

RESEARCH ARTICLE

Self-adaptive combination of global tabu search and local search for nonlinear equations

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(v1.0 released August 2009)

Solving systems of nonlinear equations is a very important task since the problems emerge mostly through the mathematical modeling of real problems that arise naturally in many branches of engineering and in the physical sciences. The problem can be naturally reformulated as a global optimization problem. In this paper, we show that a self-adaptive combination of a metaheuristic with a classical local search method is able to converge to some difficult problems that are not solved by Newton-type methods.

Keywords: Nonlinear equations; Metaheuristic; Tabu search; Hooke and Jeeves method;

AMS Subject Classification: 90C15; 90C56; 90C30

1. Introduction

In this paper, we consider solving the nonlinear system of equations

$$F(x) = 0, \quad F(x) = (f_1(x), f_2(x), \dots, f_n(x))^T \quad (1)$$

where $F : \Omega \subset \mathbb{R}^n \rightarrow \mathbb{R}^n$, $\Omega \equiv [l, u] = \{x : l_i \leq x_i \leq u_i, i = 1, \dots, n\}$ and the functions $f_i(x), i = 1, 2, \dots, n$ are continuously differentiable, using a combination of a metaheuristic with a classical local search method. Some problems in engineering, chemistry, physics, medicine and even economic areas, aim at determining the roots of a system of equations. In general, these problems are nonlinear and difficult to solve. The most famous techniques to solve nonlinear equations are based on the Newton's method [6, 10, 14, 18, 38, 45]. They require analytical or numerical first derivative information. Newton's method is the most widely used algorithm for solving nonlinear systems of equations. It is computationally expensive, in particular if n is large, since the Jacobian matrix and the solution of a system of linear equations are required at each iteration. The Quasi-Newton methods use less expensive iterations than Newton, but their convergence properties are not very different. In general, Quasi-Newton methods avoid either the necessity of computing derivatives, or the necessity of solving a full linear system per iteration or both tasks [39]. In [19], a new technique for solving systems of nonlinear equations reshaping the system as a multiobjective optimization problem is proposed. The authors applied a technique of evolutionary computation to solve

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the problem obtained after the change. In [20], the authors propose techniques for computing all the multiple solutions in nonlinear systems. Another technique to solve systems of nonlinear equations is presented in [25], where a heuristic continuous global optimization GRASP is applied. A genetic algorithm is proposed in [8]. Filled functions that guarantee convergence to global solutions are available in the literature to solve systems like (1), see for example [51] and some references therein cited. The problem of solving a nonlinear system of equations can be naturally formulated as a global optimization problem. Problem (1) is equivalent, in the sense that it has the same solution, to finding the globally smallest value of the l_2 -norm error function, usually known as merit function, related to solving the system of equations (1), defined by

$$\underset{x \in \Omega \subset \mathbb{R}^n}{\text{minimize}} M(x) \equiv \|F(x)\|_2. \quad (2)$$

Here, the global minimum, and not just a local minimum, of the objective function $M(x)$, in the set Ω , is to be found. We denote the global minimum by x^* . Classical local search methods, like Newton-type methods, have some disadvantages, when compared to global search methods. In particular

- the final solution is heavily dependent on the the initial approximation of the iterative process;
- they can be trapped in local minima;
- they require differentiable properties of all the equations of the nonlinear system.

We use the Example 1.1 below to show this local trap behavior.

Example 1.1 Consider the following system of nonlinear equations

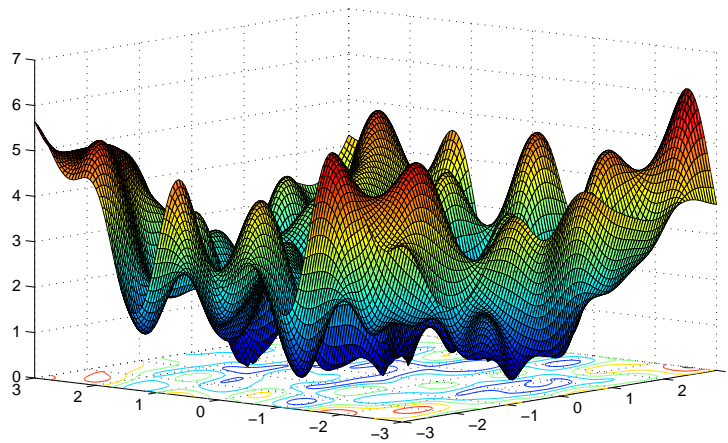
$$\begin{cases} f_1(x_1, x_2) \equiv x_1 - \sin(2x_1 + 3x_2) - \cos(3x_1 - 5x_2) = 0 \\ f_2(x_1, x_2) \equiv x_2 - \sin(x_1 - 2x_2) + \cos(x_1 + 3x_2) = 0 \end{cases} \quad (3)$$

and Figure 1 that shows the graphical representation of the l_2 -norm error function $M(x)$. The multi-modal nature of M makes the process of detecting a global minimum a difficult one. We solve Example 1.1 by `fsolve` from MATLABTM, using nine different initial approximations. In this MATLAB solver, the default trust-region dogleg algorithm with no analytical Jacobian is used [44]. The solver is able to converge to the solution only twice, although all the nine initial points are in the neighborhood of the solution. Table 1 shows the results obtained from MATLAB. The first column in the table presents the tested initial approximations and the second column lists the value of the output parameter ‘exitflag’ of MATLAB. The value ‘1’ means that the method converged to a root where the first-order optimality measure is less than a pre-specified tolerance, and ‘-2’ means that it converged to a point which is not a root and the sum-of-squares of the function values is greater than or equal to a pre-specified tolerance.

Thus, to be able to converge to a global solution, a global search strategy is required. The most important global search techniques invoke exploration and exploitation search procedures aiming at:

- i) diversifying the search in all the search space;
- ii) intensifying the search in promising areas of the search space.

Local optimization techniques guarantee globality only under certain convexity assumptions. Nonconvex problems exhibit multiple global and local (nonglobal) solutions and are more efficiently solved by global optimization methods. Preventing

Figure 1. Graphical representation of $M(x)$, from Example 1.1Table 1. Solutions obtained by `fsolve` from MATLAB for different initial approximations, for Example 1.1.

Initial approximation	'exitflag'	(f_1, f_2) at solution	n. of iterations	n. function evaluations
(0, 0)	1	(-4.1e-13, -2.4e-13)	5	18
(1, 1)	1	(3.6e-9, 2.3e-9)	6	21
(0, 1)	-2	(-3.0e-2, 9.4e-1)	27	60
(2, 2)	-2	(1.1e-1, 8.8e-1)	31	70
(-1, 1)	-2	(-5.5e-1, 6.5e-1)	55	130
(1, -1)	-2	(1.0e-2, -4.0e-2)	24	57
(-1, -1)	-2	(-5.2e-1, 1.0e-1)	31	74
(2, -2)	-2	(-1.6e-1, -5.7e-1)	32	71
(-2, -2)	-2	(-1.7e-1, -1.5e00)	34	77

premature convergence to a local solution, while trying to locate a global optimum, is an important property of global solution methods. Research concerning the problem of finding the global optimum of a continuous objective function over a compact convex set started in the early 1970's. A well-established classification of solution methods for global optimization defines two classes: the exact methods and the approximate ones. The reader is referred to papers with reviews on advances in global optimization [1, 13, 46]. Exact methods for global optimization are guaranteed to find an optimal solution within a problem dependent run time. Thus, they are able to solve a problem with a required accuracy in a finite number of steps. A theoretical analysis of convergence to a global optimum may be provided. On the other hand, approximate methods are not guaranteed to find an optimal solution although they are often able to find very good solutions in a reasonable time. Most approximate methods are stochastic. Exact methods are often known as deterministic. Unlike the stochastic methods, the outcome of a deterministic algorithm does not depend on pseudo random variables. The design of a deterministic method relies on the mathematical attributes of the optimization problem, therefore the performance depends heavily on the structure of the problem and the complexity grows very fast with problem's dimension [23, 37]. Popular deterministic approaches have been proposed within the Branch-and-Bound framework to solve certain types of nonconvex problems, see for example [3]. Other deterministic methods use concepts of DIviding RECTangle [29] and Interval Analysis. In the interval-based methods, the interval arithmetic is used to update variable bounds and compute bounds for the involved function values [23].

Another interesting idea which has been under discussion in the global optimization area, since the 1980's, uses auxiliary functions, namely Tunneling and Filled functions. The objective function is transformed into an auxiliary function using previously located local minimizers. To escape from local minimizers, gradient-based methods can then be used to descend from the local to a global one [54].

Stochastic methods rely on probabilistic elements, either in the problem data or in the algorithm itself, or in both. Compared with deterministic methods, the implementation of stochastic algorithms is often easier. In general, stochastic methods require no structural information about the problem at hand. For a challenging class of global optimization problems, the so-called black-box optimization problems, no structure is known and used when targeting the global solution. The convergence proofs for this type of methods involve the use of probability theory. Stochastic algorithms are at most able to provide a probabilistic convergence guarantee, for instance, convergence in mean square [28, 48], convergence in probability, or convergence with probability one [27]. In practice, there is no guarantee that the obtained solution is actually the global one, or by how far the algorithm has failed to locate the true global solution.

A stochastic method is based on random searches that use selected heuristics to promote the search over the feasible set and guide the choice of the most promising candidate solutions. The general consensus about the word heuristic is that this is a procedure based on commonsense rules aiming at increasing the probability of solving a specific problem and providing an approximate solution with no guarantee that it is close to the optimal solution. Although heuristics are closely associated with random search techniques, there are also deterministic heuristic methods. A selected sequence of well-known stochastic methods, in chronological order, follows: Evolution Strategy (1965), Genetic Algorithm (1975), Scatter Search (1977), Simulated Annealing (1983), Tabu Search (1986), Ant Colony Optimization (1992), Particle Swarm Optimization (1995), Differential Evolution (1997), Harmony Search (2001), Electromagnetism-like Mechanism (2003), Artificial Bee Colony (2005), Artificial Fish Swarm Optimization (2005). They are mainly inspired by nature or by evolutionary and swarm intelligence theories. A survey on stochastic methods is presented in the textbook [53].

Recent hybrid algorithms aim at combining a heuristic algorithm with a local search operator, either a deterministic or a random search, to enhance its exploitation ability [5, 21, 22, 47, 49]. A Multistart method is a hybrid, where local searches are performed starting from randomly generated points. The idea of clustering paired with multistart that appeared in the 1970's with great success aims at avoiding to perform a local search from another point that is closely associated with a found optimum solution [50]. In recent years, another type of hybrid strategy adds a second heuristic to the basic heuristic's design, for example, the Ant Colony System and the Tabu Search [30], the Harmony Search cooperating with Differential Evolution [36] or the integration of Scatter Search and Tabu Search [12].

A well-known class of global search techniques, the metaheuristics, use random procedures that invoke artificial intelligence tools and simulate nature behaviors. The word 'metaheuristics' is used to describe all approximate methods that combine basic heuristic methods into higher level frameworks to explore the search space in an efficient and effective manner. In the last three decades, these heuristics have proven to be computationally successful in solving combinatorial problems as well as continuous global optimization problems. Thus, the practical advantages of metaheuristics are their effectiveness and general applicability. Due to their random features, metaheuristics have, in general, slow convergence since they may fail to detect promising search directions in the neighborhood of a global minimum.

There are two classes of metaheuristics. A population-based heuristic that defines and maintains throughout the iterative process a set of solutions. The most known population-based heuristic is the Genetic Algorithm [17]. A point-to-point heuristic defines just one solution at the end of each iteration which will be used to start the next iteration. Simulated Annealing [21] and Tabu Search [15] are two examples of point-to-point methods. The Tabu Search (TS) is a metaheuristic developed primarily for solving combinatorial problems [15, 16]. The TS introduced by Cvijović and Klinowski [9] for continuous optimization guides the local search out of local optima and has the ability to explore new regions. It is an iterative procedure that maintains a list of the movements most recently made, avoiding in subsequent iterations the execution of movements that lead to solutions already known to have been visited. Usually, the slow convergence of TS is overcome by incorporating a classical local search strategy into the main algorithm. In general, this type of hybridization occurs in the final stage of the iterative process when the solution is in the vicinity of the global solution. An example of such method is presented in [22]. The therein proposed method, called Directed Tabu Search (DTS), uses strategies, like the Nelder-Mead method [43] and the Adaptive Pattern Search [21], to direct a tabu search.

This paper aims at assessing the performance of the metaheuristic tabu search when solving a system of nonlinear equations (1), using the function $M(x)$ as a measure of the progress of the algorithm towards the solution. According to the formulation (2), this means that the fitness of each trial solution x is assessed by evaluating the merit function M at x . And, a solution x is better than y if $M(x) < M(y)$. In this paper, and due to the reported success when solving global optimization problems of the form (2), the DTS variant of the tabu search is extended to be able to solve nonlinear systems of equations. In particular, we aim at analyzing the behavior of tabu search type methods when solving some difficult problems that are not solved by Newton-type methods.

Furthermore, we propose a new algorithm that combines DTS and a local search method in order to accelerate convergence to the solution. The issue related with the condition that defines the choice between the exploration and exploitation phases of the algorithm is also addressed with new self-adaptive weight factors. The numerical results show the goodness of the proposed self-adaptive strategies.

The organization of the paper is the following. Section 2 briefly describes the Directed Tabu Search method, Section 3 overviews the classical local search method, known as Hooke and Jeeves (HJ), and Section 4 presents the proposed combined DTS and HJ searches algorithm, and the new self-adaptive strategies to choose between the exploration and exploitation phases. The numerical results and their discussion are included in Section 5.

2. The metaheuristic tabu search

2.1 Basic tabu search

TS is an iterative process which operates in the following way. The algorithm starts with a randomly generated initial solution, x , and by applying pre-defined moves in its neighborhood, it generates a set \mathcal{Y} of solutions. The objective function to be minimized is evaluated at all solutions in \mathcal{Y} , and the best of all, y^{best} , becomes the current solution, $x \leftarrow y^{best}$ (even if it is worse than x). Accepting uphill moves, the algorithm avoids to get trapped in a local minimum. The previous procedure is repeated until a given stopping condition is reached. Further, the algorithm also stops when the solution does not improve for a specified number of iterations. To

avoid cycling, since a point already visited may be generated again, a set of points already visited are stored in a list, called Tabu List (TL). The solutions in \mathcal{Y} that belong to the TL are eliminated. This TS structure is known as short-term memory TS. The use of this type of flexible memory turns out to be advantageous, since the method is able to keep diversity, like population-based methods, in contrast to the rigid structures, like those used in Branch-and-Bound methods, or the lack of memory present in the Simulated Annealing method [7]. To improve performance, long-term memory TS structures have been proposed to record important attributes like the elite and the frequently visited solutions. The Directed Tabu Search [22] implemented in this paper for solving nonlinear systems of equations contains long-term memory structures.

2.2 Directed tabu search

The Directed Tabu Search method of Hedar and Fukushima [22] uses direct search methods in order to stabilize the search especially in the vicinity of a local minimum. Two variants of the DTS are therein proposed: one is based on the Nelder-Mead (NM) method, as a local search, inside the exploration step of the algorithm, and the other uses the Adaptive Pattern Search (APS) strategy in the exploration step. Furthermore, the Kelley's modification of the NM method [31] is still used in the therein called intensification search, in the final stage of the process. We note that the DTS method can be classified as a multi-start method. The multi-start methods are powerful search procedures to guide both global exploration and local search. The DTS method is based on three main procedures: exploration, diversification and intensification search. The structure of the DTS is shown below in Algorithm 2.1. (See details in [22].)

Algorithm 2.1 DTS algorithm

Require: randomly generated x^0

- 1: **while** the stopping criteria are not reached **do**
 - 2: Exploration search procedure
 - 3: Neighborhood search
 - 4: Local search
 - 5: Solution update
 - 6: Diversification search procedure
 - 7: **end while**
 - 8: Intensification search procedure
-

The main loop (outer cycle) of the DTS method, consisting of the exploration and diversification search procedures, begins with an initial randomly generated solution, but other initial approximation may be provided.

2.2.1 Exploration search

The exploration search aims to explore the search space Ω . It uses direct search methods as neighborhood search and local search strategies to generate trial points. These may be based on either the simplex method of NM or on the APS strategy. Here, we use the APS variant, as described in [21].

The cycling procedure is prevented not only with the standard Tabu List but also with the inclusion of two novel concepts of tabu regions. The DTS method implements four TS memory elements: The multi-ranked Tabu List, the Tabu Region, the Semi-Tabu Region and the Visited Region List. They are long-term memory

structures and are very important since they allow the method in the diversification search and intensification search procedures to behave as an intelligent search technique.

2.2.2 Diversification search

A diversification procedure aims to generate a new initial trial point outside the visited regions. The information stored in the Visited Region List (VRL) is used to direct the search towards new regions. This VRL serves as a diversification tool, with the aim of diversifying the search for areas that have not been visited in the search space.

2.2.3 Intensification search

When one of the best obtained trial solutions is sufficiently close to a global minimum, or its value has not been changed for a specified number of iterations, then the intensification search procedure is applied, at the final stage of the algorithm, to refine the best solution visited so far. In this case, DTS uses the Kelley's modification of the Nelder-Mead method [31] and [32]. A solution still closer to the global minimum is then obtained.

3. Hooke and Jeeves local search method

The derivative-free method, known as Hooke and Jeeves method, is a deterministic local pattern search method that performs, at each iteration j , a series of exploratory moves along the coordinate axes around a current approximation, x^j , in order to find a new approximation $x^{j+1} = x^j + \Delta^j s^j$, with a lower merit function value while maintaining the approximation inside the set Ω . Here, the index j is used for the iteration counter of this inner iterative process. The scalar Δ^j represents the step length and the vector s^j determines the direction of the step. The step length is reduced whenever the previous iteration is unsuccessful, i.e., when no improvement in M is obtained, and it is maintained otherwise. The HJ method also performs a pattern move whenever a successful iteration is encountered. The vector $x^{j+1} - x^j$ defines a promising direction and a pattern move is then implemented, which means that the exploratory move is carried out around the trial point $x^{j+1} + (x^{j+1} - x^j)$, rather than around the current point x^{j+1} . Then, if the coordinate search is successful, the returned point is accepted as the new point; otherwise, the pattern move is rejected and the method reduces to a coordinate search around x^{j+1} . The reader is referred to [26] for the details concerning this classical local search method.

4. Combining global tabu search and local search

The herein proposed hybridization defines an algorithm that is able to combine two types of cycles: a global and a local one. The general idea is borrowed from the algorithm presented in [14] where a classical gradient-based Quasi-Newton nonmonotone strategy is used to solve nonlinear systems of equations. The global search cycle is carried out by a modified version of DTS method, where the HJ local search method is used, instead of the NM method, as the intensification search procedure of the DTS algorithm (see Step 8 of Algorithm 2.1). On the other hand, the local search cycle uses the HJ pattern search method alone [26] (see Section 3).

The most important issue here is to decide when to carry out a global exploration of the search space, or a local exploitation in the neighborhood of a good

approximation to the solution.

4.1 Merit function sufficient reduction

Our first proposal concerned with the condition that decides which cycle should be carried out, at each iteration k , depends on a sufficient reduction obtained in the merit function, when compared with the merit function value of the previous iteration. If a pre-specified sufficient reduction is verified, a local exploitation cycle is to be required, since a fast downhill progress has been detected. Thus a promising region seems to be found by the algorithm. Otherwise, the region does not look promising yet and an exploration cycle is more appropriate where a diversifying search is to be carried out looking for a promising region with a global minimum. Figure 2 shows the general iterative structure of the combined Global Tabu Search and Local Search (GTSLs) method. Details of the procedures are discussed below.

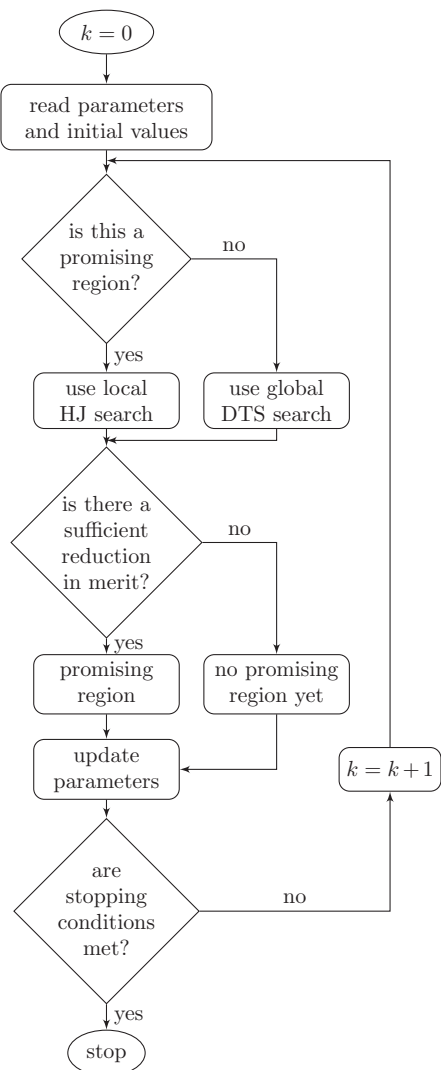


Figure 2. General flowchart of GTSLs method

In both global and local cycles of the algorithm, only an η_k -approximation x^{k+1} to the optimal solution is required, i.e., each search terminates when, at iteration k , the value of the merit function at the best solution found so far is less than η_k . The sequence of η_k values should decrease and approach zero, as k increases. The

argument is that, when the iterative process begins far from the solution of problem (1), there is no point in spending too much resources computing an approximation with high accuracy. However, as the process approaches the solution, highly accurate approximations are important to speed the overall process. Algorithm 4.1 summarizes the main steps of the proposed GTSLs algorithm.

Algorithm 4.1 GTSLs algorithm

Require: x^0 , $0 < \gamma_1, \gamma_2 < 1$, $0 < \eta^* \ll 1$, $k_{\max} > 0$; set $k = 0$, $\eta_0 = 1$, flag=0;

```

1: while  $M(x^k) > \eta^*$  and  $k \leq k_{\max}$  do
2:   if flag=1 then
3:     use local HJ search to compute an  $\eta_k$ -approximation  $x^{k+1}$ 
4:   else
5:     use global DTS search to compute an  $\eta_k$ -approximation  $x^{k+1}$ 
6:   end if
7:   if  $M(x^{k+1}) \leq \gamma_1 M(x^k)$  then
8:     set flag=1
9:   else
10:    set flag=0
11:  end if
12:  set  $k = k + 1$ , set  $\eta_k = \max\{\eta^*, \gamma_2 \eta_{k-1}\}$ 
13: end while

```

Condition $M(x^{k+1}) \leq \gamma_1 M(x^k)$ in the algorithm defines the sufficient reduction that we aim to observe in the merit function. The closer γ_1 is to one, the smaller is the required reduction. In order to guarantee that the progress has been along a downhill step and a promising region has been located, we define $\gamma_1 = 0.5$. The other parameters in the algorithm are set as follows: $\eta^* = 10^{-6}$ (accuracy of the solution) and $k_{\max} = 10n$. A successful run is registered when the algorithm stops due to the condition $M(x^k) \leq \eta^*$. Two values of γ_2 were tested, 0.1 and 0.5, see Section 5. Each iterative process, either in the local or in the global cycle of the algorithm, is called inner cycle, in contrast to the process indexed by k , called outer cycle.

4.2 Self-adaptive weight factors

Another approach is attempted to improve the efficiency of the GTSLs algorithm using a self-adaptive weight factor to check when the global exploration of the search space, or the local exploitation, is needed. At each iteration k , the weight factor is evaluated based on an ‘index’, I_k , defined by

$$I_k = \frac{M(x^0)}{M(x^k)} - 1 \quad (4)$$

that clearly shows the proximity of x^k to x^* . This ‘index’ changes according to the rate of the observed improvement on the merit function value. In the first iteration ($k = 0$), I_0 is zero but as x^k converges to x^* , I_k approaches infinity, since the required global optimum is 0. Since the self-adaptive weight factor w_k aims at checking which type of search is required, at iteration k , we use the idea of a transfer function [41]

$$w_k = \frac{1}{1 + \exp\left(-\frac{1}{\alpha I_k}\right)}, \quad (5)$$

where $0 < \alpha \leq 1$ is a constant, to define adequate weight values that vary as follows: $0.5 \leq w_k < 1$. A large weight factor, which emerges with a small I_k , means that $M(x^k)$ is very far from the merit function optimal value, and a global exploration search is required. On the other hand, a small w_k emerging from a large value of I_k means that a local exploitation search is needed since x^k is already near the optimal solution. The parameter α gives the speed of reduction in w . Figure 3 shows how w_k varies with α . Five values of α were used. The bigger the α is, the faster is the convergence to the lower bound of w . Preliminary experiments have shown that a moderate speed of reduction in w favors the efficiency of the algorithm. Another

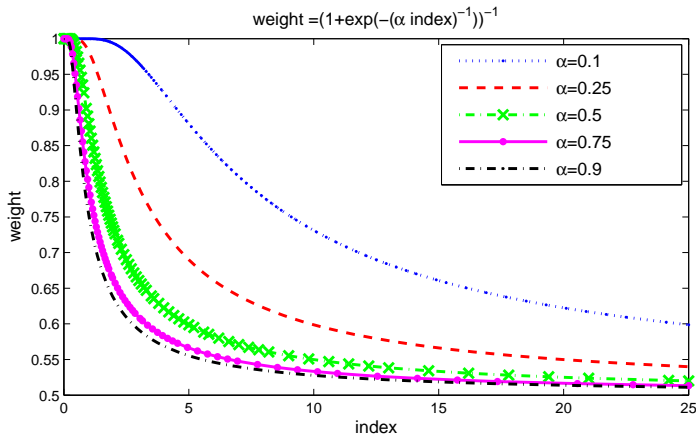


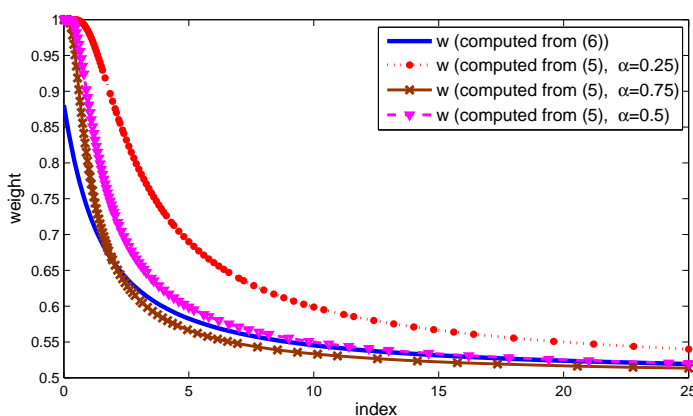
Figure 3. Weight factor w from (5) vs 'index' for five different values of α

issue is related with a reference value w_R , of the weight factor w , below which the exploration of the space is no longer necessary and the exploitation around the neighborhood of the best solution found so far is crucial for a highly accurate solution. From the numerical experiments done so far, the value 0.75 is a good choice. We remark that in this self-adaptive weight factor context, the condition on M in Step 7 of the Algorithm 4.1 is replaced by the following condition: $w_{k+1} \leq w_R$.

To overcome the need to define the parameter α in the formula for the weight factor (5), another idea borrowed from the work presented in [2] defines a weight factor ranging from 0.5 to 1 as follows:

$$w_k = 0.5 \left(1 + \tanh \left(\frac{M(x^k)}{M(x^0)} \right) \right). \quad (6)$$

The speed of reduction in the weight (6), as a function of the 'index' (4), behaves similarly to that of weight (5), when $0.5 \leq \alpha < 1$. Figure 4 depicts these behaviors. We now solve Example 1.1 to analyze the effect of parameter α in the performance of the algorithm. We compare the results of the proposed GTSLs algorithm with those of DTS and HJ used separately. The notation concerned with different versions of GTSLs is the following: GTSLs1 for the combined algorithm as presented in Algorithm 4.1, GTSLs2 for the adaptive version that uses (5) and GTSLs3 for the version using (6). See Table 2. Due to the stochastic nature of tabu search based algorithms, the problem was run 30 times and the best of the 30 obtained solutions is registered. In the table, ' k ' represents the number of outer (or main) iterations, ' k_{DTS+HJ} ' is the total number of inner iterations, ' k_{HJ} ' is the total number of iterations required by HJ alone, ' $n.f.ev.$ ' represents the average number of function evaluations, over the 30 runs, required to reach the presented solution, and ' M ' is the value of the merit function at the obtained registered solution. The solutions

Figure 4. Speed reduction comparison: weights w from (5) and (6)

reached by DTS and HJ do not have the required accuracy. We may conclude that the self-adaptive versions of the combined tabu search and local search algorithm - GTSLS2 and GTSLS3 - outperform GTSLS1.

Table 2. Results from GTSLS, DTS and HJ, for Example 1.1.

Algorithm	k	k_{DTS+HJ}	k_{HJ}	$n.f.ev.$	M	
GTSLS1	9	160	46	1186	5.0e-7	
GTSLS2	$(\alpha = 0.25, w_R = 0.75)$	7	91	1	414	3.3e-7
	$(\alpha = 0.50, w_R = 0.75)$	8	129	6	665	2.8e-7
	$(\alpha = 0.75, w_R = 0.75)$	11	188	6	985	8.8e-7
GTSLS3	$(w_R = 0.75)$	8	114	1	500	2.8e-7
	$(w_R = 0.60)$	8	122	6	645	8.2e-7
DTS	20	40	20	280	1.5e-3	
HJ	20			99	2.1e-2	

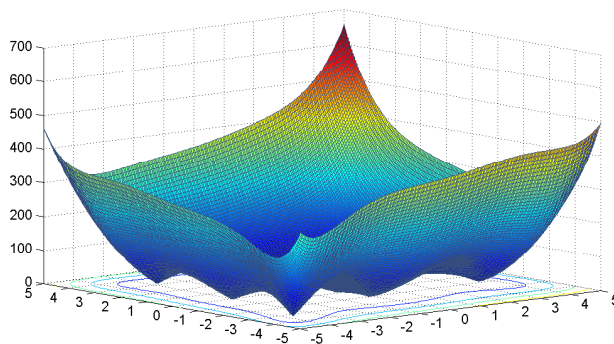
To further analyze the effect of both formulae (5) and (6) on the performance of the algorithm GTSLS, a second example is used.

Example 4.1 Consider the following system of nonlinear equations

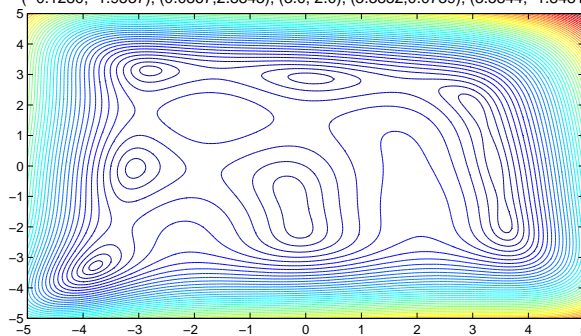
$$\begin{cases} f_1(x_1, x_2) \equiv 4x_1^3 + 4x_1x_2 + 2x_2^2 - 42x_1 - 14 = 0 \\ f_2(x_1, x_2) \equiv 4x_2^3 + 4x_1x_2 + 2x_1^2 - 26x_2 - 22 = 0 \end{cases} \quad (7)$$

where $-5 \leq x_1, x_2 \leq 5$ [51]. Figure 5 shows the graphical representation and the contours of $M(x)$ relative to this example. The merit function M has nine minima. The solutions are reported on the contour plot.

We test both versions GTSLS2 and GTSLS3, with three initial approximations, and compare with `fsolve` and the results in [51]. See Table 3. The information reported in this table is similar to that of Table 2 with the additional information concerned with the solution x^* . In the table, ‘-’ stands for unavailable information. GTSLS3 always converges to the same solution whatever the initial approximation, while GTSLS2 detects three solutions. The algorithm in [51] and the solver `fsolve` converge to three different solutions depending on the provided initial approximation.

(a) Plot of merit function $M(x)$

Solutions: $(-3.7793, -3.2832)$, $(-3.0730, -0.0814)$, $(-2.8051, 3.1313)$, $(-0.2709, -0.9230)$,
 $(-0.1280, -1.9537)$, $(0.0867, 2.8843)$, $(3.0, 2.0)$, $(3.3852, 0.0739)$, $(3.5844, -1.8481)$

(b) Contours of $M(x)$ Figure 5. Plot of $M(x)$ and contours, from Example 4.1Table 3. Results from self-adaptive GTSLs, `fsolve` and [51], for Example 4.1.

Algorithm	k	k_{DTS+HJ}	$n.f.ev.$	x^*	M
initial approximation $(-5, -3)$					
GTSLs2	6	68	380	$(-0.1280, -1.9537)$	$7.3e-7$
GTSLs3	8	116	566	$(-0.2708, -0.9230)$	$9.5e-7$
in [51]	3		-	$(-3.7793, -3.2832)$	$3.8e-7$
<code>fsolve</code>	5		18	$(-3.7793, -3.2832)$	$3.5e-10^\dagger$
initial approximation $(1, 3)$					
GTSLs2	7	84	449	$(3.5844, -1.8481)$	$6.04e-7$
GTSLs3	6	74	396	$(-0.2708, -0.9230)$	$7.3e-7$
in [51]	3		-	$(0.0867, 2.8843)$	$6.8e-7$
<code>fsolve</code>	4		15	$(0.0867, 2.8843)$	$7.1e-10^\dagger$
initial approximation $(2, 3)$					
GTSLs2	7	91	486	$(-0.2708, -0.9230)$	$2.5e-7$
GTSLs3	8	115	565	$(-0.2708, -0.9230)$	$6.6e-7$
in [51]	3		-	$(3.3852, 0.0739)$	$7.9e-7$
<code>fsolve</code>	5		18	$(3.0, 2.0)$	$5.8e-7^\dagger$

† - first-order optimality measure

4.3 Escaping from local minima

To further check whether the proposed algorithm is able to escape from local minima or from valleys that contain them, while searching for a global one, two small nonconvex optimization problems are selected from the literature (Examples 4.2 and 4.3 below [23]).

Example 4.2 This function is the well-known six-hump camelback function

$$f(x) = 4x_1^2 - 2.1x_1^4 + \frac{1}{3}x_1^6 + x_1x_2 - 4x_2^2 + 4x_2^4 \text{ with } \Omega = [-2, 2] \times [-2, 2]$$

which has six minimizers and the two global solutions are located at $(-0.089842, 0.71266)$ and $(0.089842, -0.71266)$ with a value of -1.0316 .

Example 4.3 The bi-spherical function is nonsmooth and has one global minimizer at $(1, 0)$ with $f = 0$ and a local minimizer at $(-1, 0)$ with $f = 0.1$:

$$f(x) = \min \{(x_1 - 1)^2, (x_1 + 1)^2 + 0.1\} + x_2^2 \text{ with } \Omega = [-2, 2] \times [-1, 1].$$

By allowing uphill moves, the exploration nature of the DTS algorithm searches beyond a local minimum. Furthermore, by preventing the algorithm from visiting again the points previously evaluated, a better exploration of the problem space can be enforced. We illustrate the behavior of the GTSLs3 algorithm on the two multi-modal functions over convex closed feasible sets and compare with the solver `fmincon` from MATLAB. Figure 6 contains the contours of both functions and depicts the convergence behavior starting from four different initial approximations.

Figure 6(a) shows the convergence behavior of GTSLs3 on Example 4.2 starting from the initial points, x^0 : $(2, 2)$, $(-2, 0)$, $(-1.7036, 0.7961)$ and $(1.2302, 0.1623)$. These two last points are local minima. The algorithm is able to explore all the feasible region, escape from the local minima and converge to global minima. Table 4 contains the results obtained by the GTSLs3 algorithm and `fmincon` from MATLAB for comparison. We may observe that the solver `fmincon` could not escape from the two local minima, converges just once to a global minimum and identifies a saddle point.

When starting from the initial points $(0, 0)$, $(-1, 1)$, $(-2, -1)$ and $(-0.5, 0)$, the GTSLs3 algorithm is able to converge to the global minimum of Example 4.3 in all cases, while `fmincon` locates the local one in two cases. See Table 4 for details. Figure 6(b) illustrates the iterated points obtained by GTSLs3.

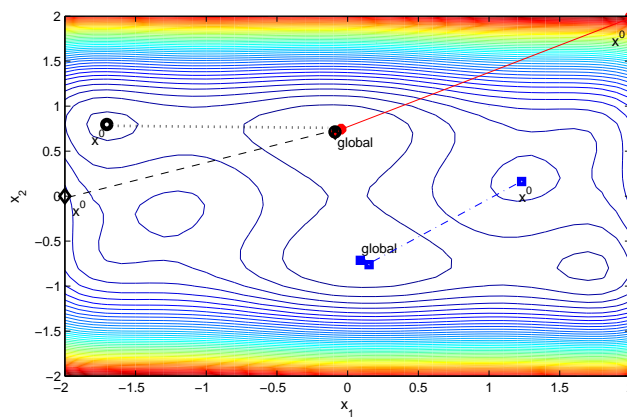
Table 4. Convergence results for Examples 4.2 and 4.3.

Ex.	x^0	GTSLs3 algorithm			fmincon from MATLAB	
		solution	f	$k(n.f.ev.)$	solution	f
4.2	$(2, 2)$	$(-0.0898, 0.7128)$	-1.03163	3 (60)	$(-0.1E-6, 0.01E-6)^\dagger$	$8E-14$
	$(-2, 0)$	$(-0.0899, 0.7127)$	-1.03163	4 (103)	$(-0.0898, 0.7127)$	-1.03163^\ddagger
	$(-1.7036, 0.7961)$	$(-0.0899, 0.7128)$	-1.03163	4 (122)	$(-1.7036, 0.7961)$	-0.2155^\S
	$(1.2302, 0.1623)$	$(0.0897, -0.7127)$	-1.03163	4 (101)	$(1.2302, 0.1623)$	2.4963^\S
4.3	$(0, 0)$	$(0.9993, 0)$	$4E-7$	2 (25)	$(1, 0)$	$7E-17^\ddagger$
	$(-1, 1)$	$(0.9996, -0.0003)$	$2E-7$	4 (108)	$(-1, 0)$	0.1^\S
	$(-2, -1)$	$(0.9997, -0.0001)$	$9E-8$	4 (93)	$(0.9999, -0.0001)$	$7E-9^\ddagger$
	$(-0.5, 0)$	$(0.9998, 0.0008)$	$6E-7$	3 (77)	$(-1, 0)$	0.1^\S

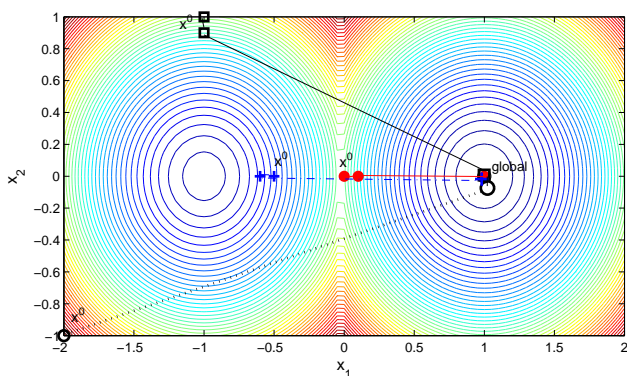
† - saddle point; ‡ - global minimum; § - local minimum.

5. Numerical results and discussion

To analyze the performance of the proposed combined Global Tabu Search and Local Search algorithm, we selected, coded in MATLABTM, and solved by the version GTSLs3, 99 test problems from the literature. For comparative purposes, the problems were also solved by `fsolve` from MATLAB, Hooke and Jeeves method,



(a) Iterated points in Example 4.2



(b) Iterated points in Example 4.3

Figure 6. Iterated points from different initial approximations

and the extended version of the Directed Tabu Search method presented in [22]. The problems in our database are referred to as P1, P2, ..., P99. They represent systems of nonlinear equations of different sizes and complexity. P99 and P49 are Examples 1.1 and 4.1 respectively. The results of the numerical experiments were obtained in a personal computer with an AMD Turion 2.20 GHz processor and 3 GB of memory.

Since the computational effort required in each outer/main iteration, by the algorithms in comparison, is different, we choose to stop the algorithms when the number of function evaluations exceeds $100n^2$. Meanwhile, if a solution is found with a merit function value less or equal to η^* ($= 10^{-6}$) (tolerance for M at the best solution found so far) the algorithms stop. This way, we aim at analyzing and comparing the accuracy of the obtained solutions.

To resume the main achievements of our numerical experiments, we show in Table 5 the results of 22 problems (from the database of 99 problems) for which `fsolve` was not able to converge to a solution with a specified tolerance within $10n$ iterations or $100n$ function evaluations (values defined by default). The output parameter 'exitflag' is '-2' for P44 and P72 and is '0' for all the others. All problems were solved with a specific initial approximation: the null vector for P50 and P52, the vector of all ones for P56, P62, P63, P70, P77 and P84, and the initial vector

provided in the literature for the remaining problems. Although the literature does not provide the bounds $[l, u]$ for most problems, we selected a specific range for each problem, as shown in Table 6. The lower and upper limits are the same for all the components of the vector x .

For the methods HJ, DTS, GTSLS3 with $\gamma_2 = 0.1$ and GTSLS3 with $\gamma_2 = 0.5$, Table 5 shows the value of M (at the best solution found after $100n^2$ function evaluations) and the required number of outer/main iterations. When the parameter γ_2 is set to 0.1 in GTSLS3, the algorithm requires less iterations and attains in general higher accuracy results than with $\gamma_2 = 0.5$. This is expected since outer iterations based on a larger tolerance reduction provide solutions with higher accuracy although computationally more expensive. Overall the accuracy of the results are not yet as we expected. We then run GTSLS3 for a maximum of $10n$ outer iterations and obtained the solutions reported in the 12th column of the table identified as ' $M_{(k \leq 10n)}$ '. The results show some improvements but further research is still needed.

To avoid the search along the coordinates during the Hooke and Jeeves local exploitation search of the GTSLS algorithm, another local search procedure will be used in the near future. Since a componentwise search requires large amounts of function evaluations, the idea proposed in the random walk with direction exploitation method, recently applied to the stochastic differential evolution algorithm [36], will be considered.

In this paper, we show that nonlinear systems of equations can be effectively solved by implementing a global optimization method to the merit function, which represents the l_2 -norm error function related to the solving of the system of equations. The application of an extended version of the metaheuristic Directed Tabu Search, proposed in [22], for solving complex and difficult nonlinear systems of equations has been analyzed and tested. We also propose a novel tabu search-type algorithm that combines the extended DTS, for a global exploration search, with the local Hooke and Jeeves search algorithm, for the exploitation search procedure. At each iteration, the choice between implementing a global search or a local search relies on self-adaptive weight factors that depend on the rate of improvement on the merit function value. The numerical results allow us to conclude that the herein proposed self-adaptive GTSLS algorithm is able to converge to the solution of some difficult problems that were not successfully solved by Newton-type methods and outperforms both HJ local search and DTS algorithms.

Acknowledgments

The authors wish to thank an anonymous referee and the editor whose comments and suggestions helped to improve the paper.

This research has been supported by CIDEM (Centre for Research & Development in Mechanical Engineering, Portugal), FCT - Fundação para a Ciência e a Tecnologia (Portuguese Foundation for Science and Technology) Project PEst-OE/EME/UI0615/2011, FEDER COMPETE (Programa Operacional Fatores de Competitividade / Operational Programme Thematic Factors of Competitiveness) and FCT Project FCOMP-01-0124-FEDER-022674.

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Table 5.: Results from `fsolve`, HJ, DTS and self-adaptive GTLSLS3.

	n	fsolve		HJ		DTS		GTLSLS3, $\gamma_2 = 0.1$		GTLSLS3, $\gamma_2 = 0.5$		
		M	$n.f.eval$	M	k	M	k	M	k	M	k	
P6	2	2.5e-3	57	1.0e-4	30	2.8e-4	28	1.0e00	3	1.6e-4	1.0e00	3
P9	2	7.0e00	49	7.0e00	73	9.1e-4	28	8.0e-5	6	8.9e-7	3.0e-4	13
P25	10	1.4e+1	1001	8.8e-1	455	4.3e-1	291	1.0e-1	7	5.9e-2	8.8e-2	21
P26	3	6.6e-1	106	6.6e-1	47	1.7e-4	70	7.0e-7	7	5.9e-7	3.1e-5	15
P27	3	1.0e-1	109	7.9e-2	77	7.9e-2	67	2.0e-2	6	7.9e-2	8.4e-2	16
P44	33	2.2e00	1170	2.2e00	47	9.4e-1	1055	1.3e00	32	1.1e00	9.4e-1	40
P45	33	4.0e-1	3306	2.6e-1	77	1.6e00	883	2.7e-1	8	2.7e-1	2.7e-1	24
P50	2	2.5e-1	47	2.5e-1	37	2.5e-1	26	2.5e-1	3	2.5e-1	2.5e-1	4
P52	5	2.2e-3	261	3.7e-2	237	3.3e-1	137	1.1e-1	4	8.5e-5	5.6e-1	4
P53	6	2.8e00	427	2.8e00	296	1.4e-2	134	2.2e-3	6	1.0e-4	1.9e-2	15
P55	6	1.4e+2	385	1.4e+2	40	1.4e+2	66	1.1e+7	2	1.4e+2	7.8e+4	10
P56	8	4.9e-1	609	2.6e-6	190	3.7e-5	219	2.8e-6	7	2.8e-6	3.6e-6	20
P62	6	4.7e-2	379	7.0e12	5	1.2e10	50	1.2e14	1	5.9e+2	9.6e11	1
P63	6	6.2e-3	427	3.0e-4	277	4.1e-6	157	2.5e-6	12	7.4e-6	2.4e-7	20
P64	8	3.3e00	657	4.1e-5	99	4.8e-5	235	1.3e-5	13	8.0e-5	1.1e-5	22
P70	10	2.6e+1	981	2.1e+1	61	1.0e+1	288	2.1e+1	3	1.0e+1	3.6e+1	20
P72	51	2.7e00	1989	2.7e00	47	1.8e00	1611	6.6e-1	50	9.4e-1	6.6e-1	57
P77	2	4.6e00	45	4.6e00	36	7.4e-1	25	7.4e-1	7	7.4e-1	7.4e-1	14
P80	3	6.3e-2	124	1.6e-5	140	9.0e-4	69	9.3e-6	7	9.1e-6	1.7e-4	17
P84	3	5.0e-4	124	3e-21	24	3.9e-13	19	7.6e-9	2	2.3e-8	3.4e-8	1
P88	10	7.5e-1	66	3.9e-2	479	5.6e-2	288	1.6e-2	5	7.5e-4	1.9e-2	16
P96	2	2.8e00	47	1.0e-6	41	2.9e-5	27	6.9e-7	7	5.7e-7	1.9e-5	16

REFERENCES

Table 6. Characteristics of the problems.

	n	$[l_i, u_i]$	reference	Problem's name in cited paper
P6	2	[-10,10]	[42]	P3-Powell badly scaled function
P9	2	[-10,15]	[42]	P2-Freudenstein and Roth function
P25	10	[10,50]	[14]	D6-Shifted and augmented trigonometric function with an Euclidian sphere
P26	3	[-10,10]	[14]	D7-Diagonal of three variables premultiplied by a quasi-orthogonal matrix
P44	33			
P27	3	[-10,10]	[14]	D8-Diagonal of three variables premultiplied by an orthogonal matrix, combined with inverse trigonometric function
P45	33			
P50	2	[-10,10]	[25]	pp. 2004
P52	5	[-20,20]	[40]	Combustion of propane chemical equilibrium equations
P53	6	[-3,3]	[42]	14-Wood function
P55	6	[-10,10]	[45]	Semiconductor boundary condition
P56	8	[-10,30]	[4]	2.3-The human heart dipole
P62	6	[0,60]	[24]	Problem 2
P63	6	[-10,10]	[35]	Example 2
P64	8	[-10,15]	[11]	Equation 3.1
P70	10	[-100,100]	[52]	Example 4.1-Nonlinear resistive circuit
P72	51	[-5,5]	[14]	D7-Diagonal of three variables premultiplied by a quasi-orthogonal matrix
P77	2	[0,3.5]	[8]	Example 1
P80	3	[0,4]	[42]	5-Beale function
P84	3	[0,1]	[34]	Example 6.2
P88	10	[-10,10]	[33]	Example 2-The Beam problem
P96	2	[-10,10]	[18]	pp. 498 (Problem N4)