Elitist Self-Adaptive Step-Size Search in Optimum Sizing of Steel Structures

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Abstract—This paper covers application of an elitist self-adaptive step-size search (ESASS) to optimum design of steel frame structures. In the ESASS two approaches are considered for improving the convergence accuracy as well as the computational efficiency of the original technique namely the so called self-adaptive step-size search (SASS). Firstly, an additional randomness is incorporated into the sampling step of the technique to preserve exploration capability of the algorithm during the optimization. Moreover, an adaptive sampling scheme is introduced to improve the quality of final solutions. Secondly, computational efficiency of the technique is accelerated via avoiding unnecessary analyses during the optimization process using an upper bound strategy. The numerical results demonstrate the usefulness of the ESASS in the sizing optimization problems of steel frame structures.

Keywords—Structural design optimization, optimal sizing, metaheuristics, self-adaptive step-size search, steel frames.

I. Introduction

Daily life is full of instances which need decision making about the best possible solution. By using the shortest path to reach the destination, shopping with a certain budget, or ordering our daily tasks, implicitly we try to find an optimum solution. Generally, time and cost limitations are the two common limitations in real life optimization instances. Similar to frequent daily problems, the field of engineering design includes a wide range of optimization problems as well. Even, it can be mentioned that engineering design without optimization is indeed not meaningful [1]. In particular the optimum design of a structural system is an attempt to find the best arrangement of solution variables that yields a minimum weight or cost design. Furthermore, for practical aspects the final design should satisfy a set of design constraints imposed with respect to a standard code. Basically, the main categories of traditional structural optimization techniques are mathematical programming [2] and optimality criteria [3, 4] approaches.

The well known shortcomings of traditional optimization methods are that these techniques are gradient-based and therefore typically work on the basis of continuous solution variables. Furthermore, computing the gradients of highly nonlinear objective functions of practical instances becomes another difficulty when dealing with these techniques. The most recent category of structural optimization techniques is referred to as non-traditional stochastic search methods or metaheuristics. These algorithms, such as genetic algorithms, particle swarm optimization, ant colony optimization, etc., are basically nature inspired approaches, which borrow their working principles from natural phenomena [5]. Different from traditional optimization techniques, metaheuristic algorithms do not perform any gradient based search and are able to handle both discrete and continuous solution variables. In addition, the stochastic nature of metaheuristics makes it more probable to find a near optimum solution even for complicated practical optimization instances. Since the optimization approaches based on metaheuristics are robust and successful in locating the optimal solutions, these algorithms can efficiently be employed for solving practical structural optimization problems. The state-of-the-art reviews of metaheuristics as well as their applications in structural design optimization can be found in Refs. [6-8].

Although meta-heuristic algorithms are generally conceived to be successful in locating promising solutions for challenging engineering optimization problems, the slow rate of convergence towards the optimum and the need for a high number of structural analyses are known as the main shortcomings of these techniques in practical structural design optimization. Mostly response computations of designs sampled during a search process mostly occupies 85-95% workload of a metaheuristic technique [9], and therefore large number of structural analyses substantially increases the total computing effort. One solution to this, is to reduce the total computational time by taking advantage of high performance computing methods, such as parallel or distributed computing techniques [9]. The idea in this approach is to distribute the total workload of the optimization algorithm amongst multiprocessors of a single computer or within a cluster of computers connected to each other via local area network. Another approach, which is more straightforward and easier to apply, is to develop efficient strategies for diminishing the number of structural analyses required in the optimization process. The latter, can be performed through developing efficient optimization techniques capable of locating reasonable solutions using less computational effort. Recently, an upper bound strategy (UBS) is proposed in Kazemzadeh Azad et al. [10], where unnecessary structural analyses are avoided during the course of optimization through a simple
and efficient mechanism. The key issue in the UBS is to identify those candidate solutions which have no chance to improve the search during the iterations of the optimum design process. After identifying those non-improving solutions, they are directly excluded from the design population without any structural analysis performed, resulting in a significant saving in computational effort [10].

Self-adaptive step-size search (SASS) algorithm is a recently proposed optimization technique based on a self-adaptive hill-climbing strategy [11]. In addition to its ability for tackling practical optimization problems, the facts that it has a simple algorithmic structure and needs relatively a small number of parameters for implementation are amongst the advantageous features of this technique. In Nolle [12] the SASS algorithm is successfully employed to find the optimum profiles for a simulated rolling mill. Nolle [13] also applied this algorithm to automated Langmuir probe tuning problem and reported numerical results indicating the favorable application of the technique. Later, Nolle and Bland [14] demonstrated the promising performance of the SASS in automatic optimization of standard engineering design problems.

This study covers application of a recently developed ESASS algorithm [15] to discrete sizing of steel frame structures. In the ESASS two approaches are considered for improving the convergence accuracy as well as the computational efficiency of the original technique. Firstly, an additional randomness is incorporated into the sampling step of the technique to preserve exploration capability of the algorithm during the optimization. Moreover, an adaptive sampling scheme is introduced to improve the quality of final solutions. Secondly, computational efficiency of the technique is accelerated via avoiding unnecessary analyses during the optimization process using an upper bound strategy. The numerical results demonstrate the usefulness of the ESASS in the sizing optimization problems of steel frame structures.

II. Problem Formulation

For a steel structure composed of \( N_m \) structural members collected in \( N_d \) member groups, the sizing optimization problem can be formulated as follows.

The objective is to find a vector of integer values such as \( I \) (Eq. 1) representing the sequence numbers of steel sections assigned to \( N_d \) member groups

\[
I^* = [i_1, i_2, ..., i_{N_d}] \tag{1}
\]

to minimize the penalized weight \( f(X) \) of the structure,

\[
f(X) = W(X) + \phi(X) \tag{2}
\]

where,

\[
W = \sum_{i=1}^{N_d} \rho_i A_i \sum_{j=1}^{N_j} L_i
\]

In the above equations \( A_i \) and \( \rho_i \) are the length and unit weight of the steel section adopted for member group \( i \), respectively, \( N_i \) is the total number of members in group \( i \), \( L_j \) is the length of the member \( j \) which belongs to group \( i \), and \( \phi(X) \) is the penalty function employed for handling the constraints. The optimization constraints consist of the limitations imposed on overall structural response and behavior of individual members which are addressed in the numerical examples section.

III. The SASS Algorithm

The present section covers the optimum design procedure based on the SASS algorithm [14]. The algorithm has a relatively simple outline, which consists of the following steps:

Step 1. Initial population: Form an initial population by spreading \( m \) solution candidates over the design space. Each solution candidate \( X_i \) is referred to as a particle \( P_i \) \((i=1, ..., m)\) in the SASS algorithm and is considered as a vector of \( n \) design variables, i.e., \( P_i = (v_{i1}, v_{i2}, ..., v_{in}) \).

Step 2. Evaluation of the initial population: Calculate the objective function value of each particle through Eq. (2). The fitness value of each particle is computed by either inverting its objective function value, or subtracting it from a constant number chosen large enough to yield always a positive value for all particles.

Step 3. Selecting a particle for improvement: Select a particle for improvement in an optimization cycle. In this process each particle \( P_i \) \((i=1, ..., m)\) is selected once according to its sequence number in the population, and the improvement of this particle is performed as discussed in the following steps.

Step 4. Defining a maximum step size vector \( S_{\text{max}} \): For each particle \( P_i \) selected in the previous step, choose two different particles \( P_k \) and \( P_j \) randomly from the population to define the neighborhood of the particle \( P_j \) based on a maximum step size vector \( S_{\text{max}} \).

\[
S_{\text{max}} = (S_{\text{max}1}, S_{\text{max}2}, ..., S_{\text{max}n}) \tag{4}
\]

\[
S_{\text{max}j} = |v_{ij} - v_{kj}| \quad \text{for} \quad j=1, 2, ..., n \tag{5}
\]

where, each component of \( S_{\text{max}} \) is equal to the absolute value of the difference between the corresponding design variables
in the particles $P_i$ and $P'_i$.

Step 5. Sampling: Sample a new particle $P'_i$ in the neighborhood of the selected particle $P_i$ based on $S_{\text{max}}$ using Eqs. (6) and (7),

$$v'_ij = v_{ij} + \text{step}_{ij}$$ \hspace{1cm} (6)

$$\text{step}_{ij} \in [-S_{\text{max}}, S_{\text{max}}] \quad \text{for } j=1,2,\ldots,n$$ \hspace{1cm} (7)

where $v_{ij}$ and $v'_ij$ are the $j$-th design variable in the particles $P_i$ and $P'_i$ respectively, and step$_{ij}$ is any number randomly chosen between the range $[-S_{\text{max}}, S_{\text{max}}]$ using a uniform distribution.

Step 6. Evaluation of the sampled particle: Calculate the fitness value of the newly sampled particle $P'_i$.

Step 7. Updating: Compare the sampled particle $P'_i$ with the original particle $P_i$ based on their fitness values. If $f(P'_i) < f(P_i)$ then $P_i$ is updated and replaced by $P'_i$, otherwise $P_i$ is retained.

Step 8. Termination: Go to Step 3 until a stopping criterion is satisfied, which can be imposed as a maximum number of iterations or no improvement of the best design over a certain number of iterations. It should be noted that one cycle in SASS is composed of $m$ iterations.

iv. The ESASS Algorithm

A reformulation of the SASS algorithm is proposed in Ref. [15] to improve the efficiency of the algorithm in structural design optimization problems. The resulting enhancement of the technique is referred to as elitist self-adaptive step-size search (ESASS) algorithm. The ESASS algorithm exhibits some superiority with respect to its standard variant in terms of both convergence accuracy and computational efficiency. In the following the enhancements in the ESASS algorithm are described in details.

In the SASS algorithm typically the components of step size vector $S_{\text{max}}$ are large in the initial cycles due to a random generation of the initial population, and they tend to decrease adaptively with the convergence of the population as the search goes on. This self-adaptive nature of the algorithm is intended to provide a suitable search mechanism by sampling new particles in a restricted, yet more favorable region of the design space in the following cycles. However, the performance of the algorithm is investigated through numerical examples, it is observed that the $S_{\text{max}}$ values tend to become very small or even zero after a certain number of cycles, resulting in negligible or sometimes no changes in the generated particles. It follows that exploration ability of the algorithm vanishes in time, leading to degenerated or sometimes totally disabled search process by the SASS algorithm. As a remedy to this problem, Eq. (6) is somewhat modified in the proposed ESASS algorithm. An additional term, rand$_{ij}$, based on a standard normal distribution, $N(0,1)$, with a mean of zero and standard deviation of 1, is used in each iteration with a probability of $R_p$ as follows:

$$v'_ij = v_{ij} + \text{step}_{ij} + \text{rand}_{ij}$$ \hspace{1cm} (8)

$$\text{rand}_{ij} = \begin{cases} N(0,1) & \text{if } u_{ij} < R_p \\ 0 & \text{if } u_{ij} > R_p \end{cases}$$ \hspace{1cm} (9)

where $u_{ij}$ is a uniform random number selected between 0 and 1. The rationale behind Eqs. (8) and (9) is to facilitate stochastic changes in the generation of new particles to keep alive the exploration capability of the algorithm especially when the $S_{\text{max}}$ values are decreased to unnecessarily low values. However, not all components of the particle are subjected to stochastic change, instead this is controlled by the probability $R_p$. In addition, the use of a normally distributed random number in this formulation ensures that the small perturbations occur more often than the large ones.

On the other hand, some recently developed metaheuristic optimization algorithms based on elitist strategies have been found to be very efficient in locating optimum or near-optimum solutions while tackling complicated design optimization problems [16-18]. For instance, two enhanced metaheuristic algorithms [17, 18] that are specifically developed by the authors for handling sizing optimization problems work fundamentally on the basis of an elitist strategy where the new candidate solutions are generated in the vicinity of the current best design.

An attempt is made to utilize an elitist strategy in the ESASS algorithm where the sampling of new particles (Step 5) is encouraged in the neighborhood of the best-so-far particle in accordance with Eq. (10),

$$v'_ij = v^{'\text{best}}_{ij} + \text{step}_{ij} + \text{rand}_{ij}$$ \hspace{1cm} (10)

where $v^{'\text{best}}_{ij}$ refers to the $j$-th component of the best particle $P^{\text{best}}$, found so far in the optimization process. It should be noted that Eqs. (8) and (10) offer two competitive formulations to be used in place of Eq. (6) for sampling new particles in searching the design space. Apparently, a more explorative search is provided with Eq. (8), whereas Eq. (10) motivates a more exploitative search by benefitting from previously visited best solution.

To combine these two useful search features in an efficient
manner, an adaptive sampling scheme with the following pseudo-code is developed in the ESASS algorithm:

\[
\text{if } \left( u_i \leq R_s \right) \text{ then} \\
\quad P_{\text{selected}} = P_i \\
\quad \text{- Sample new particle } P'_i \text{ using Eq. (8)} \\
\text{else} \\
\quad P_{\text{selected}} = P_{\text{best}} \\
\quad \text{- Sample new particle } P'_i \text{ using Eq. (10)}
\]

where

\[
R'_i = \begin{cases} 
R_i' & \text{if } P_{\text{best}} \text{ is not improved at iteration } t \\
R_i' + 0.01 & \text{if } P_{\text{best}} \text{ is improved by Eq.(11) at iteration } t \\
R_i' - 0.01 & \text{if } P_{\text{best}} \text{ is improved by Eq.(13) at iteration } t
\end{cases}
\]

In the proposed adaptive sampling scheme a new particle \( P'_i \) is generated by applying either one of these two sampling equations (Eq. 8 and Eq. 10) probabilistically. Here the sampling probability parameter \( R_s \in [0,1] \) controls the sampling scheme to be implemented when generating a new particle. For each particle, a uniform random number \( u_i \) is generated anew between 0 and 1, and at times when \( u_i \leq R_s \), the new particle is sampled using Eq. (8), otherwise it is generated using Eq. (10). It follows that the probability of sampling a new particle with Eq. (8) and Eq. (10) is \( R_s \) and \( 1 - R_s \) respectively. The \( R_s \) parameter is initially set to 0.5 to give an equal chance to either sampling scheme in the beginning. However, when iteration of a cycle is completed (i.e. when \( n \) number of particles are sampled and evaluated in a cycle) \( R_s \) is updated adaptively using Eq. (11), and this way the search is biased towards the sampling scheme that exhibits a better performance at the previous iteration.

In Eq. (11), \( R'_i \) and \( R'^{t+1} \) represent the sampling probability parameters at cycles \( t \) and \( t+1 \), respectively. Accordingly, if the best design \( P_{\text{best}} \) is improved by a particle sampled using Eq. (8) at the previous iteration, \( R_s \) is increased by 0.01; otherwise if \( P_{\text{best}} \) is improved by a particle sampled using Eq. (10), then \( R_s \) is lowered by 0.01. No update of \( R_s \) is carried out if \( P_{\text{best}} \) is not improved at the previous iteration.

On the other hand the computational efficiency of the ESASS algorithm is accelerated via the recently developed UBS method [10]. In this approach, basically the penalized weight of a current solution is considered as an upper bound limit for the net weight of a newly generated solution. Accordingly, a new solution with a net weight greater than this limit is excluded from the structural analysis stage. This strategy is used in the ESASS algorithm as follows. Here, after a new particle \( P'_i \) is sampled in Step 5 in the vicinity of a selected particle \( P_i \) or \( P_{\text{best}} \), first the net weight of \( P'_i \), i.e. \( W(P'_i) \), is calculated only; not the penalized weight. This computation is straightforward and can be done with a trivial computational effort. If \( P'_i \) has a net weight smaller than or equal to the penalized weight of the selected particle \( f(P_{\text{selected}}) \), the structural analysis of the new particle is processed and its penalized weight is computed. In the opposite case, i.e. \( W(P'_i) > f(P_{\text{selected}}) \), however, the upper bound rule is activated and \( P'_i \) is automatically excluded from the structural analysis phase required for response computations in Step 6.

v. Numerical Example

The design example covered in this section is a 135-member steel frame. The optimum solution found for this structure with the ESASS algorithm is compared to those achieved using other contemporary metaheuristic algorithms.

A. 135-Member Steel Frame

The first optimization instance is a 3-story steel frame depicted in Figure 1, composed of 135 members including 66 beam, 45 column and 24 bracing elements. The stability of structure is provided through moment-resisting connections as well as inverted V-type bracing systems along the \( x \) direction. For practical fabrication requirements the 135 members of the frame are collected under 10 member groups. For design purpose, the frame is subjected to 10 load combinations to ASCE 7-98 [19]. Further details for this example can be found in Ref. [10].

Discrete sizing of the frame is previously carried out in Ref. [10] via some contemporary metaheuristics, i.e. the upper bound strategy (UBS) integrated big bang-big crunch algorithm (UBB-BC), as well as its two enhanced variants i.e. UBS integrated modified and exponential big bang-big crunch algorithms (UMBB-BC and UEBB-BC). Moreover, this instance is also solved in Ref. [20] using a UBS integrated particle swarm optimization algorithm (UPSO).

Table 1 presents a comparison of optimum solutions located using different algorithms. As can be seen from this table, the ESASS yields a design weight of 44.33 ton for this example. Other solutions obtained are 38.91 ton by UEBBBC, 45.67 ton by UMBBB-BC, 47.3 ton by UBB-BC, and 55.66 ton by UPSO. These design weights are obtained using 1542 by ESASS, 1235 analyses by UEBB-BC, 1794 analyses by UMBBB-BC, 880 analyses by UBB-BC, and 1574 analyses by UPSO. It can be observed that the ESASS algorithm shows a promising performance which is comparable to the aforementioned contemporary enhanced optimization algorithms both in terms of solution quality as well as computational efficiency. Furthermore, the ESASS
optimization algorithm needs few parameters for implementation.

![Figure 1. 135-Member Steel Frame](image)

### TABLE I. COMPARISON OF RESULTS FOR 135-MEMBER STEEL FRAME

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### VI. Conclusion

In this paper application of the recently developed ESASS algorithm to optimum design of steel skeletal structures is presented. Basically, in the ESASS two strategies are considered for improving the convergence accuracy as well as the computational efficiency of the original technique. On the one hand, an additional randomness is incorporated into the sampling step of the technique to preserve exploration capability of the algorithm during the optimization. On the other hand, an adaptive sampling scheme is introduced to improve the quality of final solutions. Furthermore, as a result of integrating the UBS with the ESASS algorithm the total number of required structural analyses is reduced. The numerical investigations indicate a promising performance of the ESASS algorithm with an acceptable level of comparability to the contemporary enhanced optimization algorithms both in terms of solution quality as well as computational efficiency.

**References**


