Meta Model Based EA for Complex Optimization

Maumita Bhattacharya, Member IEEE

Abstract—Evolutionary Algorithms are population-based, stochastic search techniques, widely used as efficient global optimizers. However, many real-life optimization problems often require finding optimal solutions to complex high-dimensional, multimodal problems involving computationally very expensive fitness function evaluations. Use of evolutionary algorithms in such problem domains is thus practically prohibitive. An attractive alternative is to build meta-models or use an approximation of the actual fitness functions to be evaluated. These meta-models are order of magnitude cheaper to evaluate compared to the actual function evaluation. Many regression and interpolation tools are available to build such meta-models. This paper briefly discusses the architectures and use of such meta-modeling tools in an evolutionary optimization context. We further present two evolutionary algorithm frameworks which involve use of meta models for fitness function evaluation. The first framework, namely the Dynamic Approximate Fitness based Hybrid EA (DAFHEA) model [14] reduces computation time by controlled use of meta-models (in this case approximate model generated by Support Vector Machine regression) to partially replace the actual function evaluation by approximate function evaluation. However, the underlying assumption in DAFHEA is that the training samples for the meta-model are generated from a single uniform model. This does not take into account uncertain scenarios involving noisy fitness functions. The second model, DAFHEA-II, an enhanced version of the original DAFHEA framework, incorporates a multiple-model-based learning approach for the support vector machine approximator to handle noisy functions [15]. Empirical results obtained by evaluating the frameworks using several benchmark functions demonstrate their efficiency.

Keywords—Meta model, Evolutionary algorithm, Stochastic technique, Fitness function, Optimization, Support vector machine.

I. INTRODUCTION

The optimization of complex, high-dimensional, multimodal problems often poses a problem which in fact depends on the structure inherent in the problems. Examples of such problem domains include large-scale finite element analysis (FEA), computational fluid dynamics (CFD), engineering design problems etc. For example, optimization problems involving engineering design are often characterized by features that make them highly computationally expensive to be solved by evolutionary algorithms or even other standard nonlinear optimization techniques. One of the most critical of these features is that the functions used to define such optimization problems are often computationally intensive. In engineering design optimization problem domains, a design can be represented by a number of continuous design parameters and the potential solutions are vectors in a multidimensional vector space [27]. Determining the fitness of these potential solutions generally involve some form of simulation to compute the relevant physical properties of the object represented by the solution vector and consolidate them into a single measure of merit, including the information on the status of the constraints if necessary. In such problems, the run-time for a single function evaluation could be in the range from a fraction of a second to hours of supercomputer time. The computational expense of evaluating the functions that define the problem would necessarily be incurred for each iteration, within any iterative optimization algorithm and in a population-based search algorithm like EA, for the entire population or part of the population, as is required, in each iteration. Considering such prohibitive computational costs, a feasible alternative is to build approximate models, within an optimization context, since these approximate models are order of magnitude cheaper to run compared to the actual function evaluations [8,10,11]. Many regression and interpolation tools could be used to construct such meta models, (e.g., least square regression, back propagating artificial neural network, response surface models, etc.) which provide less accurate, but more efficient (in terms of computational cost) measures of the merit of the fitness functions.

However, it can not be denied that accuracy of the result is a major risk involved in using meta-models to replace actual function evaluation [23, 25, 26, and 27]. When it is infeasible to precisely judge when, where and how much of such replacement is optimal, using a controlled approach holds the answer. The use of approximate model to speed up optimization dates all the way back to the sixties [14]. The most widely used models being Response Surface Methodology [17], Kriging models [9] and artificial neural network models [5]. The concepts of using approximate model vary in levels of approximation (Problem approximation, Functional approximation, and Evolutionary approximation), model incorporation mechanism and model management techniques [27].
In the multidisciplinary optimization (MDO) community, primarily response surface analysis and polynomial fitting techniques are used to build the approximate models [16, 23]. These models work well when single point traditional gradient-based optimization methods are used. However, they are not well suited for high dimensional multimodal problems as they generally carry out approximation using simple quadratic models.

In another approach, multilevel search strategies are developed using special relationship between the approximate and the actual model. An interesting class of such models focuses on having many islands using low accuracy/cheap evaluation models with small number of finite elements that progressively propagate individuals to fewer islands using more accurate/expensive evaluations [7]. As is observed in [27], this approach may suffer from lower complexity/cheap islands having false optima whose fitness values are higher than those in the higher complexity/expensive islands. Rasheed et al. in [10, 11], uses a method of maintaining a large sample of points divided into clusters. Least square quadratic approximations are periodically formed of the entire sample as well as the big clusters. Problem of unvaluable points was taken into account as a design aspect. However, it is only logical to accept that true evaluation should be used along with approximation for reliable results in most practical situations. Another approach using population clustering is that of fitness imitation [27]. Here, the population is clustered into several groups and true evaluation is done only for the cluster representative [8]. The fitness value of other members of the same cluster is estimated by a distance measure. The method may be too simplistic to be reliable, where the population landscape is a complex, multimodal one.

Jin et al. in [25, 26] analysed the convergence property of approximate fitness based evolutionary algorithm. It has been observed that incorrect convergence can occur due to false optima introduced by the approximate model. Two controlled evolution strategies have been introduced. In this approach, new solutions (offspring) can be (pre)-evaluated by the model. The (pre)-evaluation can be used to indicate promising solutions. It is not clear however, how to decide on the optimal fraction of the new individuals for which true evaluation should be done [6]. In an alternative approach, the optimum is first searched on the model. The obtained optimum is then evaluated on the objective function and added to the training data of the model [1, 22, and 6]. Yet in another approach as proposed in [25], a regularization technique is used to eliminate false minima.

It is obvious that incorporation of approximate models may be one of the most promising approaches to realistically use EA to solve complex real life optimization problems, especially where: (i) Fitness computation is highly time-consuming, (ii) Explicit model for fitness computation is absent, (iii) Environment of the evolutionary algorithm is noisy etc. However, considering the obvious risk involved in such approach, an EA with efficient control strategy for the approximate model and robust performance is welcome.

While dealing with complex real world optimization problems, expensive function evaluations can be feasibly used only in a limited manner. Research on using surrogate models should focus on:
1) Minimizing uncertainty in approximate estimation
2) Employing corrective measures
3) Exploring ways to exploit the approximate knowledge for improving the optimization technique.

The hybrid evolutionary algorithm framework, DAFHEA (dynamic approximate fitness based hybrid evolutionary algorithm) addresses some of the above issues [14]. DAFHEA replaces expensive function evaluation by its support vector machine (SVM) approximation. The concept of merit function [22] is borrowed to maintain diversity in the solution space using approximate knowledge. However, the assumption used in the original DAFHEA is that the training samples for the meta-model are generated from a single uniform model. This does not cover situations, where information from variable input dimensions and noisy data is involved. DAFHEA-II attempts to correct this by using a multi-model regression approach. The multiple models are estimated by successive application of the SVM regression algorithm. Retraining of the model is done in a periodic fashion.

The original DAFHEA framework is similar to other existing models in that it uses an approximation model to partially replace expensive fitness evaluation in evolutionary algorithm. An explicit control strategy (a cluster-based on-line learning technique) to improve reliability of using such approximate models to reduce expensive function evaluations was introduced. Also the approximate knowledge thus generated is exploited to avoid premature convergence (one of the major impediments of using evolutionary algorithm to solve complex real life optimization problems). However, the major constraint associated with DAFHEA is that it treats the solution space as one comprising of information coming from a uniform model. Situations like model formation involving variable input dimensions and noisy data certainly can not be covered by this assumption. DAFHEA-II addresses this issue by using a multiple model regression approach for the SVM approximator.

The rest of the paper is arranged as follows: Section II presents a brief description of the available meta model generation tools. The basic frameworks and the functional details of DAFHEA and DAFHEA-II (the enhanced version of DAFHEA) are presented in Section III, IV and Section V. Simulation results are presented in Section VI. Finally conclusions are drawn in Section VII.

II. TOOLS FOR META MODEL GENERATION
Meta models in this context are models that are developed to approximate computationally expensive simulation codes. Functional approximation modeling generally involves finding a set of parameters for a given model to find the good, best or perfect fit between a given finite sampling of values of independent variables and associated values of dependent
variables [27]. A wide variety of empirical tools are used to generate functional approximation models [17,9,5]. Some of the commonly used ones are polynomial interpolation, DACE (design and analysis of computer model) or kriging model, artificial neural networks, regression spline etc. An important characteristic of a meta model generator is generalization. Generalization is the ability to map or predict values that were not considered in the training set while developing the model. The least square method (LSM) performs efficiently only within a small trust region and fails in terms of generalization particularly for complex polynomials with discontinuity in the target function. However, for low dimensional problems with real valued parameters, the polynomial regression models often outperform the connectionist methods. The connectionist models, like the neural networks perform better for high dimensional problems. Unlike the LSM, the kriging models are capable of capturing multiple local extrema, but at the expense of higher computational cost. Some of the popular approximations modeling tools are described below:

A. Polynomial Approximation Models

One of the popular and simple meta modeling methods is the linear polynomial regression model. This method involves finding the coefficients of the linearly combined terms in a given low order polynomial or a vector of polynomials. When the used error function is the mean square error, the Least Square Method (LSM) can be used to find the coefficients [27]. The method is also known as the response surface method when LSM is used.

The order of polynomial is important. Quadratic and cubic polynomials are most commonly used; with quadratic polynomial being most suited for the continuous, unimodal problems. Splines or piecewise polynomial fits with continuity of the polynomial being most suited for the continuous, unimodal problems.  

The second-order polynomial approximation model has the following form:

\[ \hat{y} = c_0 + \sum_{l=0}^{L} c_l x_l + \sum_{l=0}^{L} c_{l+i+j} x_l x_j \]

where, \( c_0, c_l \) and \( c_{l+i+j} \) are the coefficients to be estimated. Note that, there are \( n_r = n_1(n_1+1)/2 \) modeling terms in the quadratic polynomial, where, \( n_r \) is the number of input variables.

Both least square method (LSM) and the gradient method can be used to estimate the unknown coefficients [27].

1) Least Square Method

If \( n_r \) is the number of samples drawn from the original function and \( p = 1, \ldots, n_s \) then the polynomial model in matrix notation has the following form:

\[ y^{(p)} = c^T x^{(p)} \]

where, \( c \) is the vector of \( n_r \) unknown coefficients to be estimated.

\[ c = [c_0, c_1, \ldots, c_{n-1}] \]

and \( x^{(p)} \), the polynomial model, is a vector of length \( n_r \).

Estimating the unknown coefficients will require that \( n_r \geq n_s \). The estimation problem can now be formulated as below:

\[ y \approx X \hat{c} \]

where, \( y \) is the vector of \( n_s \) response values,

\[ y = [y_1, y_2, \ldots, y_{n_s}]^T \]

and \( X \) is the matrix of \( p \) row vectors with assumed rank of \( n_s \),

\[
X = \begin{bmatrix}
1 & x_1^{(1)} & x_2^{(1)} & \cdots & x_{n_v}^{(1)} \\
1 & x_1^{(2)} & x_2^{(2)} & \cdots & x_{n_v}^{(2)} \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
1 & x_1^{(n_s)} & x_2^{(n_s)} & \cdots & x_{n_v}^{(n_s)}
\end{bmatrix}
\]

The unique LSM solution is as follows,

\[ \hat{c} = (X^T X)^{-1} X^T y \]

where, \( \hat{c} \) denotes the estimate of \( c \) and \((X^T X)^{-1}\) exists if the rows of \( X \) are linearly independent.

2) Gradient Method

The least square method is unsuitable for high dimensional problems due to high computational expenses. Gradient method is a suitable alternative to address this problem [27]. Let the square error function for the \( k^{th} \) sample be:

\[ E^{(k)} = \frac{1}{2} (y - y^{(k)})^2 \]

The unknown coefficients may now be updated as below [27]:

\[ \Delta c_0 = -\xi \cdot (y - y^{(k)}) \]

\[ \Delta c_j = -\xi \cdot (y - y^{(k)}) x_j^{(k)} \]

\[ \Delta c_{n-i+j} = -\xi \cdot (y - y^{(k)}) x_i^{(k)} x_j^{(k)} \]

\( 1 \leq i \leq j \leq n \)

B. The DACE Model

The DACE (design and analysis of computer experiments) is a
statistically sound estimation method. This is also known as
gaussian process regression in the neural network literature
and Kriging in the geostatistics literature. The DACE model
may be understood as a two-component model that includes
a global model and a localized “deviation”. It can be expressed
as,
\[ y(x) = f(x) + Z(x) \]
where, \( f(x) \) represents a global model of the original
function and \( Z(x) \) is a Gaussian random function with zero
mean and non-zero covariance that creates a localized
deviation from the global model. \( f(x) \) is usually a
polynomial and can be considered to be an underlying
constant \( \beta \). Then the representation of DACE model
becomes,
\[ y(x) = \beta + Z(x) \]
The covariance matrix of \( Z(x) \) is expressed as,
\[ \text{Cov}(Z(x^{(i)}),Z(x^{(j)})) = \sigma^2 R(x^{(i)},x^{(j)}) \]
where, \( R \) is the symmetric correlation matrix with values of
unity along the diagonal and dimension of \( n_s \times n_s \). \( R \) is the
correlation function between any two of \( n_s \) samples. The user
may select the form of the correlation function \( R \). A choice
for \( R \) is often found in statistical literature is an exponential
function, \( R(x^{(i)},x^{(j)}) = \exp \left( -\sum_{k=1}^{n} \theta_k |x^{(i)}_k - x^{(j)}_k|^2 \right) \)
where, \( \theta_k \) is the vector of the unknown correlation
parameters. \( x^{(i)}_k \) and \( x^{(j)}_k \) are the \( k \)-th component of the
sample point \( x^{(i)} \) and \( x^{(j)} \). The prediction of \( y(x) \) can thus be
represented as,
\[ \hat{y}(x) = \hat{\beta} + \mathbf{r}^T(x) R^{-1} \mathbf{y} - \hat{\beta} \mathbf{r} \]
where, \( \hat{\beta} \) is unknown and both \( \mathbf{r}(x) \) and \( R \) depend on the
unknown parameter \( \theta \). The vector \( \mathbf{r} \) has length \( n_s \), with all
entries equal to unity. \( \mathbf{y} \) is a vector of length \( n_s \) and \( \hat{\beta} \) is the
estimated value of \( y \) given the \( n_s \) samples and the current
input \( x \). \( \mathbf{r} \) is the correlation vector of length \( n_s \) between
the given input \( x \) and the samples \( \{x^{(1)}, \cdots, x^{(n_s)}\} \)
\[ \mathbf{r}^T(x) = [R(x,x^{(1)}), R(x,x^{(2)}), \cdots, R(x,x^{(n_s)})]^T \]
The estimation of the parameters can be carried out using the
maximum likelihood method.

In DACE, a confidence interval of the estimation can be
obtained without much additional computational expense.
However, when the dimensionality is high, computational cost
raises as DACE involves matrix inversions for estimating the
output.

C. Artificial Neural Networks

Artificial neural networks have been seen to be effective
approximation tools. The artificial neural networks consist of
a large number of highly interconnected processing units, each
aggregating information from a large number of connected
peers.

The feed forward multilayer perceptrons and the radial
basis function networks are the popular ones.

1) Multilayer Perceptrons

A multilayer perceptron with one input layer, two hidden
layers and one output neuron can be expressed as [27],
\[ y = \sum_{t=1}^{n} v_t f \left( \sum_{k=1}^{K} w_{tk} f \left( \sum_{i=1}^{s} w_{ik} x_i \right) \right) \]
where, \( n \) is the number of inputs, \( K \) and \( L \) are the
number of hidden nodes, and \( f \) is called the activation
function, which usually is the logistic function.
\[ f(z) = \frac{1}{1 + e^{-az}} \]
where, \( a \) is a constant.

2) Radial Basis Function Network

The radial basis functions are powerful interpolation tools for
multidimensional problems. The output of a radial basis
function network is a weighted sum of radial basis functions
\( \phi_j \), each being a local function (usually Gaussian) of the
distance \( \|x - c_j\| \) between the input vector \( x \) and a “basis
function centre”, \( c_j \).

\[ y(x) = \sum_{j=1}^{n} w_j \phi \left( \| x - c_j \| \right) \]
where, \( \| \| \) is typically an Euclidean norm and \( w_j \) are the
weights or the unknown coefficients to be determined.

The normalized RBF network can be expressed as,
\[ y(x) = \frac{\sum_{j=1}^{n} w_j \phi \left( \| x - c_j \| \right)}{\sum_{j=1}^{n} \phi \left( \| x - c_j \| \right)} = \frac{\sum_{j=1}^{n} w_j u \left( \| x - c_j \| \right)}{\sum_{j=1}^{n} \phi \left( \| x - c_j \| \right)} \]
where, \( u \left( \| x - c_j \| \right) \) is known as the normalized radial basis
function.

D. Support Vector Machines

The conceptual background of single approximation model
building using support vector machines has been outlined in
Section V. A.
E. Some Comparative Information

It is hard to compare the performances of the different approximation models as performance can be problem dependent and also there are several criteria that need to be considered. However, the most important ones are the accuracy, both on the training and the test data, computational complexity and transparency [27]. One of serious problem is the introduction of false optima. A desirable tradeoff may be that of lower approximation accuracy if the model is used in global optimization. Some methods for prevention of false minima in neural network are available [27].

In [27] Jin has suggested the following general rules for model selection. It is recommended to implement first a simple approximate model, for example, a lower order polynomial model to see if the given samples can be fit with reasonable accuracy. If it fails a model with higher complexity such as higher order polynomials or neural network models should be considered. However, for high dimensional problems with small number of samples, a neural network model is generally preferable. In case of neural network models, in particular a multilayer perceptrons network, the model complexity should be controlled to avoid over-fitting. The gradient descent based method may found to be of slow convergence in some cases. The RBF networks show superior performance both in terms of accuracy and training speed for some problems [27].

The subsequent sections describe two EA frameworks that use SVM regression to generate the meta-models.

III. THE DAFHEA FRAMEWORK

The DAFHEA framework includes a global model of genetic algorithm (GA), hybridised with support vector machine (SVM) as the approximation tool. Expensive fitness evaluation of individuals as required in traditional evolutionary algorithm is partially replaced by a SVM approximation (regression) model. Explicit control strategies are used for evolution control, leading to considerable speedup without compromising heavily on solution accuracy. Also the approximate knowledge about the solution space generated is used to maintain population diversity to avoid premature convergence.

While approximation is not a new idea in accelerating iterative optimization process, DAFHEA focuses on controlled speedup to avoid detrimental effects of approximation and also exploiting approximate knowledge to improve optimization solution. The following section presents the basic algorithm structure of DAFHEA.

A. The Basic Algorithm Structure

Step One: Create a random population of \( N_c \) individuals, where, \( N_c = 5 \times N_a \) and \( N_a \) = actual initial population size.

Step Two: Evaluate \( N_c \) individual using actual expensive function evaluation. Build the SVM approximate model using normalised expensive function evaluation values as training set for off-line training. (Use of normalised values in the training set appears to improve performance of meta model, reducing effects of unnaturally high or low values). SVM hyper-parameters are initially tuned based on this training set.

Step Three: Select \( N_b \) best individual out of \( N_c \) evaluated individuals to form the initial GA population.

Remarks: The idea behind using five times the actual EA population size (as explained in Step One) is to make the approximation model sufficiently representative at least initially. Since initial EA population is formed with \( N_a \) best individuals out of these \( N_i \) individuals, with high recombination and low mutation rates, the EA population in first few generations is unlikely to drift much from its initial locality. Thus it is expected that large number of samples used in building the approximation model will facilitate better performance at this stage. Also using the higher fitness individuals, chosen out of a larger set should give an initial boost to the evolutionary process.

Step Four: Select parents using suitable selection operator and apply genetic operators namely recombination and mutation to create new generation.

Step Five: Use SVM approximation model to compute fitness of new generation individuals based on approximate evaluation. Form \( m \) distance-based (considering spatial distribution of individuals) clusters in the new population space. If for some \( n \) clusters, the standard deviation \( \sigma \geq \text{Predefined Threshold} \), rearrange solution space into \( m+n \) clusters. Compute a merit function \( f_m(x) \) as below:

\[
f_m(x) = f_a(x) - \rho_1 \sigma_i - \rho_2 d_y - \rho_3 s_i
\]

where, \( f_a(x) \) is the predicted fitness function value. \( \sigma_i \) is standard deviation (in terms of objective value) for the \( i^{th} \) cluster and \( d_y \) is the normalized minimum Euclidean distance of \( j^{th} \) point of \( i^{th} \) cluster from the all truly evaluated points so far [21]. \( s_i \) is the sparseness of the \( i^{th} \) cluster. \( \rho_1 \), \( \rho_2 \) and \( \rho_3 \) are scaling factors for \( \sigma_i \), \( d_y \) and \( s_i \) respectively.

\[
s_i = \frac{\text{No of individuals in cluster } i}{\text{Dimension of individual}}
\]

Step Six: Dynamically update the approximate model as below:

1) Identify the cluster containing the optimum based on approximation.

2) Perform expensive evaluation for the approximate optimum and its \( k \) nearest neighbours. Also perform expensive evaluation for the centroid of all other data clusters and their \( k \) nearest neighbours. Expand neighbourhood for true evaluation until a point is found in each space dimension such that % error \( \delta \leq \text{Predefined threshold} \). Here,

\[
\delta = \frac{a_x - \hat{a}_x}{a_x} \times 100
\]
where, \( a_i \) = True value of the \( i^{th} \) neighbour and 
\( a_{ip} \) = Predicted value of the \( i^{th} \) neighbour and \( \max i = k \).

3) Add the newly evaluated points to approximate model training set to update model.

**Step Seven:** When termination/evolution control criteria are not met, repeat Step Four to Step Seven.

It must be noted, the optimum is considered based on the original predicted value \( f_o(x) \). For all other purposes fitness based on the merit function \( f_m(x) \) is considered. Periodic parameter tuning of the SVM approximation model was incorporated, though no specific criterion was used.

The following section presents the DAFHEA-II framework that incorporates the enhancement to tackle situations where the inputs to the meta model generation tools are generated by different models.

## IV. THE DAFHEA-II FRAMEWORK

As in the original DAFHEA framework, DAFHEA-II [Figure I] includes a global model of genetic algorithm (GA), hybridised with support vector machine (SVM) as the approximation tool. Expensive fitness evaluation of individuals as required in traditional evolutionary algorithm is partially replaced by SVM approximation (regression) models. Evolution control is implemented by periodic expensive evaluations, leading to considerable speedup without compromising heavily on solution accuracy. Also the approximate knowledge about the solution space generated is used to maintain population diversity to avoid premature convergence.

DAFHEA-II is specifically suited for applications involving information that could be considered generated by more than one model. As in original DAFHEA, this framework also focuses on controlled speedup to avoid detrimental effects of approximation and exploiting approximate knowledge to improve optimization solution. The following section presents the basic algorithm structure of DAFHEA-II, while Section IV explains its major functional aspects.

### A. Basic Algorithm Structure

The proposed DAFHEA-II framework is introduced in the context of unconstrained optimization problems. Figure (1) schematically presents the algorithm.

![The DAFHEA-II Framework](image)

*/ The basic algorithm for the DAFHEA-II framework */

```
Procedure DAFHEA_II
{
  initialize population matrix, \( gen = 0 \) and set \( \alpha, \beta \)
  call actual function evaluation
call Procedure train_SVM to generate approximation models
while (\( gen < \alpha \))
{
  \( gen = gen + 1 \)
  rank solutions based on fitness
  retain actual elite
  apply crossover and mutation to generate offspring
  call Procedure predict_SVM to approximate the fitness of the offspring
  retain approximate elite
  if (\( gen \mod \beta = 0 \)) then
  {
    call actual function evaluation
    call Procedure train_SVM
  }
  rank solutions based on fitness
  get the best solution
} /* This procedure estimates multiple models from the training data set */
Procedure train_SVM
{
  initialize data set=population in current generation with fitness resulting from actual evaluation
while (!stopping criterion)
{
  /* Estimate the dominant model describing majority of the candidates in data set */
  apply robust regression to data set
  /* Partition the available data */
  { analyse available data to separate others from majority based on their distance from the dominant model
    remove subset of data generated by the dominant model from the data set
  }
} /* This procedure selects the most likely model for each member of the population to find its corresponding estimate */
Procedure predict_SVM
{
  while (population member)
  {
    determine appropriate model for the given test sample \( z = (x, y) \) using a distance measure from each model
  }
}
```
In the above framework (see Figure 1) \( \alpha \) is the number of predetermined generations and \( \beta \) is the predetermined retraining frequency.

**B. The Implementation of the DAFHEA-II Framework**

The detailed implementation of the DAFHEA-II [15] framework is as given below:

**Step One:** Create a random population of \( N_c \) individuals, where \( N_c = 5 \times N_a \) and \( N_a \) = actual initial population size.

**Step Two:** Evaluate \( N_c \) individual using actual expensive function evaluation. Build the SVM approximate models using the candidate solutions as input and the actual fitness (expensive function evaluation values) as targets forming the training set for off-line training. Details of the Multiple Model Formation technique is described in Section IV.

**Step Three:** Select \( N_e \) best individual out of \( N_c \) evaluated individuals to form the initial GA population.

**Remarks:** The idea behind using five times the actual EA population size (as explained in Step One) is to make the approximation model sufficiently representative at least initially. Since initial EA population is formed with \( N_e \) best individuals out of these \( N_c \) individuals, with high recombination and low mutation rates, the EA population in first few generations is unlikely to drift much from its initial locality. Thus it is expected that large number of samples used in building the approximation model will facilitate better performance at this stage. Also using the higher fitness individuals, chosen out of a larger set should give an initial boost to the evolutionary process.

**Step Four:** Rank the candidate solutions based on their fitness value.

**Step Five:** Preserve the elite by carrying over the best candidate solution to the next generation.

**Step Six:** Select parents using suitable selection operator and apply genetic operators namely recombination and mutation to create children (new candidate solutions) for the next generation.

**Step Seven:** The SVM regression models created in Step two are applied to estimate the fitness of the children (new candidate solutions) created in Step six. This involves assignment of most likely or appropriate models to each candidate solution. The details of this prediction mechanism are given in Section IV.

**Step Eight:** The set of newly created candidate solutions is ranked based on their approximate fitness values.

**Step Nine:** The best performing newly created candidate solution and the elite selected in Step five are carried to the population of the next generation.

**Step Ten:** New candidate solutions or children are created as described in Step six.

**Step Eleven:** Repeat Step seven to Step ten until either of the following condition is reached:

i. The predetermined maximum number of generations has been reached; or

ii. The periodic retraining of the SVM regression models is due.

**Step Twelve:** If the periodic retraining of the SVM regression models is due, this will involve actual evaluation of the candidate solutions in the current population. Based on this training data new regression models are formed. The algorithm then proceeds to execute Step four to Step eleven.

**Remarks:** The idea behind using periodic retraining of the SVM regression models is to ensure that the models continue to be representatives of the progressive search areas in the solution space.

**V. MAJOR FUNCTIONAL ASPECTS OF DAFHEA-II**

The following sections detail the major functional aspects of the DAFHEA-II framework.

**A. Single Approximation Model Formation Using SVM Regression**

The theoretical background of support vector machine is mainly inspired from statistical learning theory [24]. Major advantages of the support vector machines over other machine learning models such as neural networks, are that there is no local minima during learning and the generalization error does not depend on the dimension of the space. Also the fast learning ability of the SVM regression [4, 2] model is a desirable property for on-line learning. In DAFHEA (both original and enhanced versions), as the approximation model has to be rebuilt frequently to be representative of the progressing solution space, this is an important criterion for model selection.

Let us consider the problem of approximating the set of data,

\[
D = \{(x^1, y^1), ..., (x^l, y^l)\}
\]

with a linear function,

\[
f(x) = w \cdot x + b, \quad w, x \in R^n, b \in R
\]

The construction of a model is reduced to the minimization of the following regularized \( \varepsilon \)-insensitive loss function:

\[
L = ||w||^2 + C \cdot \frac{1}{l} \sum_{i=1}^{l} \max\{0, |y_i - f(x_i)| - \varepsilon\}
\]

where \( \varepsilon \) is the tolerable error, \( C \) is a pre-specified regularization constant and \( f \) is the function to be estimated.

The minimization of (6) is equivalent to the following constrained optimization problem, giving the optimal regression function as:

\[
\min \frac{1}{2} ||w||^2 + C \cdot \frac{1}{l} \sum_{i=1}^{l} (\xi_i + \xi_i^*)
\]

subject to \((w \cdot x_i) + b - y_i \leq \varepsilon + \xi_i^*)\) \hspace{1cm} (8)

\[
y_i - ((w \cdot x_i) + b) \leq \varepsilon + \xi_i^*
\]

(9)
\[ \xi_i, \xi^*_i \geq 0, \quad i = 1, \ldots, l \]

where \( \xi_i \) and \( \xi^*_i \) are slack variables representing upper and lower constraints on the output of the system.

Thus, quadratic-programming techniques can be applied to solve the minimization problem.

In the enhanced version of DAFHEA (DAFHEA-II), a multiple model regression approach is used. The technique used closely follows the approach described in [21].

B. Multiple Model Regression

The multiple model regression involves the following two stages:

i. The \textit{training/learning phase}, involving creation of the models based on training data.

ii. The \textit{prediction phase}, involving assignment of the most likely model to each candidate data and estimation of improved response using the selected model.

1) The Training/Learning Phase

Let us consider a finite number of samples or training data \((x_i, y_i), (i = 1, \ldots, n)\) [21]. The learning involves two objectives:

(a) to estimate \( N \) target models from a set of possible models:

\[
\hat{f}_m(x, \omega^*_m), (\omega^*_m \in E_m, m = 1, \ldots, N)
\]

Where \( E_m \) is a parametric space for model \( m \). Each model estimate approximates the corresponding target model \( f_m(x, \omega^*_m) \rightarrow tr_m(x) \).

(b) to partition available training data set into \( N \) subsets, where each subset belong to an appropriate model. The input \((x)\) and/or output \((y)\) space will be thus partitioned into \( N \) disjoint regions.

It is clear from the above discussion that the creation of multiple models here can be viewed as a generalization of the traditional single-model estimation. Traditional regression is applied to estimate appropriate regression-like models in a progressive manner while partitioning the data set into subsets at the same time.

2) The Prediction Phase

Using a single model approach, estimating a response \( \hat{y} \) for a given test input \( x \), simply amounts to deducing \( \hat{y} = f(x, \omega^*) \), where \( f(x, \omega^*) \) is a model predetermined based on the training data. In case of multiple model estimation, first an appropriate model has to be selected for the test input \( x \) and then the response \( \hat{y} \) can be computed as \( \hat{y} = \hat{f}_i(x, \omega^*_i) \), where \( \hat{f}_i(x, \omega^*_i) \) is the specifically chosen model for \( x \). However, it is not possible to select a model using \( x \) alone, as there may be overlapping of input domains for different models. Thus, selection of model should be based on the \((x, y)\) values of the test data as described in [21]. For a given sample of test data \( z = (x, y) \) generated by
an unknown model \( U \) and a set of models estimated during training stage:

\[
f_i(x, \omega^*_i), x \in X_i (i = 1, \ldots, N)
\]

(12)

to determine the appropriate or most likely model the distance between the test sample and each of the models in (12), has to be computed. Each model in (12) is defined as a region in the input (x) space and the mapping \( f_c : x \rightarrow y \) in this region. Therefore, the distance may be defined in the input (x) space or in the y-space, or some combination of the two.

3) Exploiting Approximate Knowledge to Avoid Premature Convergence

The basic idea is to encourage exploration along with exploitation in the initial generations. The underlying notion is to achieve this by attaching a merit point to:

i. An individual that belongs to a subset that is furthest from the set that is associated with the dominant model;

ii. An individual coming from a sparsely populated, i.e., inadequately represented subset.

The merit function \( f_m(x) \) is conceptually similar to the one used in our earlier work [14].

VI. EXPERIMENT RESULTS AND DISCUSSIONS

The performance of the proposed algorithm is tested on five popular benchmark test functions (see Table I): namely, Spherical, Rosenbrock, Rastrigin, Schwefel and Ellipsoidal. These benchmark functions in the test suite are scalable and are commonly used to assess the performance of optimization algorithms [12]. For all five functions except Rosenbrock the global minimum is \( f(x) = 0 \) at \( x^n = 0 \). Rosenbrock has a global minimum of \( f(x) = 0 \) at \( x^1 = 1 \).

All simulations were carried out using the following assumptions: The population size of 100 was used for all the simulations, where \( n \) is the number of variables for the problem; for comparison purposes three sets of input dimensions are considered; namely, \( n = 5, 10 \) and 20. For all three cases, tenfold validation was done with the number of generations being 1000; the SVM regression models were trained with five times the real GA population size initially and all the simulation processes were executed using a Pentium 4, 2.4GHz CPU processor. Tables II, III and IV show the comparative statistics of the various simulations runs using canonical GA model which uses only actual function evaluations and the proposed DAFHEA and DAFHEA-II models which use actual function evaluations sparingly. We report the results for the 5-D (dimension), 10-D (dimension) and 20-D (dimension) scenarios. The reported results were obtained by achieving same level of tolerance for both canonical GA and the proposed model. For comparison purpose, results reported in [12] were considered (see Table II, III and IV).

It is clear from the depicted results that the proposed DAFHEA and DAFHEA-II models effectively reduce the number of actual evaluations for all the benchmark function in our test suite. It is true that the formation and maintenance of the regression models incorporates additional computational expense. However, this approximation based evolutionary algorithm model is not proposed for regular optimization problems where actual function evaluation is not a matter of concern. Complex real world problems involving very expensive function evaluations will benefit from such approximation based algorithms even when the reduction in the number of actual evaluations is relatively low.

VII. CONCLUSIONS

Reduction in the number of function evaluations to reach an optimum or an asymptotic optimum is of prime importance in the context of application of evolutionary algorithm to real world problems. This can drastically lower the computational expense of using EA to solve complex optimization problems. Use of approximation or meta models to replace actual functions is an attractive choice to address this issue. In this paper brief description of tools to generate such meta models has been outlined. Two evolutionary algorithm frameworks that replace actual function evaluation by SVM regression tool generated meta model evaluation, have been presented. The second framework is an enhanced version of the first. In this, a multiple model approach for support vector machine regression is used to develop the approximate models. The algorithms showed reliable performance in terms of accuracy and the overhead cost towards developing and maintaining the meta-model is not alarmingly high. Since this overhead is expected not to increase much with increased problem complexity, both the versions of DAFHEA should lead to considerable speed up for complex real life problems. As mentioned earlier the DAFHEA-II [15] framework is an enhancement of the original DAFHEA [14] to extend its application to problems involving uncertain fitness functions. The enhanced framework is capable of solving complex real world optimization problems where the input information is expected to be generated by multiple models instead of a single model. Future research may investigate the possibilities of reducing the overhead incurred by developing and maintaining the regression models.

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