporating the current best vector $\underline{x}_{\text{hest.G}}$. This feature can be useful for

non-critical objective functions. Fig. 3 illustrates the vector-generation process defined by (16). The construction of \underline{u} from \underline{v} and $\underline{x}_{i,G}$ as well as the decision process are identical to DE1.





Fig.3: Two dimensional example of an objective function showing its contour lines and the process for generating <u>v</u> in scheme DE2.

Competing minimization methods

In order to compare the DE method with other global minimizing strategies, we looked for approaches where the source code is readily available, which are known to be powerful and which are capable of coping with nonlinear and non differentiable functions. Two methods in particular piqued our interest. The first was the annealed version of the Nelder&Mead strategy (ANM) [10] which is appealing because of its adaptive scheme for generating random parameter deviations. When the annealing part is switched off, a fast converging direct search method remains which is especially useful for non-critical objective functions. The basic control variables in ANM are T, the starting temperature, TF, the temperature reduction factor and NV, the number of random variations at a given temperature level.

The second method of interest was Adaptive Simulated Annealing (ASA) [8] which claims to converge very quickly and to outperform genetic algorithms on the De Jong test suite [9]. Although ASA provides more than a dozen control variables, it turned out that just two of them, TEMPERATURE_RATIO_SCALE (TRS) and TEMPERATURE_ANNEAL_SCALE (TAS), had significant impact on the minimization process. We will compare both ANM and ASA to DE1 and DE2. During our research we also wrote an annealed version of the Hooke&Jeeves method [5] and tested two Monte Carlo methods [3] one of which used NP parallel vectors and the differential mutation scheme of DE. Although these approaches all worked, they quickly turned out not to be competitive.

The Testbed

Our function testbed contains the De Jong test functions as presented in [9] plus some additional functions which present further distinctive difficulties for a global minimizer:



1) First De Jong function (sphere)

$$f_1(\underline{x}) = \sum_{j=0}^{2} x_j^2; \qquad x_j \in [-5.12, 5.12]$$
(17)

 $f_1(\underline{x})$ is considered to be a very simple task for every serious minimization method. The minimum is $f_1(\underline{0}) = 0$.

2) Second De Jong function (Rosenbrock's saddle)

$$f_{2}(\underline{x}) = 100 \cdot (x_{0}^{2} - x_{1})^{2} + (1 - x_{0})^{2}; \qquad x_{j} \in [-2.048, 2.048]$$
(18)

Although $f_2(\underline{x})$ has just two parameters, it has the reputation of being a difficult minimization problem. The minimum is $f_2(\underline{1})=0$.

3) Third De Jong function (step)

$$f_{3}(\underline{x}) = 30. + \sum_{j=0}^{4} \lfloor x_{j} \rfloor;$$
 $x_{j} \in [-5.12, 5.12]$ (19)

For $f_3(\underline{x})$ it is necessary to incorporate the constraints imposed on the x_j into the objective function. We implemented this according to the min-max formulation (8). The minimum is $f_3(\underline{-5}\underline{\epsilon})=0$ where $\epsilon \in [0,0.12]$. The step function exhibits many plateaus which pose a considerable problem for many minimization algorithms.

4) Modified fourth De Jong function (quartic)

$$f_{4}(\underline{x}) = \sum_{j=0}^{29} (x_{j}^{4} \cdot (j+1) + \eta); \qquad x_{j} \in [-1.28, 1.28]$$
(20)

This function is designed to test the behavior of a minimization algorithm in the presence of noise. In the original De Jong function, η is a random variable produced by Gaussian noise having the distribution N(0,1). According to [9], this function appears to be flawed as no definite global minimum exists. In response to the problem, we followed the suggestion given in [9] and chose η to be a random variable with uniform distribution and bounded by [0,1). In contrast to the original version of De Jong's quartic function, we also included η inside the summation instead of just adding η to the summation result. This change makes $f_4(\underline{x})$ more difficult to minimize. The functional minimum is $f_4(\underline{0}) \leq 30 \cdot E[\eta] = 15$, where $E[\eta]$ is the expectation of η .

5) Fifth De Jong function (Shekel's Foxholes)

$$f_{5}(\underline{x}) = \frac{1}{0.002 + \sum_{i=0}^{24} \frac{1}{i + \sum_{j=0}^{1} (x_{j} - a_{ij})^{6}}} ; \qquad x_{j} \in [-65.536, 65.536]$$
(21)

with
$$a_{i0}=\{-32, -16, 0, 16, 32\}$$
 for $i = 0, 1, 2, 3, 4$ and $a_{i0}=a_{i \mod 5, 0}$
as well as $a_{i1}=\{-32, -16, 0, 16, 32\}$ for $i = 0, 5, 10, 15, 20$ and $a_{i1}=a_{i+k, 1}, k=1, 2, 3, 4$
The global minimum for this function is $f_{6}(-32, -32) \cong 0.998004$.

6) Corana's parabola [8], [13]

$$f_{6}(\underline{x}) = \sum_{j=0}^{3} \begin{cases} 0.15 (z_{j} - 0.05 \operatorname{sgn}(z_{j}))^{2} \cdot d_{j} & \text{if } |x_{j} - z_{j}| < 0.05 \\ d_{j} \cdot x_{j}^{2} & \text{otherwise} \end{cases}; x_{j} \in [-1000, 1000]$$
(22)
with $z_{j} = \left\lfloor \left| \frac{x_{j}}{0.2} \right| + 0.49999 \right\rfloor \cdot \operatorname{sgn}(x_{j}) \cdot 0.2$
and $d_{j} = \{1, 1000, 10, 100\}$

 $f_6(\underline{x})$ defines a paraboloid whose axes are parallel to the coordinate axes. It is riddled with a set of holes that increase in depth the closer one approaches the origin. Any minimization algorithm that goes strictly downhill will almost always be captured by the holes. The minimum here is $f_6(\underline{x}) = 0$, with $|x_i| < 0.05$, j=0,1,2,3.

7) Griewangk's function [14]

$$f_{7}(\underline{x}) = \sum_{j=0}^{9} \frac{x_{j}^{2}}{4000} - \prod_{j=0}^{9} \cos\left(\frac{x_{j}}{\sqrt{j+1}}\right) + 1; \qquad x_{j} \in [-400, 400]$$
(23)

Like test function $f_6(\underline{x})$, $f_7(\underline{x})$ has many local minima so that it is very difficult to find the true minimum $f_7(\underline{0}) = 0$.

8) Zimmermann's problem [15]

$$f_8(\underline{x}) = 9 - x_0 - x_1;$$
 $x_j > 0, j=1,2$ (24)

with
$$(x_0 - 3)^2 + (x_1 - 2)^2 \le 16$$
 (25)

and
$$\mathbf{x}_0 \cdot \mathbf{x}_1 \le 14$$
 (26)

Finding the minimum $f_8(7,2)=0$ poses a special problem, because the minimum is located at the corner of the constrained region defined by (24), (25) and (26).

9) Polynomial fitting problem

$$f_{9}(\underline{\mathbf{x}}, \mathbf{z}) = \sum_{j=0}^{2k} \mathbf{x}_{j} \cdot \mathbf{z}^{j} \text{ , k integer and >0,}$$
(27)

is a polynomial of degree 2k in z with the coefficients x_j such that

$$f_{9}(\underline{x}, z) \in [-1, 1] \qquad \text{for} \quad z \in [-1, 1]$$
(28)

and

$$f_9(\underline{x}, z) \ge T_{2k}(1.2)$$
 for $z = \pm 1.2$ (29)

with $T_{2k}(z)$ being a Chebychev Polynomial of degree 2k. The Chebychev Polynomials are defined recursively according to the difference equation $T_{n+1}(z) = 2z \cdot T_n(z) - T_{n-1}(z)$, n integer and > 0, with the initial conditions $T_0(z)=1$ and $T_1(z)=z$. The solution to the polynomial fitting problem is, of course, $f_9(\underline{x}, z) = T_{2k}(z)$, a polynomial which oscillates between -1 and 1 when its argument z is between -1 and 1. Outside this "tube" the polynomial rises steeply in direction of high positive ordinate values. The polynomial fitting problem has its roots in electronic filter design [16] and



challenges an optimization procedure by forcing it to find parameter values with grossly different magnitudes, something very common in technical systems. In our test suite we employed

$$\Gamma_{8}(z) = 1 - 32z^{2} + 160z^{4} - 256z^{6} + 128z^{8}$$
(30)

with

$$T_{s}(1.2) \cong 72.6606669$$
 (31)

as well as

$$T_{16}(z) = 1 - 128z^{2} + 2688z^{4} - 21504z^{6} + 84480z^{8} -$$

$$180224z^{10} + 212992z^{12} - 131072z^{14} + 32768z^{16}$$

$$T_{16}(1.2) \cong 10558.1450229.$$
(33)

with

and used the weighted sum (7) of squared errors in order to transform the above constrained optimization problem into an objective function to be minimized. The starting values for the parameters were drawn randomly from the interval [-100,100] for (30), (31) and [-1000,1000] for (32), (33).

Test Results

We tried to optimize each of the four algorithms by experimenting to find the control settings which provided fastest and smoothest convergence. Table I contains our choice of control variable settings for each minimization algorithm and each test function along with the averaged number of function evaluations (nfe) which were required to find the global minimum.

f _i (<u>x</u>)		ANM				ASA			DE1				DE2 (F=1)			
i	Т	TF	NV	nfe	TRS	TAS	nfe	NP	F	CR	nfe	NP	λ	CR	nfe	
1	0	n.a.	1	95	1.10 ⁻⁵	10	397	10	0.5	0.3	490	6	0.95	0.5	392	
2	0	n.a.	1	106	1.10 ⁻⁵	10000	11275	6	0.95	0.5	746	6	0.95	0.5	615	
3	300	0.99	20	90258	1·10 ⁻⁷	100	354	10	0.8	0.3	915	20	0.95	0.2	1300	
4	300	0.98	30	-	1.10 ⁻⁵	100	4812	10	0.75	0.5	2378	10	0.95	0.2	2873	
5	3000	0.995	50	-	1.10 ⁻⁵	100	1379	15	0.9	0.3	735	20	0.95	0.2	828	
6	5·10 ⁶	0.995	100	-	1.10 ⁻⁵	100	3581	10	0.4	0.2	834	10	0.9	0.2	1125	
7	10	0.99	50	-	1.10 ⁻⁵	0.1	-	30	1.	0.3	22167	20	0.99	0.2	12804	
8	5	0.95	5	2116	1.10 ⁻⁶	300	11864	10	0.8	0.5	1559	10	0.9	0.9	1076	
9(k=4)	100	0.95	40	(391373)	1.10 ⁻⁶	1000	-	30	0.8	1	19434	30	0.6	1.0	14901	
9(k=8)	5·10 ⁴	0.995	150	-	1.10 ⁻⁸	700	-	100	0.65	1	165680	80	0.6	1.0	254824	

 Table I:
 Averaged number of function evaluations (nfe) required for finding the global minimum. A hyphen indicates misconvergence and n.a. stands for "not applicable".



If the corresponding field for the number of function evaluations contains a hyphen, the global minimum could not be found. If the number is enclosed in parentheses, not all of the test runs provided the global minimum. We executed ten test runs with randomly chosen initial parameter vectors for each test function and each minimization.

When the global minimum was 0, we defined the minimization task to be completed once the final value was obtained with an accuracy better than 10^{-6} . For $f_4(\underline{x})$, we chose a value less than 15 to indicate the global minimum and a value less than 0.998004 in the case of $f_5(\underline{x})$.

Conclusion

The Differential Evolution method (DE) for minimizing continuous space functions has been introduced and shown to be superior to Adaptive Simulated Annealing (ASA) [8] as well as the Annealed Nelder&Mead approach (ANM) [10]. DE was the only technique to converge for all of the functions in our test function suite. For those problems where ASA or ANM could find the minimum, DE usually converged faster, especially in the more difficult cases. Since DE is inherently parallel, a further significant speedup can be obtained if the algorithm is executed on a parallel machine or a network of computers. This is especially true for real world problems where computing the objective function requires a significant amount of time.

Despite these already promising results, DE is still in its infancy and can most probably be improved. Further research might include a mathematical convergence proof like the one that exists for Simulated Annealing. A theoretically sound analysis to determine why DE converges so well would also be of great interest. Whether or not an annealed version of DE, or the combination of DE with other optimization approaches is of practical use, is still unanswered. Finally, it is important for practical applications to gain more knowledge on how to choose the control variables for DE.



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