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A Comparative Review of Approaches to Prevent Premature Convergence in GA

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\textbf{ARTICLE INFO}

\textbf{ABSTRACT}

This paper surveys strategies applied to avoid premature convergence in Genetic Algorithms (GAs). Genetic Algorithm belongs to the set of nature inspired algorithms. The applications of GA cover wide domains such as optimization, pattern recognition, learning, scheduling, economics, bioinformatics, etc. Fitness function is the measure of GA, distributed randomly in the population. Typically, the particular value for each gene start dominating as the search evolves. During the evolutionary search, fitness decreases as the population converges, this leads to the problems of the premature convergence and slow finishing. In this paper, a detailed and comprehensive survey of different approaches implemented to prevent premature convergence with their strengths and weaknesses is presented. This paper also discusses the details about GA, factors affecting the performance during the search for global optima and brief details about the theoretical framework of Genetic algorithm. The surveyed research is organized in a systematic order. A detailed summary and analysis of reviewed literature are given for the quick review. A comparison of reviewed literature has been made based on different parameters. The underlying motivation for this paper is to identify methods that allow the development of new strategies to prevent premature convergence and the effective utilization of genetic algorithms in the different area of research.

1. Introduction

Genetic Algorithms (GAs) are search and optimization algorithms based on natural selection and genetics. Genetic Algorithms are one of the most popular algorithms in the categories of evolutionary algorithms. The basic principles of GAs were initially developed by Holland \cite{1} and further carried by De Jong and Goldberg. Goldberg and Michalewicz have given the detailed overview and implementation of GAs in various fields \cite{2} \cite{3}.

GA operates on a population of solutions represented by some encoding. During the implementation process of GAs every solution or individual is assigned a fitness which is the measure of GAs. The fitness of each individual is directly related to the objective function for optimization problem \cite{4}. Then, using the reproduction, crossover and mutation operators the individual population can be modified to a new population \cite{5}.

In GAs, the searching is iteratively guided by the fitness of the current parent generation. Whenever, we apply GAs for an optimization problem, it runs over thousands of individual, each represents a solution. The obtained solutions are evaluated and recombined to get offspring. It has been proven in \cite{1} \cite{2} \cite{3} \cite{6} that the previous generations details are only implicitly and partially preserved in the current generation. Hence, the regeneration can be hard to manage because of the following reasons:

- a) In regeneration an individual population appears that had already been seen in the search space.
- b) Genetic drift can affect the search for optimization in a wrong direction.
- c) As a result, a big part of the search space is left unexplored for the global optimum.

GAs has gained popularity because it can be applied in a wide range of problems, including multimodal function optimization, machine learning, pattern recognition, image processing, natural language processing, grammar inference \cite{4} \cite{5} \cite{8} and many more. But the behaviors of GAs have not been found as adaptive as expected. A common problem is that of premature convergence \cite{7}.

The primary interest of this paper is to show the detailed survey of different approaches proposed and applied for solving the premature convergence problem in GAs.

Section 2 offers a detailed description of different factors affecting GA search. In this section we discussed diversity of population and selective pressure. The aim of section 3 is to provide the theoretical framework to understand the genetic mechanism of evolutionary search process. This section briefly describes various theories such as Schema theory, Markov chain theory, and Statistical mechanism. This section also discusses and presents the important literatures to understand the genetic dynamics through Markov chain and statistical mechanics. The major parts of this paper contained in section 4. This section gives the detailed and comprehensive survey on approaches of handling premature convergence within GAs. In addition, strengths and weaknesses have been provided for each approach. The surveyed research works have been organized according to the publication. Surveyed researches have been summarized in section 5 using various parameters. The conclusions are drawn in...
2. Factors Affecting GAs

A genetic algorithm is a method of optimization based on Darwin’s natural selection theory. Darwin’s theory was based on survival of the fittest to refine a set of solution in an iterative manner [2]. Darwin’s theory says that evolution occurs if and only if three conditions met [9] (see Figure 1). The conditions shown in Figure 1 are necessary; it means that if due to any reason any one condition does not meet in the regeneration process, natural selection will cease at that generation.

Fig. 1. Necessary and Sufficient Conditions of Darwin’s Theory

GA starts with a set of solution which is represented by chromosome. By using an appropriate selection technique a solution from one population is picked depending on the fitness and used to form a new offspring. This process repeated until GA reached to the threshold or the maximum number of generations. Figure 2 shows the flowchart of the Standard Genetic Algorithms (SGA).

Fig. 2. Flowchart of Standard Genetic Algorithm [8]

The performance of GA largely depends on the crossover and mutation operations. Hence, adapting the suitable value of crossover and mutation is very important because it guides the search process and maintains the diversity in the population [19].

As discussed, GA is the population based search method. Therefore, selection of the population is the critical step because if population has not been selected in an intelligent manner then it will be difficult to get the appropriate answer to the problem. Typically, selection done at two places, one is in the beginning known as Initial Population selection or generation of random initial population and the second one is the selection of population for the next generation. Hence, before applying the selection or generation of population one has to take some factors into account as shown in the Figure 3. In this paper, we are considering two important factors, namely Population Diversity and Selection Pressure.

In order to reach to the global optima and explore the search space adequately, maintaining population diversity is very important. Also, it had been explained clearly that degree of population diversity is one of the main causes of premature convergence [27] [28]. It is necessary to select the best solution of the current generation to direct the GA to reach to the global optimum. The tendency to select the best member of the current generation is known as selective pressure. Selection pressure is also a key factor that plays an important role in maintaining genetic diversity. A proper balancing is required between genetic diversity and selection pressure to direct GA search to converge in a time effective manner and achieve global optima. High selective pressure reduces the genetic diversity; hence in this situation premature convergence can occur while the little selective pressure prohibits GA to converge to an optimum in reasonable time.

3. Theoretical Framework

This section provides an overview of different theories presented to address theoretical concerns related to evolutionary theory. There are various tools and methods available such as Schema theory, Markov chain theory, Dimensional analysis, Order statistics, Quantitative genetics, Orthogonal functional analysis, Quadratical dynamical systems, and Statistical physics. We have concentrated our study on three methods, namely Schema theory since it is considered as the fundamental theory, Markov chain theory since it is helpful in setting the boundaries for convergence and many more features and Statistical mechanics theory.

3.1. Schema Theory

Holland [136] presented Schema theory and it was accepted as fundamental theory to understand GA. It was widely used until the early 1990’s. It was claimed that in a population of ‘n’ chromosomes, GA process O(\(n^2\)) schemata into each generation. This property is known as intrinsic/implicit parallelism [1]. A schema is a similarity template used to describe a subset of string displays similarity at certain string position. It can be formed by ternary alphabet \{0,1,*\}, where ‘*’ represents a notation symbol shows the description of all possible similarities among string of a particular length and alphabet [137]. Length and order of schema were used to understand the working of schema theory. Length can be defined as the distance between first and last defined position in the schema while the order of schema is the number of defined positions. The GA modeled in schema theory known as canonical GA, which works on binary strings.
Although the schema theory was considered as fundamental theory, but it was criticized by many researchers. Some of them are pointed out as:

- It is not suitable to explain the dynamical or limit behavior of GA.
- Implicitly separation of problem to some extent.

The ignorance of these assumptions leads to the “Building block hypothesis” which clearly explains the behavior of GA [2]. In addition, Béryer [139] presented an alternative method to show how GA operates. Reeves and Rowe [138] presented the detailing of criticism of schema theory in their book “Genetic Algorithms: Principles and Perspectives (A Guide to GA theory). One more reason of replacement of schema theory is the suitability of Markov chain theory in the field of evolutionary algorithms (EAs).

### 3.2. Markov Chains Theory

The term Evolutionary Algorithm (EA) stands for a family of stochastic problem solver based on principles that can be found in biological evolution [140]. A stochastic algorithm can be considered as a random sequence of events \( E = \{ X_1, X_2, X_3, \ldots \} \) occurring in time, where each \( \{ X_i \} \) represents a random variable and the possible values that these variables can take are called the states of the system [138]. A Markov chain is a random sequence of variables \( X_0, X_1, X_2, \ldots \) satisfies the Markov property, i.e. the simplest kind of dependence within a stochastic process is when the distribution of time \( t \) depends only on what happened at the time \( t-1 \) [138]. In statistical terms, this is known as “memory less property” and can be defined as [142]:

\[
P(X_{n+1} = x | X_n = x_n, \ldots, X_0 = x_0) = P(X_{n+1} = x | X_n = x_n)
\]

Rudolph [141] presented the matrix property for Markov chain theory as given below:

A square matrix \( P = (p_{ij}) : N \times N \) is said to be

- Nonnegative if \( p_{ij} \geq 0 \), \( \forall i, j \in \{1, \ldots, N\} \)
- Positive if \( p_{ij} > 0 \), \( \forall i, j \in \{1, \ldots, N\} \)

A nonnegative square matrix \( P = (p_{ij}) : N \times N \) is said to be

- Primitive, if \( \exists n \in \mathbb{N} : P^n \) is positive
- Reducible, if \( P \) can be brought into the form (with square matrices \( C \) and \( T \))

\[
\begin{pmatrix}
C & 0 \\
R & T
\end{pmatrix}
\]

By applying the same permutations to rows and columns.

- Irreducible, if it's not reducible

- Stochastic, if \( \sum_{j=1}^{N} p_{ij} = 1 \), \( \forall i \in \{1, \ldots, N\} \)

A stochastic matrix \( P = (p_{ij}) : N \times N \) is said to be

Matrices that describe the transition probabilities in a Markov chain are stochastic matrices and will be referred to as a “transition matrix” [142]. In the next paragraph, we have provided a brief review of Markov chain theory which shows the applicability of this model.

Jeffery Horn [160] used Markov chains to analyze the stochastic effects of the niching operator of a niched GA. It was proved that a Markov chain is an effective tool to study the dynamic nature of a niched GA. The effect of “Absorbing Markov chain” and “Ergodic Markov chain” was shown to estimate the convergence of a niched GA [160]. Eiben et. al. [154] studied classical GA and simulated annealing (SA) and presented Abstract Genetic Algorithm (AGA) which generalize and unifies GA and SA. For the better clarification purpose, similarity and dissimilarity between classical GA and AGA was presented. In addition, Markov chain model was used to show the evolution of AGA. Leung et. al. [158] discussed the ability of the GA search by applying zero mutation probability and the relationships between premature convergence and effect of GA parameters were presented. Using Markov chain, the reasons of choosing zero mutation probability when analyzing premature convergence was demonstrated. Jun and Xinghuo [156] showed theoretical analysis to show the condition for the convergence of EA. The necessary and sufficient conditions for the convergence of EA to the global optimum were derived with the help of Markov chain theory to describe the limiting behavior. In addition, the upper and lower bound of the convergence rate required for the EA was explored. Stark and Spall [155] presented computable rate of convergence for GA via Markov chain analysis. Huang [145] extended the work presented in [146] by applying Markov model developed by Nix and Vose [144]. Reeves and Rowe [138] presented the applicability of Markov chain theory to describe the limiting distribution of simple GA in which they explained the effect of selection, mutation and crossover. In addition, modeling GA uses Markov chain theory was presented using suitable example. Auger [143] studied the convergence property of self-adaptive evolutionary strategies via phi-irreducible Markov chain. Basically, he used recurrent Markov chains to prove the existence of critical size of population, which was calculated using different parameters of algorithms and determines the convergence rate of the algorithm. Braak [148] presented Differential Evolution Markov Chain (DE-MC) which combines the essential features of differential evolution (DE) presented in [149] [150] [151] for global optimization over real parameter space with Markov chain Monte Carlo (MCMC) [152] [153] to solve an important problem of MCMC in real parameter spaces, namely that of choosing an appropriate scale and orientation for jumping distribution. Clark [142] applied this theory for modeling stochastic optimization algorithm. Hédéer and Bitó [157] implemented a GA to provide special site diversity to simultaneously optimize downlink and uplink the signal to interference and noise ratio (SINR) value in broadband fixed wireless (BFWA) access system. For the convergence analysis, i.e. to determine the maximum number of iterations they used Markov chains where every state represents a possible population containing different terminal station (TS) and base station (BS) assignment set as individuals. Suzuki [147] suggested large deviation principle approach for GA using Markov chains. Poli et. al [159] presented exact schema theory and Markov chain model Genetic Programming (GP) and variable length GA for the homologous crossover. In the absence of mutation the
relationship between approximate and exact schema theory for different forms of homologous crossover was shown.

Although Markov chain model was implemented for convergence analysis. But it was found limited by the complexity of calculations and it does not show the predictive description of genetic dynamics [140] [161].

3.3. Statistical Mechanics

It is a branch of mathematical physics. It can be used to study the state of uncertainty of a mechanical system using probability theory which works on the average behavior of the system [161] [162]. Prügel et. al [163] and Shapiro [164] described the GA using statistical mechanics. Shapiro [164] showed statistical formulation of GA to study the dynamics of GA searching and explained how statistical mechanics theory addresses the limitations of Schema and Markov chain theory. Two sources were identified in studying the dynamics of evolutionary approaches as given below [164]:

- The dynamics are stochastic – selection involves sampling and mutation and crossover involve probabilistic changes to the strings.
- The dynamics acts on the space of possible populations, a space of astronomical dimensionality

It was said that by deriving a deterministic set of equations, one can describe the evolution which must be sufficiently simple to allow for some type of study. There are many previous works available which proves the applicability of this theory [138] [165] [166] [167] [168] [169].

4. Approaches for Preventing Premature Convergence

This section gives the detailed explanation of the different approaches given by researchers for managing the risk of premature convergence in GAs. We have reviewed various literatures and presenting twenty four approaches with their strengths and weakness.

4.1. Crowding Method

Crowding was introduced by De Jong [38] to eliminate the most similar individual whenever a new one enters in a subpopulation. The aim of presenting this technique was to maintain the population diversity and to fix the premature convergence. It was applied in the survival solution step of genetic algorithms. The primary objective of introducing it in the survival selection is to decide which individual among those in the current population and their offspring individuals will pass to the next generation.

Mahfoud [12] presented a modified version of the De Jong’s strategy of replacing offspring of the most similar parents. The modified version of crowding was found effective in maintaining population diversity and preventing premature convergence. This approach works as follows:

Let, \( P_1 \) = Next generation population, \( P_i \) = First individual, \( c_i \) = Second individual (child), \( d(x, y) \) = distance between two individuals \( x \) and \( y \).

Then,

\[
\begin{align*}
\text{if } (d(p_1, c_2) + d(p_2, c_1)) &< (d(p_1, c_2) + d(p_2, c_1)) \\
P_1 &\leftarrow \text{Winner between } p_i \text{ and } c_i \\
\text{Else} \\
P_1 &\leftarrow \text{Winner between } p_i \text{ and } c_2 \\
P_2 &\leftarrow \text{Winner between } p_2 \text{ and } c_1
\end{align*}
\]

Three variations of crowding approach were presented, namely Standard Crowding [38], Deterministic Crowding [12] and Restricted Tournament Selection (RTS) [84]. Standard crowding was found to be limited in multimodal function optimization. Deterministic crowding (competition between children and parents of identical niches) approach was proposed by Mahfoud [12] to address the limitations [17] of the standard crowding approach suggested by De Jong [38]. The resulting order of complexity of deterministic crowding was \(O(N)\). RTS adapts standard tournament selection for multimodal function optimization. In case of RTS two elements from the population was picked to apply reproduction operators and then a random sample of CF (crowding factor) individual was taken from the population as in standard crowding. This procedure was repeated \(N/2\) times. The order of complexity of RTS is \(O(CF \cdot N)\) which varies from \(O(N)\) to \(O(N^2)\) depending on the value of CF [85].

4.1.1 Strengths and Weaknesses

This method was found effective for problems of all levels of difficulties, especially for multimodal function optimization. Using this, one can solve problems that are much more difficult than those solvable by traditional based hill-climbing. It uses a random sample of CF individual where CF is the crowding factor rather than fitness proportionate selection to resolve the selection pressure. There is no need of sharing function which is the key element of the sharing approach proposed by Holland [1] and further expanded by Goldberg and Richardson [16]. Selection of good sharing function needs a-prior knowledge of the objective function characteristics [74].

The problem with this approach is that it may lose lower optima which may lie on a crossover path to higher optima. Maintaining diversity in the population was the main goal of this approach and algorithm was successful to some extent, but stochastic errors by the low value of crowding factor affect the performance. Maintenance of bitwise diversity is the problem. It does not model the method by which a population arrives at a stable mixture of species, but instead strives to maintain the diversity of the preexisting mixture [75]. Handling replacement errors were the main concern with this technique. In addition, writing replacement rules are not an easy task.

4.2. Incest Prevention Algorithm

To avoid premature convergence of GA two approaches were introduced, namely the dynamic application of crossover and mutation operator and the population partial re-initialization [31]. The primary objective of applying the crossover operation is to exchange certain information between two parent chromosomes at the same time one must keep in mind that important information should not be lost. Hence, the selection of mating pair is a very important step in removing the insufficiencies of GAs.

Nicoara [31] and Eshelman and Schaffer [32] used Hamming distance as a measure for selecting the mating pair. The idea is that if Hamming distance is above a threshold then selects the parent chromosome for mating otherwise no. Before starting, the
threshold is set to a high value and during running of GA if no pair was chosen with the initialized threshold, the value can be decreased.

A detailed formula (equation (1)) was given for crossover operator and mutation operators (equation (2)). Equation (2) is the modified formula as given in [33]. Results of [31] were compared against three other genetic algorithms, namely NSGA-II [34], Canonic genetic algorithm and Elitist genetic algorithm. For the simulation a well-known job shop scheduling problem was implemented and performances of NSGA-DAR (Non-dominated Sorting Genetic Algorithm-Dynamic Application of genetic operators and population partial re-initialization) were tested on the n10 test instance and the obtained results were found encouraging.

\[
\Pi(x) = \begin{cases} 
1 & \text{if } D \text{ dominates } \mathbf{P}_1 \text{ and } \mathbf{P}_2 \\
\max(1 - \frac{k_1 x^t}{g}, 0.5) & \text{if } D \text{ dominates one parent and no dominance relation with other parents} \\
\max(0.5, \frac{k_2 x^t}{g}) & \text{if } D \text{ is dominated by at least one parent} \\
0 & \text{if no dominance relation exist between } D \text{ and } \mathbf{P}_1, \mathbf{P}_2, \text{otherwise.}
\end{cases}
\]

(1)

Shimodaira [78] employed the concept of incest prevention and proposed a new genetic algorithm Diversity Control-oriented Genetic Algorithm (DCGA). Incest prevention was applied in survival selection and this increased the diversity by giving individuals with father Hamming distance from the best solution a higher probability of survival [77]. Matsui [79] incorporated two new selections scheme, namely correlative tournament selection (CTS) for reproduction and correlative family based selection (CFBS) for survival. CTS choose the mate with higher fitness and Hamming distance from a set of candidates.

\[
\Pi(x) = \begin{cases} 
1 & \text{if } M(x) \text{ dominates } x, N(x) \text{ or } M(x) \text{ not valid} \\
\max(1 - \frac{k_2 x^t}{g}, 0.5) & \text{if no dominance relation exist between } x \text{ and } M(x)
\end{cases}
\]

(2)

A similar approach was proposed by Fernandes and Rosa [80] called as negative assertive mating into GA with varying population size (GAVaPS). They showed that GAVaPS mimic natural species’ behavior by selecting parents, according to parenthood or phenotype similarity. A strategy of choosing different crossover methods was proposed in [81]. It was suggested that one can choose the appropriate crossover mechanism to avoid premature convergence on the basis of the couple’s Hamming distance. In contrast to incest prevention, three new methods of selection of mating pairs for GA were suggested [82]. It was tried to mimic inbreeding in nature which restricts individuals to mate with one separated by a minimal Hamming distance. Experimental results showed that the proposed approach could enhance exploitation, reduce the disruption of schema and perform better than a conventional genetic algorithm (CGA) and incest prevention in certain problems.

4.2.1 Strengths and Weaknesses

This approach works effectively when combined with multiple crossovers on multiple parents. Therefore, it was given a name Multiple Crossover on Multiple Parents with Incest Prevention (MCMPIP) [76]. It was found effective for multimodal function optimization. In addition, it avoids premature convergence and improves the solution at the expense of convergence speed [77]. The approach suggested in [31] is helpful in two fold. First, dynamic application of genetic operators takes the advantages of the identification of the operators to promote the beneficial effects of all the available operators during all evolution. Second, the population partial re-initialization achieves the main goal only in the critical situation, when the risk of premature convergence is big enough which saves the computational cost.

The drawbacks of this approach are: it suffers with the problem of an increase in computation cost for calculating the individual’s Hamming distance. Comparison of individual Hamming distance with threshold increases the computational cost.

To overcome the problem (additional cost requires in computing the individual’s Hamming distance) of incest prevention, Craighurst and Martin [83] considered the family tree to disallow incest by adhering to the ancestry based incest tree. The level of incest law helps in allowing or disallowing the mating among family members. Experimental results prove its effectiveness on solution quality and diversity maintenance. This approach showed the slow convergence rate and was sensitive to the mutation rate while incest prevention yielded a satisfactory outcome.

4.3. Scheduled Sharing Approach

Palmer, and Stephen [15] presented a nature inspired approach known as Scheduled Sharing to prevent premature convergence by applying an extension called as sharing to standard GA. This approach handles premature convergence by requiring organisms within a niche to compete with their neighbors. Initially, one picks the niche size as large as possible to encourage the organism to spread across the search space. Thereafter, exponentially decays and allow gradual clustering of the organism. This approach is the extension to the standard sharing algorithm [16] [17]. The approach given in [15] over [16] [17] is the modification of fixed niche sharing to handle more difficult and realistic test function.

Palmer and Stephen [15] not only worked on sharing algorithm, but also they worked on sampled sharing, which removes \(O(n^2)\) communication between processors by standard sharing. Experimental results prove that scheduled sharing can delay the convergence as long as require which continues the search process efficiently. To calculate the convergence of a population search space can be divided into subspace or buckets. Suppose, \(x_i\) represent the percentage of the population of \(i^{th}\) bucket, then the percentage of the total population can be calculated as:

\[
\text{Percentage of Total Population} = \sum_i x_i^2
\]

(3)

Equation (3) is an important measure to decide the convergence of a population, higher value represents that the population is more concentrated in one or a few buckets while the lower measure shows that populations are more distributed across the buckets. Two classes of landscapes were used, namely Easy and Difficult landscapes generated by the weights \(W\) of the sub-functions. Easy landscapes are those have a small number of local optima with wide basins of attraction while difficult landscapes have just opposite of the easy landscapes. Experimental results prove that both Scheduled Sharing and Non-
Sharing algorithms performed better than the Random Search algorithm at all population sizes on both classes of landscapes and prevent premature convergence of a genetic algorithm [15].

4.3.1 Strengths and Weaknesses

This approach was found effective on large population genetic optimization of rugged two dimensional landscapes. It is helpful in improving offline performance by requiring organism within a niche to compete with their neighbors. It was shown that it allows the application of sharing for complex problems where there is no definable best niche size, without violating black box principle in the calculation of niche size. This approach takes the advantages of simulated annealing in calculating the niche size by applying an exponential decrease in the schedule. It reduces the computational cost by allowing gradual clustering of organism in areas of greatest interest. It provides flexible control of the convergence of a GA. This approach allows the feedback to define a problem dependent sharing schedule.

Although this technique outperformed over random search and standard sharing techniques. But still there are some limitations.

- The key concern of this approach is finding the appropriate schedule for applying sharing.
- It requires prior knowledge to write the appropriate sharing function which is a difficult task because of a real optimization problem, no information about the search space and distance between the optima is generally available.
- It was found that this approach improves the performance of standard sharing approach by incorporating the concept of cluster analysis and dynamic niching. But still the computational time to obtain the fitness of individuals dominates the computational cost of comparisons.
- It was accepted that to make the schedule sharing more effective, a difficulty detector is needed which might be used to decide upon the schedule [15].

4.4. Migration Model and nCUBE Based Approach

Balakrishnan [24] presented a Migration Model to maintain the diversity and variation in the genetic property of individual population which keeps the GA search process alive. The behavior of migration model is very similar to an ordinary communication model, which allows individuals to migrate from one population to another. Therefore, migration model helps in finding global solution and prevent premature convergence. Features of nCUBE model were used to design the migration model because it has several reasons as listed in [24].

Three different scenarios namely Best Individual, Mean Individual and Random Individual were suggested to migrate an individual from one population to another. It was observed that once an individual reached to target population the migrated population gets welcomed in a different ways, such as Warm, Neutral and Cold welcome. Cold welcome was used for the simulation in [24].

Two experiments were conducted to ensure the effectiveness. The first experiment tests the suitability of migration of individual based on different migration strategies while the second experiment gives the answer to the scalability issue of the system [24]. The simulation results of both the experiments were found effective to prevent premature convergence. In addition, it improves the performance of GA to solve the complex optimization problem.

4.4.1 Strengths and Weaknesses

The present approach was powered by maintaining independent population and allowing inter population migration to maintain the diversity and alleviate premature convergence. nCube model was used for the experiment which improves the performance of GA search. In addition, it was evident that this approach is one of the best approaches for complex optimization problems.

But still there are some key issues in applying this technique. First, setting the migration interval play an important role because both large and small intervals being desirable. If the migration interval is too large then it increases the possibility of premature convergence whereas the small migration interval increases the interaction among the population which results in an increase in the computation time. Second, making decision for an individual to migrate out of a population is a crucial factor. Third, how to choose number of independent populations because solution status will be affected if more independent populations are available.

4.5. Cooperation Based Approach

Cooperation based genetic algorithm to prevent premature convergence was presented in [18]. The fact which was shown in [18] is that the hardness of a problem inherently related to the way we represent the problem. Therefore, the difficulty level of the problem can increase or decrease based on the representation techniques used. To prove this fact 1’s counting problem (max1 problem) was implemented and the difficulty level showed using different representation techniques.

The idea of cooperating genetic algorithm says that instead of using only one representation apply varieties of GA concurrently each optimizing the same object but using its own representation. In other words, after a fixed number of generations, each GA provides other GA’s with its best local optima found. Experimental results were found effective for solving those problems which was either difficult or cannot be solvable by sequential GA.

4.5.1 Strengths and Weaknesses

Fonlup, Preux, and Robilliard [18] claimed that the varieties of representation techniques could be helpful in avoiding premature convergence. Experiments were conducted to prove the claim which showed the effectiveness of the presented approach. Various representation techniques are the key concept in removing the correlation between the neighboring individual and maintains the diversity of the population. This approach improves the speed without applying parallelism. An exchange in individuals creates new paths of research and allows GA to choose the new created paths.

The weaknesses with this approach are: a) selection of the appropriate representation techniques to maintain the diversity in the population because how one can say that the new representation technique will work effectively. (b) In the presented model [18] it was considered that after 20 generations each GA broadcasts to every GAi, it’s 10 individual, but what is the guarantee that same assumption will work for any other problem. (c) Dealing with more than one representation is not an easy task.

4.6. Syntactic Analysis of Convergence

The average Hamming distance of a population was used as a syntactic metric to prevent premature convergence and predict the convergence time [29] [30]. The primary goal of using syntactic metric is to get the probabilistic bound on the time
convergence of GAs. It was shown that the analysis of flat function provides worst-case time complexity of static functions which develops a theoretical basis for the premature convergence.

It was observed that mutation is one of the ways of maintaining diversity of the population [29]. But the high mutation rate raises problems. In addition, it also increases the risk to destroy good schemas as bad ones. These observations were worked as motivational factors to come-up with new approaches.

Therefore, it was suggested that in place of increasing mutation rates, pick an individual and add its bit complement to the population [29]. The aim of doing this was to ensure that each allele is present and the population spans the whole encoded space of the problem. The presence of complementary individuals also makes a GA more resistant to deception. Depending on the assumptions about the search space, one can pick individual to be complemented. Various approaches were suggested for doing this such as probabilistic, random selection and others. It was found that randomly selecting the individual to be complemented makes the least number of assumptions and it may be the best strategy in the absence of other information. For conducting the experiment minimum fitness individual were replaced by either the complement of a randomly picked individual in the current population or the complement of the current best individual.

If \( l_i \) represent the number of bits needed to represent variable \( i \) in a deceptive problem, then the functions used in [29] can be described using equation (4).

\[
\text{Deceptive}(x_i) = \sum_{i=1}^{l_i} \begin{cases} x_i & \text{if } x_i \neq 0 \\ 2^{l_i+2} & \text{if } x_i = 0 \end{cases}
\] (4)

For 10-bit and 20-bit deceptive problem \( \text{Dec}_1 \) and \( \text{Dec}_2 \) were used which can be represented as:

\[
\text{Dec}_1: \text{Deceptive}(x_1^2, x_2^2)
\] (5)

\[
\text{Dec}_2: \text{Deceptive}(x_1^2, x_8, x_5, x_3, x_4^2)
\] (6)

In equation (5) and (6) superscripts denote the number of bits needed for each variable in \( \text{Dec}_1 \) and \( \text{Dec}_2 \) respectively.

The results obtained experimentally were compared with classical GA (CGA) and a GA with compliments (GAC), which proves that GAC easily reached to the optimum whereas CGA fails to find the optimum.

Louis and Gregory [29] accepted that the GAC does not give guarantee for an optimal solution to all problems, whereas Messy GA [45] can make much stronger guarantees. But the drawback of the messy genetic algorithm is its computational complexity. It was also shown that GAC performed much better than the CGA on Ackley’s One Max [46] and no worse on the De Jong test suite [38]. Also, it was agreed that GA hard problems that are both deceptive and epistatic may need masked crossover or other length independent recombinant operators to be solved successfully using complements [47].

A time convergence model was presented to predict the time to convergence [29]. The string similarity assumption was made to develop this model. This model works successfully for unimodal function. The behavior of multimodal function, genetic drift and others can also be predictable using this model. The model starts with the equation (7) which computes the change in Hamming average per generation.

\[
h_{t+1} = f(h_t)
\] (7)

Where, \( h_t \) = Hamming average in generation \( t \) and \( h_{t+1} \) = Hamming average in the next generation.

Two observations were made for choosing the function \( f(h_t) \). The first observation says that “the schema theorem along with the similarity assumption indicates \( f(h_t) \) is linear (see equation (8)).

\[
h_{t+1} = ah_t + b
\] (8)

The second observation says that “without mutation the final Hamming average is zero” which implies that \( b = 0 \). By putting this value in equation (8), we will get.

\[
h_{t+1} = ah_t
\] (9)

The general solution of the equation (9) is given as:

\[
h_t = \begin{cases} \frac{1}{a} h_0 & t = 0 \\ a^t h_0 & t > 0 \end{cases}
\] (10)

The value of \( a \) can be obtained by keeping track of the Hamming average, while running a genetic algorithm [29]. It was shown that the model works effectively and a good prediction of time complexity is possible.

4.6.1 Strengths and Weaknesses

The primary strength of this approach is that it provides the time limits to the convergence of the GA. In addition, combining fitness prediction with Hamming average prediction provide the details about how much progress is possible with a GA along with bounds on how much remaining work can be done. Syntactic analysis model forms the basis to see the effect of selection and mutation on the behavior of GA. It was evident from the result that good predictions of time complexity are possible, even from a rough model that uses easily computable syntactic information.

The problems with this approach are: mutation rate affects the performance of the GA in the later generation. In addition, computing upper bound and lower bound is not cost effective process. It was accepted that qualitative prediction may not be possible with syntactic information [29][30].

4.7. Pygmy Algorithm

Rayan [43] described a new selection scheme to reach to optimal solution. Actually, he suggested Pygmy algorithm to deal with multi-objective problems. This algorithm was originally applied to the problem of evolving minimal sorting networks, which must not only be able to sort numbers, but also in as few less number as possible [43]. This approach uses two fitness functions and maintains two lists of parents, one for each criterion of the problem. It was clearly shown that if the population using Pygmy algorithm then the chances to converge to an optimal solution is more compared to other techniques uses a fitness function which is a sum of the assumed criteria. In [44]
Pygmy algorithm was applied for handling premature convergence in the case of grammar induction. In order to implement Pygmy algorithm and to select the resulting grammar with minimum number of rule principle was used which was considered as another fitness criterion to create a second parent list.

4.7.1 Strengths and Weaknesses

This approach uses a less computational cost compare to the previous approaches such as Crowding [38], Sharing [16], restricted mating, spatial mating and others. It reduces the risk of losing genetic material compared to the methods which reward parsimony [86] or try to combine the two measures [87] run the risk of individuals trading off various parts of the fitness function. Pygmy algorithm maintains two lists of fitness which removes the possibilities of inbreeding between individuals with similar performance. In addition, this approach avoids the chances for individuals to breed with close relatives like siblings or even parents of opposite sex.

The weakness with this approach is that in some situations it is not well suited for multi-objective problems. In order to get the solution of multi-objective problems an individual must solve two or more smaller problems of the same nature to solve the main problem. Pygmy algorithm does not address some of the sub problems simply because individuals in the same group could not mate with each other.

To address the above issue Ryan [43] presented Racial Harmony. It was evident in [43] by applying races among the individual in the same group the performance of Pygmy algorithm was improved. Racial Preference Factor (RPF) was used as a measure of probability that an individual will choose an individual from another race when selecting a mate.

4.8. Adaptive Probability Based Approach

In many previous works [68] [88] [89] [90] the idea of adapting crossover and mutation was applied for the betterment of GA performance. Schaffer et al. [88] implemented a crossover mechanism where the crossover points were chosen depending on the performance of the generated offspring. Davis [89] suggested mechanism of adapting probabilities based on the performance of the operators. He allotted higher probabilities to the operators which create and cause the generation of better strings. The technique presented in [89] was developed in the context of a steady state GA [90]. Fogarty [90] showed the effect of varying mutation rate over generations. In his work he showed that the exponential decrease in mutation rate shows better performance for a single application. Whitley et al. [68] presented that the probability of mutation is a dynamic variable parameter which can be determined by the Hamming distance between the parent solutions. It was evident in [68] that adapting mutation probability dynamically significantly improves the performance.

Adaptive Genetic Algorithm (AGA) (Srinivas and Patnaik, 1994) is one of the efficient methods for optimizing the multimodal function [19]. This approach differs from the existing approaches [89] [90] as crossover and mutation probability was determined for each individual as a function of its fitness. Whitley et al. [68] implemented adaptive mutation approach in the context of steady state GA, whereas AGA is based on generational replacement strategy. AGA employ best solution protection strategy which resolves the high level disruption while steady state GA applies populationary elitism in which there is no need to protect the best solution.

To prevent premature convergence during the implementation of AGA a probability based approach was adopted which varies the probabilities of crossover and mutation depending on the fitness value of the solution. A detailed design was given for suitably choosing the crossover ($P_c$) and mutation ($P_m$) probabilities [19]. The objective of this approach was not only to present a mechanism to choose crossover and mutation probabilities but also present a suitable technique to prevent the GA from getting trapped at a local optimum. It was shown that the adaptive value of crossover and mutation should be from 0.0 to 1.0 and 0.0 to 0.5 respectively. It was proven that the low value of the crossover probability and high value of the mutation probability lead to premature convergence or a shift from genetic algorithms to random search. These facts help in preventing premature convergence of the GA.

Extensive experiments were conducted to show the effectiveness of AGA. For the experiment various optimization functions were used such as De Jong’s functions known as spiky function ($f_5, f_6, f_7$), Order-3 Deceptive problem, Traveling Salesman Problem (TSP) and many more. The outcome of the experiment was found to be encouraging.

4.8.1 Strengths and Weaknesses

The main strength of this approach is that it provides a basis for adapting the crossover and mutation probability depending on the fitness value of the solution, i.e. for low fitness value assign high value of $P_c$ and $P_m$ and vice-versa. AGA protects best solution from the high levels of disruption which receives zero $P_c$ with small amount of $P_m$. This information is helpful in reducing the extra computation cost of applying crossover. In this approach a criteria were set as “all solution with a fitness value less than the average fitness value of the population have $P_m = 0.5$ which shows that sub-average solutions are completely disrupted and totally new solutions are created” this criteria was also found effective in saving the computational cost and helps GA not to get trapped at a local optimum.

As discussed, this approach gives a concrete basis for selecting the crossover and mutation probability. It was found that most of the individuals were trapped at a local optimum because the exploration effect of mutation was low. To address this issue Jung (2009) presented a Selective Mutation based approach (discussed in section 4.20) [35].

4.9. Social Disaster Technique

Grefenstette [91] attempted Travelling Salesman Problem and suggested the concept of greedy crossover. The crossover operator implemented was called “greedy crossover” because it always prefers locally cheaper paths. Picking a locally cheaper path was based on the knowledge about the problem and showed promising results than more expensive one. It was evident in [91] that crossover based on knowledge, improve convergence, but it might derive to local minima and leads to premature convergence.

To address the above issue Social Disaster Technique (Kureichick, Miagkikh and Topchy, 1996) was suggested which considers the whole population [42]. Kureichick et. al. [42] directed their efforts to avoid the following two reasons:

- Immoderate crossover greediness
- Low influence of random factors intended to make the population get out of local minima.
The concept of super-individual was suggested which takes over the charge of the population. Catastrophic operator was used in this method to maintain the population diversity. Packing and judgment day operator was used for the experiment. The working of packing operator is to apply separation, i.e. separates all the tours have the same length and only one remains unchanged, whereas in case of judgment day operator only the best instance remains unchanged. In addition, warp operators were used in population, which introduces partial or full randomization.

An experiment was conducted on the Travelling Salesman Problem (TSP) using the benchmarks given by Oliver’s 30 [93], Eilon’s 50 and Eilon’s 75 [92]. It was shown that the thirty town problem was solved easily while 50 and 75 town TSPs was found harder in the presented approach. The result of 5 and 75 TSPs was compared against the results given in [92] and it was observed that the suggested approach showed the shorter tour.

4.9.1 Strengths and Weaknesses

The method described above brought significant improvements, some of them are: introduced new operators provide the facility to use the population pool effectively. It is suitable for the small population size problem. Warp operator brought the partial or full randomness which helps the population to get out of the local minima. Decrease in the greediness of the crossover operator increases the variation among the population.

The weakness of this approach is: there was no sufficient information is provided to tell up to what extent greediness of crossover should be decreased. This approach does not fit for complicated problems, i.e. for large population size problems this approach consumes more time to converge.

4.10. The Island Model Genetic Algorithm

Vose [94] and Whitley et. al [95] presented exact models of an SGA using infinite population assumption. One derivation of these models was presented in [95] while it was extended to look at finite populations using Markov models [94]. It was found that the weakness of the infinite population model was that one could not take into account the sampling bias introduced by using finite populations whereas the finite population Markov model was too expensive to execute very small problems. Therefore, it was suggested that using a distribution taken from a finite population one can introduce finite effects into the infinite model.

Whitley et. al. [69] suggested an application of the infinite population model to see the behavior of multiple SGA running parallel. These SGA was allowed to exchange information time to time. It was shown in previous research that parallel GA yield better performance than serial [70] [71] [72]. Island Model was used to implement GA on both serial and parallel machines [65] [66] [67] [68]. In an island model each machine executes a GA and maintains its subpopulation for directing the search. It was observed that an island model uses multiple populations for each machine in parallel implementation which helps to preserve genetic diversity, since each island can potentially follow a different search trajectory through the search space [69].

In order to implement island model two parameters were introduced, namely migration interval and migration size. The number of generations between migrations was represented by a migration interval while migration size is the number of individuals in the population to migrate. Each machine in the island model periodically exchange a portion of their population using a process known as migration.

The function $G$ was used to represent trajectory of the infinite population SGA, which works in a vector $p$ [94]. $p^t$ represent the population at the time $t$ which shows the sampling distribution of a finite or infinite population. The search process enters into new generation using function $G$ as depicted in equation (11).

$$p^{t+1} = G(p^t)$$

(11)

Where, $p^{t+1}$ represent next generation an infinitely large population. It also represents the expected sampling distribution of a finite population [69]. The construction details of $G$ was given by Whitley [96]. It was observed that the behavior of the function $G$ was exactly correspond to the SGA provided the infinite large population. The details about the finite and infinite population were covered in varying population size (small and large) [69].

Whitley et. al. [69] used function $G$ to develop an idealized Island GA. The idea was total population was divided into a number of subpopulations called as “islands”. These islands are responsible to execute a version of GA on each subpopulation. Each island manages a separate copy of $G$ for SGA using 1-point crossover. The crossover rate chosen was 0.6. It was found that migration occurs between subpopulation in a restrictive manner using temporal and spatial considerations.

![Fig. 4](https://example.com/island_model)  
*Fig. 4: Basic organization of Island model using six islands. For first migration after X generation migrate most fit 5% from Island-1 to Island-2 (immediate neighbor on the left). As soon as Island-2 receive the migrated data immediately delete the least fit 5% of the population the same process repeated to every other island excluding itself.*

The migration strategy applied in Island model is depicted in Figure 4. We have used six islands to demonstrate the working. From the Figure 4 it is clear that all the islands are arranged in a ring. For the first migration Island-1 migrate most fit 5% after X generation to its immediate neighbor in the left i.e. Island-2. As soon as Island-2 receives the migrated information it deletes the least fit 5% of its population. For the second migration Island-2 migrate the most fit 5% to its immediate neighbor to its left i.e. Island-3 after X generations. This migration process occurs every X generation until each island has send one set of strings to every other excluding itself then the process repeated [69].
Experiments were conducted using two separable and non-separable problems. Two separable problems were chosen because they represent examples from a well-known set of linearly separable test problem while the non-separable problem was chosen to demonstrate how the added interaction between parameters can affect the GA performance. GENITOR was used as a search engine for the experiment [73]. The experimental results shown in [69] prove that the Island Model genetic algorithms outperform and maintain the population diversity, avoid premature convergence and reach to the global optima.

4.10.1 Strengths and Weaknesses

Island GA helps to preserve genetic diversity because every island potentially follows a different search trajectory through the search space [69]. An abstract model was presented which support it to outperform single population models on linearly separable problems. It is suitable for large total population size problems.

This model drawn the attention of researchers since it offers some degree of independence and hence explore different regions of the search space while at the same time sharing of information allowed. The concept of migration was found effective which helps in maintaining population diversity and the key to resolving premature convergence.

The limitation of this approach is the migration interval, i.e. after how many generations one should apply migration of best fit individual from one island to another. Another factor which was not defined clearly was migration size. For the presented experiments 5% best fit individual was migrated but what is the guarantee that it will be suitable for all types of optimization problems. It was accepted that the presented approach could show very complex and unexpected behavior when applied to separable problem.

4.11. Shifting Balance Theory in Dynamic Environment

Wright [20] presented Shifting Balance Theory (SBT). The ill behavior of evolutionary system was observed, i.e. getting trapped at a local optimum. It was explained that for a fitness landscape there might be two peaks or even more. In rugged field of this character (very many peaks), selection will easily carry the species to the nearest peak, but there may be innumerable other peaks which are higher but separated by “valleys”. Wineberg [97] showed he thought of evolution as a mechanism by which the species may continually find its way from lower to higher peaks. The term “trial and error” was suggested as exploratory mechanisms which take the species to the next higher peak, despite the harsh selection pressures of the valley forcing the species back to the original hill [98].

Hartl and Clark [98] presented a good summary of SBT in their book “Principles of Population Genetics”. They suggested that subdivision of the population into a set of small semi-isolated demes increases the possibility for demes to explore the full range of the adaptive topography and reach to the highest fitness value on a convoluted adaptive surface. Reduction of large population into smaller demes reduces the possibility of random genetic drift of allele frequencies and provides the flexibility to explore the adaptive topography more or less independently. It was found that the random genetic drift in any subpopulation leads to temporary reduction in fitness. This could be rectified by selection in a large size population which leads the search process to pass through a “valley” of reduced fitness and possibly end up “climbing” a peak of fitness higher than the original. It increases the size if a subpopulation reached to the adaptive peak on the fitted surface. In addition, migration of individual genes will be higher and therefore the favorable gene combinations are gradually spreading throughout the entire set of subpopulations by means of interdeme selection.

Hartl and Clark identify three distinct phases and demonstrated the role of each phase [98] as depicted in Figure 5.

Wright’s Demes was one of the significant contributions in justifying the introduction of a novel GA implementation particularly for parallel GA. It was observed that both fine-grained and coarse grained model (Island model) is presented when creating parallel GA. These techniques use “demes” to indicate a mating pool of individuals locally clustered geographically [97]. We have already discussed Island model (section 4.10) in which each processor runs an SGA on its own subpopulation or deme.

Cohoon, Martin and Richards [99] initiated the association of the Wright’s SBT with Island model style of parallel GA. Cohoon et. al. [100] justified the applicability of island model. He presented the idea of punctuated equilibria for parallel GA using Wright’s theory [20] and Eldrege and Gould’s theory [101] of punctuated equilibria. Similar to island model, many researchers worked on a fine grained model using Wright’s theory [102] [103] [104] [105] [106] [107]. In this case, each processor is given only a single member of the population. Each member is permitted to mate with its nearest neighbors or possibly within a small neighborhood. The basis of this design was the fact that fast communication may be achieved among neighbor. The neighbors can be arranged using various topologies [97]. Here, diffusion of genetic materials occurs through neighborhood overlap which was identified as demes.

Although Wright’s theory was accepted and applied in several applications. But this theory was found controversial. Coyne, Barton and Turelli [108] questioned about the validity and
importance of SBT. Coyne et al. [112] studied both theory (Fisher’s theory and Wright’s theory) and rejected the idea that “Wright’s SBT theory has played a major role in adaptive evolution”. Instead Darwin’s and Fisher theory [111] “adaptation is natural selection acting on differences among individuals without genetic drift, population subdivision and differential migration playing the vital roles hypothesized by the SBT” was accepted [112]. Peck, Ellner and Gould [109] and Wade and Goodnight [110] presented a framework to support the SBT theory and claim that the dismissal of SBT is premature.

As discussed the SBT theory was controversial still Oppacher and Wineberg [21] presented the modified version of SBT in which the original goal was to avoid defects inherent in its original formulation and prevent premature convergence in GAs. The new algorithm implemented is known as Shifting Balance Genetic Algorithm (SBGA) which improves the performance in a dynamic environment. Two different types of population groups, namely Core group and Colonies were used in the modified SBT. The core group is the large central population, while a smaller group called as colonies. The main job of colonies is finding the search space of the landscapes that the core group is unable to find. In this way one can prevent premature convergence. To develop a complete system few measures were considered such as population diversity (measured by equation (12)), and distance (measured by equation (13)). Distance represents the distance between a single chromosome and population in gene space, is a modification of the diversity measure [21].

\[
\text{Diversity}(P) = \frac{1}{\ln(N-1)} \sum_{i=1}^{N} \sum_{j=1}^{N} \text{H}D(p_i, p_j)
\]  

(12)

\[
\text{Distance}(c, P) = \frac{1}{\ln N} \sum_{i=1}^{N} \text{H}D(c, p_i)
\]  

(13)

Where, \(L\) represents the length of a chromosome, \(N\) is the size of the population \(P\), \(p_i\) is the \(i^{th}\) chromosome in the population, \(\text{H}D\) represents the Hamming distance function. The term \(\text{H}D(c, P)\) shows the Hamming distance between a single chromosome \(c\) whose distance is being measured and population \(P\).

One more measure was considered known as containment (equation (14)). It was considered as a measure of the extent to which one population is contained within another.

\[
\text{Containment}(A, B) = \frac{1}{M} \sum_{i=1}^{M} \text{WithinDistance}(a_i, B)
\]

\[
= \frac{1}{MN} \sum_{i=1}^{M} \sum_{j=1}^{N} d_p(a_i, b_j)
\]  

(14)

Where,

\[
\text{withinDistance}(c, P) = \frac{1}{N} \sum_{i=1}^{N} d_p(c, p_i)
\]

\[
d_p(a, b) = \begin{cases} 
1 & \text{if} \quad \text{Distance}(a, P) < \text{Distance}(b, P) \\
0 & \text{otherwise}
\end{cases}
\]

\(a_i\) and \(b_j\) is the \(i^{th}\) and \(j^{th}\) member of the population \(A\) and \(B\) respectively, and \(M, N\) represents the population size of \(A\) and \(B\) respectively.

Table 1 presents the state of affairs of the colony and the remedy suggested to keep colony away from the core. The measure of containment (containment function) can be used for this purpose which can be calculated using equation (14).

![New Population after reproduction of colony](image)

Fig 6: Structure of new population after reproduction of colony.

It is depicted in Figure 6 that when the colony is ready for reproduction the new population was divided into two halves. Member’s of the first part was selected using the regular fitness function, whereas the member of the second part was selected based on the distance from the core group using the fact that the greater the distance higher the fitness.

Unlike the SBT theory of immigration of colony members, sending members from the core to the colony, Oppacher and Wineberg [21] suggested no migration from the core to the colony. Although the colony still sends migrants back to the core. It was observed that colony members were selected randomly, stochastically or using elite subgroup for immigration into the core based on their fitness. Adding the colony members into core resulted in a temporary increase in the size which could be reduced to normal size during the reproduction by exerting selection pressure on the core. As the migration of colony members disrupts the core group (see Table 1), hence to rectify this issue, time (migration interval) was given for each colony to evolve potentially useful members.

To ensure the effectiveness of the implemented approach two experiments were conducted using the \(F2\) function from the De Jong test suite composed with the one dimensional version of the Griewangk function known as \(F8\). The combined function was used for the experiment called as \(F8F2\) function. The primary aim of the first experiment was to check the premature convergence while the second experiment was conducted to ensure the dynamic environments. The experimental results were found significantly better and encouraging than the standard GA to prevent premature convergence and also run in a dynamic environment.

<table>
<thead>
<tr>
<th>S.N.</th>
<th>State of affairs</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>If colony remains completely outside the core.</td>
<td>No action, freely evolve</td>
</tr>
<tr>
<td></td>
<td></td>
<td>using regular fitness function, allow looking potentially interesting area ignored by the core.</td>
</tr>
<tr>
<td>2.</td>
<td>If colony moves or trying to move into area searched by the core group.</td>
<td>Pushed colony away from the core group.</td>
</tr>
<tr>
<td>3.</td>
<td>If the colony is at the outside boundary of the core</td>
<td>Push colony gently away</td>
</tr>
<tr>
<td>4.</td>
<td>If the colony is almost entirely contained in the core</td>
<td>Apply strong pressure to push it into novel territory</td>
</tr>
</tbody>
</table>

Table 1: State of affairs of colony and their remedy
4.11.1 Strengths and Weaknesses

The main strength of this approach was to resolve premature convergence and up to an extended this approach works successfully. This approach was strengthened by providing the concept of colonies to search areas of fitness landscapes missed by the core group. The flow of colony member of the core group after a time interval increases the diversity of the core and helps in preventing premature convergence. In addition, it was found suitable in a dynamic environment. In SBGA, if the peak shifts away from the core group the colony are already in an area away from the core.

Oppacher and Wineberg [21] assumed no migration between colony and core and remedial approaches were suggested for the colony member if enters in the core search area still some of the colony member jumps into the core group which might be problematic in managing the diversity. This approach suffers with the problem of migration interval, i.e. when to migrate the colony member. Lastly, it was accepted that there is no dynamic model available of the dynamic environment.

4.12. Random Offspring Generation Approach

Rocha and Jose [22] presented a new approach to remove the insufficiency of GA. This approach is known as Random Offspring Generation (ROG). By looking at the flowchart shown in Figure 7 one can easily understand the working of ROG process. The flowchart clearly demonstrates the reason of occurring of premature convergence in GA. The answer is for a large amount of individual is using the similar genetic material during the reproduction process. In other words, we can say that for the crossover operation same genetic materials were used. In this case reproduction process will not show any effect because the offspring bred are nothing but simply clone of their parents.

Therefore, to prevent premature convergence in GA, ROG process could be applied before applying the reproduction operations. The idea is first test the similarity of genetic material, if genetic materials found similar than generate the random offspring which will code a random solution for the problem otherwise apply reproduction operation. In [22] two different strategies were used in ROG namely 1-RO (one offspring per two parents) and 2-RO. In case of 1-RO, only one offspring will be generated and other one obtained from their parent to reach to the result while in 2-RO both the offspring will be randomly bred.

ROG strategy was applied to a selected set of Travelling Salesman Problem (TSP) [23]. For the experiment TSP problem was used as a benchmark. The data obtained from the experiments were compared with other approaches such as Adaptive Mutation Rate (AMR) and Social Disaster Techniques (SDT) on various crossover operators. The comparative results prove that the technique presented in [22] is far better to prevent premature convergence to local optima and hence improve the performance of GAs.

4.12.1 Strengths and Weaknesses

It was shown experimentally that the presented approach yields the best results compared to adaptive mutation rate (AMR) (section 4.20) and social disaster technique (SDT) (section 4.9). As observed AMR showed the solution with a similar degree of quality as obtained by SGA whereas SDT in either of its form didn’t show any improvement in the solution. Another important merit of this approach is the simple nature in implementation.

Rocha and Jose [22] accepted that the weakness of this approach is the computational overheads disregarded since there is no change in the selection methodology compared to other approaches such as Crowding [38] (section 4.1), Sharing [16] and Scheduled Sharing [15] (section 4.3).

4.13. Chaos Operator Based Approach

Juan, Zixing, and Jianqin [26] presented a novel GA (Figure 9 shows the flow chart) which contains chaos operator within the framework of Markov chain to prevent premature convergence.

![Fig. 7. Genetic Algorithm flowchart with a Random Offspring generation process to prevent premature convergence](image)

![Fig. 8: Key factors play an important role in GAs performance improvement](image)
Based on the analysis of the previous research, it was suggested that two key factors (convergence speed and premature convergence) play a crucial role in improving the performance of GAs as shown in Figure 8. In addition, it was explained that one can improve GA by the modification of population size; mutation probability and fitness function reduce the probability of losing gene and maintain population diversity. But increasing the population size, increases the workload as well. Therefore, a dynamic approach for increasing the population size was applied [26]. A dynamic increase in the population size means that applying selections, crossover and mutation up to a certain range of the population and if premature convergence occurs, then increase the population size and then perform the operations again on the modified population size.

It was observed that increasing the population size by generating random individuals affects the performance of the GA searching. Hence, chaotic search was implemented (see Figure 9) to enlarge the population size dynamically for the GA in which equation (15) was used as the chaotic operator.

$$x_{n+1} = ax_n(1-x_n) \tag{15}$$

The iterator in equation (15) may be called as Logistic equation. If the value of $a=4$ then the system is totally in chaotic state where $x$ might reach to any state in $(0, 1)$. It was found that equation (15) is highly sensitive to the initial state. Minute differences say $\delta$ may cause divergence after a period of time.

Juan, Zixing, and Jianqin [26] suggested ways to manipulate the chaotic operator which is drawn below:

- Judge the concentrating degree of individuals. For example, compute the difference between the average fitness value and the maximal fitness value as suggested by Srinivas and Patnaik [19] (section 4.8).
- If the fitness of individual reaches to a certain degree, then use the equation (15) to produce a new population together with the current optimal individual as the initial state $x_0$.
- The genetic operations on the new population to reach to the optimum (threshold) or maximum generation was applied. During this procedure, concentrating degree of the every generation was checked.

Caponetto, Riccardo et. al. [113] showed the effect of chaotic sequence for convergence analysis of evolutionary algorithms. The idea presented was based on the substitution of the random number generator with chaotic sequence [113]. Coelho [114] presented a novel Quantum-behaved PSO (QPSO) using chaotic mutation operator. QPSO replaced random sequences and used the application of chaotic sequences based on chaotic Zaslavskii map [114]. The simulation results showed that QPSO was a powerful tool in diversifying the population, which leads in preventing premature convergence to local optima.

Juan, Zixing, and Jianqin [26] implemented the presented approach for multimodal function optimization. It was found that the suggested approach performed better for multi-modal function optimization compared with mono-modal function optimization. The results found to be effective as far as convergence speed is concerned to reach into the optimal state and preventing the premature convergence.

4.13.1 Strengths and Weaknesses

This approach works effectively for avoiding premature convergence using chaotic operator. The significance of chaotic operator is given in various literatures [113] [114] [115] [116] [117]. As chaos is best suited operator for the highly unstable motion of deterministic system in finite phase space and it disallow the two identical populations even if the initial states obtained by algorithm are very close. These features preserve the population diversity and help GA search process to reach to the global optima. Since, chaos operator ensures the diversity and when combines with crossover and mutation operators (Figure 9) improve the convergence speed.

Although, it was shown that the present approach works on realizing the relationship between premature convergence and parameters of GA. Chaotic searching increases the population size dynamically, which introduces the population diversity and prevent premature convergence. But increase in population size brought a tremendous amount of calculation and increases the computation cost. A serious investigation is required to resolve these issues.

Affenzeller & Wagner [25] presented a new generic evolutionary algorithm to prevent premature convergence by introducing a new selection scheme. The new selection mechanism maintains the genetic diversity by applying the features of self-adaptive steering of selection pressure. In addition, this selection model not only avoids the premature convergence, but also it enables the intuitive condition to detect premature convergence. The presented selection technique was combined with Segregative Genetic Algorithm (SEGA) [55] developed by Michael Affenzeller (2001). SEGA is an advanced GA which introduces parallelism to improve global solution quality [25].

There is no manageable model exist for handling the selection pressure within the theory of genetic algorithms [56]. Hence, Self-Adaptive Selection Model was introduced in which the concept of virtual population was given to handle the selection pressure (see Figure 10). The aim of applying this model was to maintain the degree of population diversity. But, to apply this selection model it is necessary to adjust parameters for the actual selection pressure which requires lots of parameter tuning to steer the search process.

The working of selection model can be divided into the following steps:

Algorithm: Self-Adaptive Selection Model

POP: Population, P1 and P2: Parents, FITT_CHLD: Fitness of child population, FITT_PRNT: Fitness of parent population

S1 POP ← population at 1st generation

S2 Choose parents P1 and P2 for crossover using appropriate selection mechanisms such as a roulette wheel, linear rank or some kind of tournament selection etc.

S3 Before starting GA search do

If FITT_CHLD is better than FITT_PRNT then using Rechenberg’s 1/5 success rule for evolution strategies.

Filled up the claimed ratio (SuccRatio) at (i + 1)th generation with successful individuals i.e. simply fill \([\text{POP} \times \text{SuccRatio}]\).

Having filled up the claimed ratio (SuccRatio) of the next generation with successful individuals, simply fill up the rest on the next generation \((\text{POP} \times (1 - \text{SuccRatio}))\) with individual arbitrarily chosen from the pool of individuals that were also created by crossover but did not reach the success criterion.

S4 Update the size of the population as:

\[ \text{POP} = \text{POP} \times \text{SuccRatio} + \text{POP} \times (1 - \text{SuccRatio}) \]

Step3 (S3) was given to check the adoption of Rechenberg’s success rule for GA, which says that a child is successful if the fitness of the child (FITR CHLD) is ‘better’ than the fitness of its parents (FITR PRNT). The meaning of better was considered as “if it surpasses the fitness of the weaker, the better, or it’s some kind of mean value of both?” [25]. Therefore, the concept of simulated annealing (SA) was applied for making appropriate decision. Using this model one can predict the premature convergence as follows: during running GA if one cannot find \((\text{SuccRatio} \times \text{POP})\) then in this situation children outperform their own parent which causes the situation of premature convergence. In addition, \((\text{MaxSelPress} \times \text{POP})\) also shows premature convergence. MaxSelPress represent the upper limit of selection pressure.

The algorithm implemented using the above selection mechanism is known as Self Adaptive Segregative Genetic Algorithm with Simulated Annealing aspects (SASEGASA). SASEGASA added two new features in the basic GA.

a) A new variable was introduced for handling the selection pressure. It also maintains the diversity of the population.

b) A separation of the population. This concept actually increases the size search space. For getting the final result join the subpopulation at the end. The concept of categorizing the whole population into subpopulation is very similar to divide and conquer [57] [58].

By adding these two features the process of SASEGASA will end up with a population, including all the genetic information sufficient for locating a global optimum [25]. In order to test the effectiveness of the method, experiments were conducted on the travelling salesman problem and multimodal combinatorial optimization problems with the aim to see the increase in the number of subpopulation at the start of the evolutionary process.

Fig. 10. Flowchart for introducing virtual population in the new selection scheme into a Genetic Algorithm [25].
allows achieving scalable improvement in terms of global convergence [25].

4.14.1 Strengths and Weaknesses

The obtained results showed that SASEGA works effectively in avoiding premature convergence. Adding two new features to SEGA help in maintaining diversity and locating global optima. Virtual population was found effective in handling the selection pressure while the separation of population into subpopulation helped in reducing the toughness of the problem.

It was shown that virtual population and the separation of the population helped GA not trapped at local optima but at the same time these two features could be considered as weakness of this approach. Separation of population increases the population size which leads to a huge increase in computation time. It was suggested that parallel computer architecture will be suitable to apply this approach.

4.15. Multi-Combinative Strategy

Multi-Combinative strategy was implemented to avoid premature convergence in a real/binary like coded genetic algorithm (RBCGA) (Achiche, Balazinski, and Baron, 2004) [59] [60]. RBCGA was used in automatic generation of fuzzy knowledge bases (FKBs). This strategy uses varieties of crossover techniques which were applied to the same couple of parents. Two ways were suggested to perform multi-combinative reproduction in GA namely multi-parent recombination i.e. multi-crossover on multiple parents (MCMP) and multiple crossovers per couple (MCPC) [61] [62] [63] [64]. Both the crossover strategies were implemented to maintain the genetic information from parents within GA search.

Three principle crossover mechanisms such as multi-crossover, premises/conclusion and blended crossover α and uniform mutation were applied. For the selection the following technique was applied: Randomly generate a real value \( R \) in the interval \([0,1]\). \( R \) is then multiplied by \( S \) where \( S \) is the sum of fitness values of the individual population. Starting with the best individual of the population, the fitness values are summed till the result is higher than \( R S \). The last added individual is assumed as a potential parent.

It was suggested that exploration and exploitation can be used to influence the search process which helps in avoiding the premature convergence and reaching to the global optima. It was shown that the best ways of applying exploration and exploitation is: apply exploration at the early stage of the evolution, relaxing in the middle and exploitation at the end [59].

Multi-combinative approach was applied to 3D surfaces of the type. Three different surfaces such as Sinusoid surface \( z = \sin(x, y) \), Spherical surface \( z = x^2 + y^2 \) and Hyper-tangent surface \( z = \tanh(x(x^2 + y^2)) \) of different complexities were tested to ensure the effectiveness of this approach. The experimental results showed that MCPC improve the behavior of RBGA in preventing premature convergence while generating fuzzy knowledge bases using small population size. In addition, it was shown that MCPC resolves the problems of single crossover applications like fast convergence, lack of diversity and reaching a plateau.

4.15.1 Strengths and Weaknesses

Different combinations of multi-combinative evolution achieved better results in dealing with RBGA with automatic generation of fuzzy knowledge bases. Creating multiple offspring from the same parents introduces diversity and variation and keeps the search process alive. Also, the exploration/exploitation balance influences the results.

Although this approach showed promising results, we have noticed two issues with this approach. First, increase in the complexity when generating multiple offspring from the same pair of parent’s genes. Second, how to deal with too many look-alike individuals at the latter stage of the evolution.

4.16. GA using Self-Organizing Maps

Amor and Achim [6] presented a novel genetic algorithm known as “Genetic Algorithm using Self-Organizing Maps (GASOM)” to address the problem of premature convergence by intelligent exploration technique. They used self-organizing maps to mine the data from the evolutionary process. Self-organizing Map (SOM) was used to perform a nonlinear mapping from high dimensional input space onto a two dimensional grid.

Generally speaking, during the implementation of standard GAs only the current population is stored. But GASOM says that if we maintain the data of the previous generation, we could gain valuable insight into the way that GA works. By maintaining the historical data we can get the information about the problems encounter in the different generation and then we can come up with the important conclusions in order to guide and improve the search process.

In order to maintain previous generation information an intelligent and efficient technique is required, which can mine and store data. Here, we need to process the incoming data in such a manner by which a new chromosome should not lead to an entire recalculation of previous computed knowledge. In other words, we can say that we want to process high dimensions and independent data entities. Therefore, SOM were used to project high dimensional lattice because this projection does not have to be repeated when new data points are evaluated. Also, SOM stores data in a table and uses less memory. The usability of SOM’s to visualize different entities of evolutionary algorithms was shown in [10].

The training of SOM carried out in the following steps:

a) Generate the initial weight \( m_{i1}, m_{i2}, \ldots, m_{in} \) of prototype vector.

b) Repeat step (c) to (g) until the maximum number of iterations is reached.

c) Randomly pick the weight vectors \( m_i \) of different types.

d) Ensure that random selection of weight vector should be uniformly distributed; if not, then the learning will be biased.

e) Evaluate the distance (in Euclidean sense) of all the prototype vectors.

f) Check for the winning neuron \( b \) also known as a Best Matching Unit (BMU) with the prototype closest to \( x \) as:

\[
\|x - m_i\| = \min_{i} \|x - m_i\| \quad (16)
\]

g) Update the prototype as:

\[
m_i(t+1) = m_i(t) + \alpha(t)h_n(t)[x - m_i(t)] \quad (17)
\]

Where, \( t \) = current iteration, \( \alpha(t) \) =learning rate at \( t \) \( h_n(t) \) = neighborhood kernel centered at winning unit and represented as:
Fig. 11. A trained SOM in prototype representation (a), fitness landscape representation (b) and search history representation (c) [6]

$$h_{bi}(t) = \exp \left( -\frac{||r - b||^2}{2\sigma^2(t)} \right)$$

(18)

h) Ensure the quality of SOM:

$$E(i) = E(j) \quad \forall i, j \in U$$

Where, $E(i) =$ count function returns the number of times a neuron $i$ was activated.

Each evaluated individual was mapped to the nearest map unit available and store the outcome using the frequency tables, namely Population Distribution Table (PDT) and Search History Table (SHT). PDT was used to store the information of the activation frequencies w.r.t. the current population represented as $E_p(i)$ while SHT keep the track of activation frequencies w.r.t. all individuals computed during evolution represented as $E_h(i)$. Both tables were used to test the diversity. To test the diversity in the current population PDT table was used. Higher the diversity in the population means the higher number of different activated neurons.

On the other hand, SHT gives us an insight into the course of evolution. Visual representation techniques were used for interpretation (see Figure 11).

Figure 11(a) shows a trained SOM of $10 \times 10$ units. Training were done on 16-bits binary chromosomes, represented by a pattern of 16 pixels visualized as $4 \times 4$ block which displays the weight vector of that particular unit. Here, black pixel was represented by binary 0 while binary 1 used to represent the white pixel.

If a unit’s color is entirely white or black then the corresponding weight vector can be represented as either ‘1111111111111111’ or ‘0000000000000000’ respectively.

Figure 11 (a) also shows that there is a cluster of chromosomes onto neighboring neurons. Figure 11 (b) shows the fitness landscape for the 16bit 1s-counting problem where light color represents the good units, while the dark color represent the worse unit which proves the observation. At this stage a GA was implemented to avoid the dark region and interesting in getting the light color i.e. the higher fitness region. Figure 11 (c) shows that the representation of fitness landscape for 1s-counting problem after the implementation of GA where low fitness regions were avoided. Figure 11 (c) also shows that each unit is centered on a high fitness region in the lower right and the upper left corner of the SOM.

GASOM uses two ranks for computing the fitness of a chromosome. One gets the value of the first rank by ordering the population w.r.t. to the fitness values while the second rank results from the novelty of the individual population. Novelty of individual means the total number of similar chromosomes already encountered during the evolution. The novelty factor of current chromosome is represented as. If $b_i$ represent the BMU on the trained SOM then novelty can be obtained using the function as given in equation (19).

$$\text{novelty}(c) = 1/ E_h(b_i)$$

(19)

Individual fitness score (IFS) can be obtained by taking the sum of both the rank.

$$\text{IFS} = R1 + R2$$

(20)

Where, $R1$ and $R2$ represent the first and second ranks respectively.

Equation (20) was used to provide two chances to individual population to survive. The first possibility is based on the good performance on the problem under consideration while the second possibility is based on the exploration of new subspace. The activation frequencies of exploited regions were increases drastically during high exploitation. This results in decrease of the novelty factor. Then many individual having low novelty vanishes from the population and provide space to the individual having higher novelty. Hence, an increase or decreases in exploitation also affect an increase of decrease of exploration. In order to regain the lost diversity reseeding operation were applied by introducing individual have high novelty factor. Reseeding process was done in a two-step.

a) Create a reseeding pool, which consists of a set of buckets distributed uniformly. Each bucket represents a particular neuron on the SOM. Each bucket keeps the information about the newly creating chromosome and represents its BMU.

b) Then the neurons having lowest activation frequencies are determined. In some situation individual are sampled from respective buckets and inserted into the population. To keep the population size fixed kills the population with lower fitness.

It was shown clearly that the SOM reseeding technique maintains the high exploratory power. But there should be a proper balance between exploration and exploitation to gain the effective and efficient implementation of a genetic algorithm. Therefore, counting the number of different activated neurons in
the population distribution table was suggested to find the amount of exploration in the current state of the GA. Thereafter, a balance function was used to find out the number of neurons that should be activated in the current state of the search process.

If \( n \) is the number of fitness evaluation already done, \( n_{\text{max}} \) represent the maximum number of fitness evaluations and \( \|U\| \) is the total number of neurons of the SOM. Then the optimal number of activation units \( r \) can be calculated as:

\[
r = \left( 1 - \frac{n}{n_{\text{max}}} \right) \|U\| \tag{21}
\]

To achieve an effective exploration strategy there should be a good combination of the new fitness assignment scheme and the SOM reseeding operator, because if we will apply only reseeding operator then the new individuals would in most cases die immediately and disappear from the population. The reason is we didn't take novelty of an individual into account. Also, if we generate the novel chromosomes then there is no guarantee that it will rely on reproduction operators and only by chance produce new genetic material.

GASOM was implemented on Royal Road function [11], Deceptive F9 function [12] and the generalized Rosenbrock's function [2] and the obtained results shows that the performance of GASOM is far better than other approaches. In addition, experimental results were compared with other approaches such as standard steady-state GA, deterministic crowding [12] and the novel hybrid replacement scheme of Lozano et al. [13]. The results of GASOM were compared against Steady state, Crowding and Hybrid genetic algorithm using average fitness obtained for the test functions. It was observed that GASOM yields the best results out of the tested GA for Deceptive F9 function and the generalized Rosenbrock's function, whereas Hybrid GA performed slightly better than GASOM on the Royal Road function. However, the performance of hybrid method deteriorates on the other two functions.

4.16.1 Strengths and Weaknesses

The comparative analysis explains that GASOM is well suited for preventing premature convergence. It performed well on a deceptive problem where other GAs gets trapped at local optima. Also, it maintains good balance between exploration and exploitation. Visualization of GASOM is easier because it offers the use of SOM for the state of evolution.

The fundamental issues one can come across during the implementation of GASOM are maintaining the size of SOM and the effect of variable length neurons on the performance of GASOM.


In the previous literatures several approaches were tried to maintain the genotypic diversity of the population by modifying different parameters during the implementation of GA. As discussed, replacement strategy plays an important role in the population diversity management [12] [13] [38]. Goldberg and Richardson [16] and Palmer and Smith [15] implemented sharing function and scheduled sharing respectively, which modify the fitness of individuals based on their genotypes similarity. Tanese et al. [67] modified population structure in which individuals have a location and are restricted to interact to interact with their neighbors. The drawbacks with these approaches are that they work on genotypic comparison of an individual; it works effectively with bit strings, then the genetic programs and therefore limited in discovering the solution within the basins of attraction of the initial population. Breaking out the basin of attraction can be achieved by introducing new randomly generated individual in the population. Hence, a chance of a premature convergence problem is fairly common when running GA multiple times with different random number seeds, whereas restarts the search process at least increase the chances of finding the global optimum when the random individual generator can produce every point the fitness landscape, then eventually the global optimal will be found in one of the initial populations, but deciding how long to run the EA before restarting becomes a challenge [41]. If few generations are used then what is the guarantee that the population will get enough time to reach to the global optimum and large number of generations will increase processing time to converge on top of a fitness peak before starting the next generation.

In order to address the discussed issues, Jian et al. [130] presented Hierarchical Fair Competition (HFC) model. HFC was based on fitness layer. The idea of HFC was to separate the population into different layers and competition, breeding, selection, and replacement was allowed only within the fitness layer very similar to Fitness Uniform Selection (FUS) EA presented by Hutter [131]. But the drawback with this approach was individual, converged to a local optimum near the top of a fitness layer prevent newer individuals in different basins of attraction from climbing through that fitness layer.

To address the insufficiencies of HFC, an adaptive version of HFC was presented known as Adaptive Hierarchical Fair Competition (AHCF) [132]. AHCF extends HFC by allowing the admission threshold (determined either initially or adaptively) of fitness levels, which was determined dynamically through evolutionary process. Admission buffer was used to collect qualified candidates synchronously or asynchronously from other subpopulation. In addition, each subpopulation hold export threshold, defined by the admission threshold of the next higher level subpopulation. Another, modified version of HFC was implemented known as Continuous Hierarchical Fair Competition (CHFC) [133]. To improve the computational capability of AHCF, Oliveira, A., et al. [134] presented Parallel Adaptive Hierarchical Fair Competition (PAHCF) in which some enhancement were suggested in the original HFC such as an adaptive determination of number of levels, more sophisticated admission threshold adaptation mechanism and multi-processor parallel implementation. Although, PAHCF showed significantly better result than HFC because of parallel implementation, but other two approaches AHCF and CHCF have not been shown to be significantly better than regular HFC [41].

The Age-Layered Population Structure (ALPS) was introduced by Hornby (2006) [41]. Unlike, HFC and any other evolutionary algorithm, ALPS is based on age measure. The key features of ALPS are [135]: it introduces parallel execution among the multiple instances of search algorithm, each age layer assigned a maximum age and not allowed to contain individuals older than that maximum age, age of an individual is based on when the original genetic material was created from random, at regular intervals the search algorithm in the first age layer restarted.

Figure 12 explains the working of ALPS. It is shown that separation of population has been done according to the individual’s age. In addition competition, breeding, selection and replacement is restricted to adjacent layers and allowed only with same aged individuals.
Fig. 12. Age layered population structure demonstrating the layered structure according to the individual’s age. Showing the competition and breeding restriction between individuals are based on the age and it is allowed with same aged individuals.

To produce an offspring and individual created through mutation or recombination start with an age of 1 plus the age of its old parents. Each layer has a maximum allowable age for an individual to be in it. This approach checks that how long the genetic materials are participating in the population of the next generation. Age Measure can be defined as the total number of generations where an individual genetic material has been evolving in the population. The age of an individual can be computed depending on the situations using equation (22), (23), (24) and (25).

**Age Measure**

\[
\text{Age of new individual} = \begin{cases} 
0 & \text{(22)} \\
1 + \text{Age of old parent} & \text{(23)} \\
1 + \text{Age of previous generation} & \text{(24)} \\
\text{Age of previous generation} & \text{(25)} 
\end{cases}
\]

Equation (22) is applicable when a new individual was generated randomly, but if we already have the randomly generated population and we have applied the evolutionary operations such as mutation and crossover then equation (23) will be applied.

When one has copied an individual for the next generation where the copied offspring is used as a parent, then we use equation (24) otherwise equation (25). The generalization equation for computing the age of an individual is given in equation (26) [135].

\[
age = I + \frac{(\text{evals}_{\text{current}} - \text{evals}_{\text{created}})}{\text{popsize}} \tag{26}
\]

The concept of age-layer was introduced to avoid competition and breeding among individuals. The last age layer can have individuals of any age while a maximum age limit was assigned to the other age layer in the population. In order to set the age-limits, different schemes were suggested such as Linear, Fibonacci, Polynomial, and Exponential. AGE-GAP parameter was used to keep the population size as small as possible.

To ensure the effectiveness, experiments were conducted and the outcome of the experiment was compared against a handful of other EAs such as Canonical EAs, Hierarchical Fair Competition Model and Multi-start EA. In all the cases the performance of ALPS-EA was found to be significantly better than others. In addition, hybridizing ALPs was introduced as a future variant to maintain the diversity of the population.

4.17 Strengths and Weaknesses

The comparative results of ALPS showed that it outperformed other EA approach. It was observed that ALPS were capable to reduce the premature convergence by continuously generating new sub-population of randomly generated individuals in its bottom layer. It uses individual’s age in forming the layered structure of population and restricts competition and breeding, helps in diversity management and give opportunity to new individuals to develop without being dominated by older ones.

The previous work on evolving antenna design optimization for other problems an EA using deterministic crowding had been found best, whereas ALPS showed better than had been produced previously [41]. Although, the author tried to improve the performance of ALPS, but still special treatment is needed to improve this strategy.

4.18. Number Structuring Approach

Sultan et al. [14] presented odd and even number structuring approach to generate the population to perform GA search. The aim of applying the number structuring scheme was to increase the diversity and preventing the premature convergence. The odd and even number structuring follows the classic binary representation, i.e. 0 and 1. This approach was implemented to solve the timetabling problem. Structuring of the chromosome was done using the sequence of odd and even numbers as shown in the Figure 13.

**Fig. 13.** Chromosome structure ‘O’ and ‘E’ indicates odd and Even respectively.

For the effective implementation of this approach the chromosome should always be in the position of holding odd and even number in each iteration. Therefore, some heuristic was applied to ensure the structure of chromosmes. In other words, a simple adjustment technique (see Figure 14) was applied to ensure the structure of chromosomes. The OEGA (Odd-Even Genetic Algorithm) which was used for the timetabling problem is shown below (see Figure 14).
4.18.1 Strengths and Weaknesses

It was observed that this approach is comparatively simpler in nature than any other approaches for preventing the premature convergence and to maintain the diversity of the population.

We have observed two issues with this approach. First, maintaining the sequence of odd and even number structure during each generation needs extra processing time. Second, it is not suitable for handling the very complex problems such as multi-modal function optimization or any others.

4.19. Hybrid Particle Swarm Optimization (PSO) and GA

Eberhart and Kennedy [123] presented Particle Swarm Optimization (PSO) and are widely applied to optimization problems due to its high search power in state spaces. Van den Bergh and Engelbrecht [40] showed that PSO is a stochastic optimization technique based on a flock of birds or the sociological behavior of a group of people. PSO is a population based method in which the population is referred as a swarm, consists of a number of individuals known as particles. The gbest was considered as a first PSO model in which each particle i in the swarm keeps the following information such as: current position \((X_i)\), current velocity \((V_i)\), best position (associated with best fitness) of the particle \((pbest_i)\) and global best position \((gbest)\). Although PSO was shown effective for solving many optimization problems such as artificial neural network training [124] [125] [126] and function minimization [127] [128]. But few limitations were observed during the implementation of PSO:

- The swarm may prematurely converge [40].
- PSO has stochastic approach and performance is largely dependent on the problem.

Hence, to remove the insufficiency of PSO, PSO using GA was suggested which takes the merits of PSO and GA respectively. PSO and GA both are used to deal with optimization problems using evolutionary strategy. In [39] modifications were made in PSO using GA to improve the performance and reach to global maxima. The advantages of PSO and GAs are shown in the Table 2.

Looking at Table 2, one can easily understand that genetic operators can be used to prevent premature convergence. Using the crossover operation information can be exchanged between two particles which improve the likelihood of searching for the global optimum. Similarly, by applying mutation to PSO population diversity can be managed. Three different hybrid approaches were proposed, namely PSO-GA (Type 1), PSO-GA (Type 2) and PSO-GA (Type 3). In PSO-GA (Type 1) the gbest particle position does not change its position over any designated time steps whereas in PSO-GA (Type 2) the stagnated pbest particles are change their position by the mutation operator of GA. But in case of PSO-GA (Type 3) the initial population of PSO is assigned by solution of GA.

Experiments were conducted on four different benchmark functions: Shaffer, Rosenbrock, Rastrigin, and Ackley. The outcomes of the experiments were found encouraging for all the three approaches, but PSO-GA (Type 1) performed better than the other two approaches and Standard PSO for Rastrigin function.

4.19.1 Strengths and Weaknesses

The experimental results showed that PSO with GA outperform the standard PSO. Hybrid PSO was found effective to find a better solution without trapping in local maximum and achieve fast convergence. Unlike, standard PSO, PSO-GA was found more flexible and robust and produce better quality result in reasonable computational time.

The drawback with this approach is that it sometimes diversifies the particle position without reaching to global optimum or produces a worse solution.

4.20. Selective Mutation Based Approach

Cavicchio [129] presented adaptive search techniques. The purpose was to show the efficiency of adaptive system. The effect of various parameters was tested, including genetic operator probabilities and types and it was accepted that genetic operators probability plays a vital role during the search process. Srinivas and Patnaik [19] suggested adaptive probability based approach for the selection of crossover and mutation probability. In addition, the power of crossover and mutation operations was shown clearly to alleviate the premature convergence. It was observed that most of the individuals get trapped at a local optimum because the exploration effect of mutation was low.

In order to address this issue Jung [35] presented Selective Mutation for GA. It was suggested that if the exploratory power of mutation probabilities is high, then the speed of GA to reach to
global optimum become slow. Hence, one has to smartly choose the mutation probabilities to address this problem.

Szeto and Zhang [120] [121] applied matrix formulation based approach and presented adaptive genetic algorithm using mutation matrix (MOGA) [120] [121]. It was shown that mutation matrix can be used in the development of adaptive GA and therefore no external input parameter require during the selection process. MOGA uses mutation probability as a function of time, fitness ranking of the chromosome and locus. The dependence is automatically determined by the statistics of the locus and the fitness ranking of the chromosome at the given time [118]. Looking at the numerical examples on Knapsack problem shown in [118] one can observe the superiority of MOGA over many traditional methods [120] [121] [122].

Spears [119] presented the importance of crossover and mutation. Nga and Szeto [118] extended the work done by Szeto and Zhang [120] by including crossover matrix. Two crossover methods were suggested, namely Long Hamming Distance Crossover (LHDC) for exploration of the solution space and Short Hamming Distance Crossover (SHDC) for the exploitation of the local search process [118]. To ensure the effectiveness of adaptive GA with crossover and mutation matrix, experiments were conducted on one-dimensional Ising model for spin glass and the result suggested the superiority of adaptive GA with crossover and mutation matrix comparing to MOGA.

In selective mutation each individual was assigned a rank as suggested in [118] [120] [121] [122]. Thereafter, mutated one bit in a part of the individual string which was selected to correspond to their ranks [35]. Few assumptions were made such as low rank individuals are far from the global optimum whereas the high rank individuals are very close to the global optimum. The effect of this assumption was that lower rank individual needs additional mutation to explore the most significant part of the individual’s string while high rank individual mutate the least significant part of the individual’s string.

To test the effectiveness of GA with selective mutation experiments were conducted on four function optimization problems, namely, a Simple Function, a Mexican Hat Function, Shafer Function 2 and De Jong Function 2. [36] [37] [38]. The obtained result shows that GA with selective mutation performs considerably better than standard GA. In addition, one can easily apply this approach in other optimization problems.

4.20.1 Strengths and Weaknesses

Experimental results prove that selective mutation is a powerful technique to alleviate premature convergence. The additional mutation in selective mutation approach (based on rank) enhanced the performance of the GA. In addition, a rule (if the rank of individuals is high, then a bit in the least significant part is changed, and vice-versa) was provided to apply mutation.

It was observed that this technique showed fast convergence when the population size was large and vice-versa. Also, additional processing time require in comparing the rank of an individual and applying mutation twice.

4.21. Elite Mating Pool (EMP) Approach

Elite Mating Pool (EMP) was suggested to prevent premature convergence in Context Free Grammar (CFG) induction problem [49]. In this approach, simple genetic algorithm (SGA) was executed multiple times on different population to choose the best solution. The EMP approach works in the following manner [49]:

- S1: Create a mating pool of ‘n’ individuals (n, very small than population size N)
- S2: Calculate the fitness of all individuals in the population.
- S3: Sort the mating pool on the fitness of the individuals.
- S4: Perform crossover of the first parent with the elite member of the mating pool and create two offspring.
- S5: Replace old weaker individuals in the mating pool with newly created offspring in sorted order.
- S6: Perform mutation on the newly created offspring.
- S7: Replace old weaker individuals in the mating pool with newly created offspring in sorted order.
- S8: Once repeat steps S4 through S7.

In this approach diversity was maintained by generating elite members capable to satisfy multiple objectives. Here, an entire generation of the chromosome was generated by the application of the elite mating pool until either the population with a new elite chromosome was evolved or a minimum number of generations were completed with converged population.

The collections of Context Free Language (CFL) as well as Regular Languages (RL) were used for conducting the experiment [49]. To test the effectiveness of EMP approach, GA generations were processed with various probabilities ranging from 0% to 50%. It was observed that EMP approach can be used to break the local optimum convergence situation of the GA.

4.21.1 Strengths and Weaknesses

EMP approach was found effective in maintaining the population diversity. Hence, reduces the risk of premature convergence, if the population converges before reaching to the threshold value. It was observed that this approach is capable of breaking the local optimum convergence.

The problem with this approach was its execution time because extra processing time requires in maintaining the separate mating pool to get the elite members. Choubey and Kharat [49] accepted that the execution time is assumed to be greater than a normal SGA approach.

In addition, it was shown experimentally that the execution time of EMP approach was higher as compared to Dynamic Application Reproduction Operator (DARO) [49].

4.22. Dynamic Application of Reproduction Operator (DARO)

Choubey and Kharat [49] suggested Dynamic Application of Reproduction Operator (DARO) for handling premature convergence in the CFG induction problem. DARO was an extension of the work done by Nicoara [31]. Equal probability was assigned to each reproduction operator combination (crossover-mutation operator combination, CMOC) to begin the genetic algorithm generations. The flowchart for DARO is shown in Figure 15, which shows the step by step flow of this approach.

Three different crossover strategies, namely two Point Crossover (TPC), Two-Point Crossover with Internal Swapping (TPCIS) and Uniform Crossover (UC) and four mutation techniques such as: Stochastic Mutation, Inverse Mutation, Block Copier with Fixed Length Mutation and Block Copier with Random Length Mutation were implemented in DARO.

The working of this approach was given as: create the initial random population and compute their fitness value. Then assign the initial probability for each CMOC using the heuristic given in equation (27).
Random population generator & Compute Fitness
Assign initial population to each CMOC using equation (27)
Arrange current population in ascending order of the fitness value
Select the first crossover and select the first mutation operator and set \( u = 1 \)
Set \( RE = 0 \)
Select parents \( P1 \) & \( P2 \) from current population and produce offspring \( C1 \) & \( C2 \) with crossover operator \( X_i \)
Perform mutation on the newly generated offspring with the mutation operator \( M_i \)
Calculate the fitness offspring say \( \text{fitness}(C1) \) & \( \text{fitness}(C2) \)
Updated \( RE \) using equation (28)
Insert the newly generated offspring into child population
Set \( P(u) = P(u) + RE \)
Select next mutation operator & increment \( u \)
all the mutation operators are applied with current crossover operator
all crossover operators applied
apply equation (30)
create new current population by overlapping the current population with child population

**Fig. 15.** Flowchart of Dynamic Application of Reproduction Operator.

\[
P(u) = \frac{1}{m \times n} \text{ and } \sum_{u=1}^{m \times n} P(u) = 1
\]

(27)

Where, \( m \) and \( n \) represents \((m \text{ crossover operators} \times n \text{ mutation operators})\)

Now, select first CMOC and set \( u = 1 \). A term Resultant Effectivity \((RE)\) was used to ensure the effectiveness of the new offspring. Initially, \( RE \leftarrow 0 \). Choose two parents say \( P1 \) and \( P2 \) and apply the first crossover operation \((X_i)\). This will produce two offspring say \( C1 \) and \( C2 \). Perform first mutation \((M_i)\) on \( C1 \) and \( C2 \), compute \( \text{fitness}(C1) \) and \( \text{fitness}(C2) \) and then update \( RE \) using equation (28).

\[
RE \leftarrow RE + \left( \frac{[\text{fitness}(P1) + \text{fitness}(P2)]}{2} \right) - \left( \frac{[\text{fitness}(C1) + \text{fitness}(C2)]}{2} \right)
\]

(28)

Insert the new created offspring in child population. The above procedure will be repeated until \( P(u) \) proportion of the population is generated using current CMOC. Then update the value of \( P(u) \) using an equation (29).

\[
P(u) = P(u) + RE
\]

(29)

\[
P(u) = \frac{P(u)}{\sum_{u=1}^{m \times n} P(u)} \text{ and } \sum_{u=1}^{m \times n} P(u) = 1
\]

(30)

Finally, create the new current population by overlapping the current population with child population using equation (31)

\[
\text{POP}_{\text{new}} = \text{POP}_{\text{current}} + \text{POP}_{\text{child}}
\]

(31)

\( \text{POP}_{\text{new}} \) preserve the diversity and therefore prevent the premature convergence. Same set of languages (languages used for EMP [49]) was considered for the experiment purpose. It was found that this method works effectively for simple languages, but there is scope for applying this method for more complex grammar sets. It was also observed that DARO consume comparatively less time than EMP.

**4.22.1 Strengths and Weaknesses**

DARO was found effective in preventing premature by empowering the GA with dynamic application of crossover and mutation operators. It was observed that this approach consumes comparatively less processing time than EMPA (section 4.21) and therefore it can be used to overcome the issues found in EMPA.

As discussed, DARO is a time effective algorithm as compare to EMPA but it is not fit for handling the situation of local optimum convergence whereas EMPA handles such situation effectively. Also, there were no fixed rules provided to decide the initial probability to reproduction operator combination while it was already suggested by Srinivas and Patnaik [19] and Jung [35].
4.23. Hybrid Strategy using EMPA and DARO

The merits of EMPA and DARO were collected to develop the Hybrid model to avoid premature convergence and maintain the diversity of population [44]. In the hybrid model, the DARO approach leads towards generation of the optimal solution. To prevent premature convergence the appropriate mix of EMPA and random breed was used to create next population [44]. The flowchart for hybrid model is shown in the Figure 16 [44].

Figure 16 demonstrates the working of hybrid model and show the effective utilization of EMPA and DARO approaches. The hybrid model uses two important elements, namely FIT_DIFF and GEN_STEP. Both the terms are equally important and helpful in making decisions (implement DARO or not). FIT_DIFF represents the fitness difference between the best and worst fitness while the GEN_STEP shows the number of generations to be processed. The second decision box shown in the Figure 16 clearly shows that until the value of FIT_DIFF is equal or more than MIN_DIFF the process of DARO will continue otherwise it predicts the situation of premature convergence. Therefore, to prevent premature convergence, hybrid model makes decision again depending on the value of GEN_STEP. That is, it helps in making the decision to apply EMPA or apply crossover operations to generate population in the next generation (Figure 16 the last decision box indicates the same) [44].

Orthogonal Array Method [50] was used for parameter selection. It was developed by Dr. Genichi Taguchi for designing experiments to investigate the effect of parameters on the mean and variance of process performance characteristic [44]. Experiments were conducted on the different language set used by Huisjen [51], Keller and Lutz [52], Tomita [53], and Dupont [54]. In addition, other languages were also implemented, namely an odd binary number over (0 + 1) *, even binary number over (0 + 1) * and {0+1}n over (0 + 1) *. The results obtained using GA with a hybrid strategy for first ten successful runs on each language set and it was observed that hybrid approach prevent premature convergence effectively. It was also seen that both (EMPA and DARO) approaches are significantly more efficient than other approach in order to maintain the performance while the other methods are found to handle the premature convergence more effectively.

4.23.1 Strengths and Weaknesses

The convergence analysis of hybrid approach with other approaches such as Simple Genetic Algorithm (SGA), Elite Mating Pool Approach (EMPA) [49], Dynamic Allocation of Reproduction Operator (DARO) [49], Incest Prevention Algorithm (IPA) [32], Pygmy Algorithm (PA) [43], Population Replacement Algorithm (PRA) [38], Random offspring Generation Algorithm (ROGA) [22] and Social Disaster Algorithm (SDA) [42] were shown and it was found that the hybrid model works more effective than other methods. Orthogonal array method was used for the selection of parameters which reduces the issues of selection of parameters as discussed with DARO. In addition, it was also observed that DARO and EMPA showed better convergence as compared to the hybrid method at the cost of convergence at local optima but the hybrid method produces the optimal result of delaying the convergence.

Although using hybrid approach the trade-off between EMPA and DARO for execution time was achieved up to a certain level, but not completely since the weakness of this approach remains the same when applying EMPA because the hybrid approach uses the features of EMPA.

4.24. Frequency Crossover with Nine Different Mutations

Ramadan [48] implemented a GA to get the solution of the travelling salesman problem (TSP) in which the concept of Frequency Crossover (FC) along with nine different mutation techniques were used to prevent premature convergence. The idea of FC is: at a time, Y different chromosomes were selected from the existing population. All the selected chromosomes were stored in an ascending order so that Y / 2 chromosomes can form a crossover group for crossover with a 100% crossover rate [48]. The working of FC can be defined as:

Algorithm: Evaluate Frequency
- **Best Chromosome**: Choose the best chromosome in crossover group.
- **Produce offspring**: apply crossover with every other chromosome.
• Evaluate frequency: The frequency values of the corresponding genes from both parents can be evaluated as:
  If genes carry the same city
    Frequency ← two
  Else
    Frequency ← one
In order to produce the new offspring first compute the frequency, i.e., look for the genes have frequency of two and copy the value into the offspring while preserving their relative position. Then remaining genes were assigned randomly in the offspring.

Figure 17 demonstrates the working of frequency crossover strategy. Since, the genes such as X10, X3, X5, X6 and X8 are at the same location in both the parents P1 and P2, therefore the frequency count (F) for the gene is 2.

<table>
<thead>
<tr>
<th>P1</th>
<th>X10</th>
<th>X1</th>
<th>X9</th>
<th>X3</th>
<th>X5</th>
<th>X4</th>
<th>X7</th>
<th>X2</th>
<th>X6</th>
<th>X8</th>
</tr>
</thead>
<tbody>
<tr>
<td>P2</td>
<td>X10</td>
<td>X2</td>
<td>X4</td>
<td>X3</td>
<td>X5</td>
<td>X9</td>
<td>X1</td>
<td>X7</td>
<td>X6</td>
<td>X8</td>
</tr>
<tr>
<td>OS</td>
<td>X10</td>
<td>X7</td>
<td>X1</td>
<td>X3</td>
<td>X5</td>
<td>X2</td>
<td>X9</td>
<td>X4</td>
<td>X6</td>
<td>X8</td>
</tr>
</tbody>
</table>

F = 2

Fig. 17. Frequency crossover strategy

The remaining genes X1, X2, X4, X7 and X9 copied randomly in the offspring because the frequency count is 1. This process preserves the information carried and accumulated over generations, but there is no guarantee that it will maintain the population diversity and hence increases the likelihood of premature convergence. Therefore, frequency crossover was considered as a stability inducer in the GA. The selected parents and the generated offspring were used to replace the existing Y-1 chromosome. The remaining (since we have Y/2 parents and Y/2-1 offspring) were immigrated [48].

At this stage, subpopulations were formed using the new chromosome Y, which was used in the mutation step. Thereafter, best chromosomes were selected from the subpopulation to apply nine different types of mutation. And then the original subpopulations were replaced by the best chromosome along with the nine offspring generated by mutation which increase the population diversity and guaranteed that the best chromosome found in that particular generation will survive for the next generation. Nine different mutation techniques were applied, which was categorized into two groups, namely Group1 and Group2. Both the group together maintain a high degree of population diversity. Group1 mutation was used for exploration because it offers a huge change in the chromosome. As Group2 mutation introduces a minor change, therefore it was used for exploitation. Three different sets of data for TSP were taken from [23] for conducting the experiment. A test of the hypothesis was done to test the effectiveness of the GA [48] and the results showed that the GA with FC and nine different mutations performed significantly better the target value. Author of [48] not only showed the effectiveness of the GA but also the effectiveness of nine different mutations were shown. In addition, it was shown clearly that how many times FC was able to enhance the existing fitness function from one generation to the next one.

4.24.1 Strengths and Weaknesses

Overall, it was shown that the frequency crossover with nine different mutations able enough to introduce the diversity in the population and reduce the effect of premature convergence. In addition, it was shown that this approach not only increases the diversity but also guarantee that the best chromosome found in any particular generation will survive for the next generation.

But the demerit of this approach is that it is applicable only for Order-Based Chromosome representation. Also, extra processing time requires in computing the frequency and then applying nine mutation strategies.

5. Summary and Analyses of Reviewed Literatures

In this section we have furnished the summary of reviewed literatures and presented in tabulated form for quick review. Table 3 shows the details about the techniques applied in each approach. Table 4 and Table 5 shows the comparative analysis based on various parameters. Using these tables one can easily and quickly get the answer of questions such as: how diversity management was applied, how the recombination techniques was applied, how to measure the similarity of an individual, what are the key terms used with any particular approach, which selection technique was applied and above all the key concerns related with the approaches. The objective of showing the comparative analysis is to develop in depth understanding of each approach which might be helpful in designing new approaches to address the issue of premature convergence.

6. Conclusions

In this paper, we have reported the different factors which can affect the working of genetic algorithms. At the early stage of the paper we have discussed the effect of population diversity and selective pressure. In addition, we have given theoretical framework for GA which covers a brief description of three important theories, namely Schema theory, Markov chain theory and Statistical theory and their applicability have been shown by providing important literatures.

The main contribution of this paper is that it presents a review of various approaches used to prevent premature convergence in genetic algorithms. Many representative works were selected from the literature for the review and analysis. The working of each approach has been reported in a comprehensive manner with their merits, demerit and the experimental details. It has been observed that the diversity of the population and selective pressure are the key factor behind the premature convergence. In addition, it has seen that much of the research made use of GA reproduction operators to handle the premature convergence. It was shown in many literature that by applying the crossover and mutation in an appropriate manner one can avoid premature convergence. A portion of the reviewed research reported where researchers had used other techniques also to control the convergence and keep the search process alive. The reviewed literatures have been summarized for a quick review (Table 3).

By looking the summary Table 3 one can easily and quickly understand the workability of different approaches reviewed. The comparative analysis of reviewed approaches has been reported in Table 4 and Table 5 based on various aspects. This paper not only covers various approaches to prevent premature convergence, but also it reports the key issues with each approach. The purpose of reporting the issues are to identify new research path to address the problem of premature convergence.

As we have reported, lots of research work had already been done in the area of genetic algorithm. But still there is scope for the development of advance and efficient algorithm to prevent premature convergence within GA search.
<table>
<thead>
<tr>
<th>Citation</th>
<th>Approach</th>
<th>Author(s), Year of publication</th>
<th>Technique used</th>
</tr>
</thead>
<tbody>
<tr>
<td>[38]</td>
<td>Crowding Method</td>
<td>De Jong, Kenneth Alan, 1975</td>
<td>Eliminate the most similar individual whenever a new one enters in a subpopulation.</td>
</tr>
<tr>
<td>[15]</td>
<td>Scheduled Sharing</td>
<td>Palmer, Michael E., and Stephen J. Smith, 1992</td>
<td>Organism within a niche to compete with their neighbors. To calculate the convergence of a population, the search space can be divided into subspace or buckets.</td>
</tr>
<tr>
<td>[24]</td>
<td>Migration Model and nCUBE Based Approach</td>
<td>Balakrishnan, Karthik, 1993</td>
<td>A migration model was introduced to maintain the diversity in the genetic property of individual population to keep the GA search alive.</td>
</tr>
<tr>
<td>[18]</td>
<td>Cooperation Based Approach</td>
<td>Fonlupt, Cyril, Philippe Preux, and Denis Robilliard, 1993</td>
<td>Instead of using only one representation, apply varieties of GAs concurrently and each optimizing the same object but using its own representation.</td>
</tr>
<tr>
<td>[29]</td>
<td>Syntactic Analysis of Convergence</td>
<td>Louis, Sushil J., and Gregory JE Rawlins, 1993</td>
<td>The average Hamming distance of a population was used as a syntactic metrics to get the probabilistic bound on the time convergence of GAs. A model was presented to predict the time to convergence.</td>
</tr>
<tr>
<td>[43]</td>
<td>Pygmy Algorithm</td>
<td>Ryan, Conor, 1994</td>
<td>Uses two fitness functions and maintains two lists of parents, one for each criterion of the problem.</td>
</tr>
<tr>
<td>[19]</td>
<td>Adaptive Probability Based Approach</td>
<td>Srinivas, M. and Lalit M. Patnaik, 1994</td>
<td>The probability based approach was adopted which varies the probabilities of crossover and mutation depending on the fitness value of the solution.</td>
</tr>
<tr>
<td>[42]</td>
<td>Social Disaster Technique</td>
<td>Kureichik, Victor M., Victor V. Miagkikh, and Alexander P. Topchy, 1996</td>
<td>Works on the whole population to maintain the population diversity. The concept of super individual was suggested which takes over the charge of the population.</td>
</tr>
<tr>
<td>[69]</td>
<td>The Island Model Genetic Algorithm</td>
<td>Whitley, Darrell, Soraya Rana, and Robert B. Heckendorn, 1998</td>
<td>Each machine executes a GA and maintains its subpopulation for directing the search which helps to preserve genetic diversity, since each island can potentially follow a different search trajectory through the search space.</td>
</tr>
<tr>
<td>[22]</td>
<td>Random Offspring Generation Approach</td>
<td>Rocha, Miguel, and José Neves, 1999</td>
<td>Test similarity of genetic material, if found similar than generate the random offspring which will code a random solution for the problem otherwise apply reproduction.</td>
</tr>
<tr>
<td>[25]</td>
<td>Self-Adaptive Selection Pressure Steering Approach</td>
<td>Affenzeller, Michael, and Stefan Wagner, 2003</td>
<td>Introduced an intermediate step for a new selection scheme (virtual population) to handle the selection pressure. In addition, a separation of population concept was used to increase the search space. In this approach the concept of simulated annealing was also used.</td>
</tr>
<tr>
<td>[6]</td>
<td>GA using Self-Organizing Map</td>
<td>Amor, Heni Ben, and Achim Rettinger, 2005</td>
<td>Self-organizing Map was used to address the premature convergence. This approach works by maintaining previous generation information. Two frequency tables were used, namely Population Distribution Table and Search History Table.</td>
</tr>
<tr>
<td>[41]</td>
<td>Age-Layered Population Structure Approach</td>
<td>Hornby, Gregory S, 2006</td>
<td>Based on age measure defined as the total number of generations where an individual genetic material evolving in the population.</td>
</tr>
<tr>
<td>[14]</td>
<td>Number Structuring Approach</td>
<td>Sultan, B. M., Ramlan Mahmud, and Muhammad Nasir Sulaiman, 2007</td>
<td>Based on odd and even number representation technique. Introduced a function readjustment p(t) to maintain the sequence of odd and even in each generation of GA.</td>
</tr>
<tr>
<td>[39]</td>
<td>Hybrid Particle Swarm Optimization and GA</td>
<td>Premalatha, K., and A. M. Natarajan, 2009</td>
<td>Uses the merits of Particle Swarm Optimization and Genetic Algorithms. Three hybrid approaches were used PSO-GA (type-1), PSO-GA (type-2), and PSO-GA (type-3).</td>
</tr>
<tr>
<td>[35]</td>
<td>Selective Mutation Based Approach</td>
<td>Jung, Sung Hoon, 2009</td>
<td>In this approach each individual was assigned a rank. For low rank individuals, apply additional mutation to explore the most significant part of the individual’s string while for high rank individual, mutate the least significant part of the individual’s string.</td>
</tr>
<tr>
<td>[49]</td>
<td>Dynamic Application of Reproduction Operator Approach</td>
<td>Choubey, Nitin, and Madan Kharat, 2011</td>
<td>Extension of the work done in [31]. Equal probability was assigned to each reproduction operator (crossover-mutation operator combination, CMOC). Three crossovers and four mutation techniques were used.</td>
</tr>
<tr>
<td>[48]</td>
<td>Frequency Crossover with Nine Different Mutations</td>
<td>Ramadan, Saleem Zeyad, 2013</td>
<td>Based on the concept of Frequency Crossover (FC) along with nine different mutation techniques.</td>
</tr>
</tbody>
</table>
Table 4. Comparison based on key factors, diversity management, similarity measurement and crossover and mutation techniques implemented

<table>
<thead>
<tr>
<th>Approach</th>
<th>Key Factors</th>
<th>Diversity Management</th>
<th>Similarity Measurement</th>
<th>Crossover and mutation technique implemented</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crowding Method</td>
<td>Generation Gap (G) and Crowding Factor (CF) had been chosen 2 to 3 by De Jong.</td>
<td>Replacement of similar individual maintains population diversity.</td>
<td>Using genotypic distance.</td>
<td>Form n/2 pairs and then crosses all pairs and mutates the offspring.</td>
</tr>
<tr>
<td>Incest Prevention Algorithm</td>
<td>Individual Hamming distance</td>
<td>Avoided mating of pairs showing similarities based on the parent’s Hamming distance.</td>
<td>Short Hamming distance indicates that individuals might be related</td>
<td>Uses heterogeneous recombination. Mating is only allowed if the Hamming distance is above the chosen threshold, which decreases as evolution proceeds.</td>
</tr>
<tr>
<td>Scheduled Sharing Approach</td>
<td>Sharing schedule, Nice size, Sample sharing.</td>
<td>Punished the members which are similar to each other by being required to share their fitness, whereas the isolated individual retains all the fitness value they achieve.</td>
<td>Using simple two dimensional Cartesian distance function or using Hamming distance between genotypes.</td>
<td>By scaling the fitness in proportion to their similarity which is defined by a sharing function.</td>
</tr>
<tr>
<td>Migration Model and nCUBE Based Approach</td>
<td>Sending protocol, migration interval, Number of independent populations</td>
<td>Migration of population from one to another</td>
<td>Spatial segregation approach</td>
<td>Using uniform crossover, mutation and migration of population</td>
</tr>
<tr>
<td>Cooperation Based Approach</td>
<td>Appropriate representation techniques and remapping.</td>
<td>By applying remapping, which may increase or decrease of the number of local optima by dynamically changing the neighborhood of individuals.</td>
<td>Using genotypic distance</td>
<td>Crossover and mutation are applied in base 4. After computing fitness chromosome genotype was turned into base 3. Null fitness was assigned to non feasible chromosomes in base 3. N-point crossover and usual mutation operator (pick an individual and add its bits complement to the population). Recombination depended on the individual’s fitness.</td>
</tr>
<tr>
<td>Syntactic Analysis of Convergence</td>
<td>Average Hamming distance, time bound for convergence</td>
<td>Operators not considered which results in less Hamming distance to offspring compared to their parents Hamming distance. The minimum fitness individual was replaced with either the complement of randomly picked individual in the current population or the complement of the current best individual.</td>
<td>Average Hamming distance also known as Hamming average.</td>
<td>Disassortive mating, breading of phenotypically different parents sometime used by plant breeders to maintain diversity.</td>
</tr>
<tr>
<td>Pygmy Algorithm</td>
<td>Two lists namely Civil Servants, Pygmies.</td>
<td>Maintain two lists namely Civil Servants and Pygmies. For a new offspring one parent was drawn from each list.</td>
<td>Phenotypic distance of parents.</td>
<td>$P_c$ and $P_m$ are adapted based on the fitness values of an individual for recombination.</td>
</tr>
<tr>
<td>Adaptive Probability Based Approach</td>
<td>Crossover Probability ($P_c$) and Mutation Probability ($P_m$), new parameters $k_1$ and $k_2$ for controlling crossover and mutation</td>
<td>Generational replacement and explicitly protects the best solution from disruption.</td>
<td>Spatial distance (based on the fitness value)</td>
<td>Modified version of standard greedy crossover called as fifty-fifty approach with a variable probability of bad solution which require proper tuning and usual mutation operators.</td>
</tr>
<tr>
<td>Social Disaster Technique</td>
<td>Fifty-fifty approach, Packing operator, Judgement_Day and wrap operators</td>
<td>By applying crossover operator modification and operators (Packing, Judgement_Day and warp operator) working together on the whole population. In short, determine the suboptimum and then apply either Packing or Judgement_Day operator together with warp operator.</td>
<td>Distance matrix and evaluation function [92]</td>
<td>Allows recombination within the island and sometimes across the island</td>
</tr>
<tr>
<td>The Island Model Genetic Algorithm</td>
<td>Migration interval, migration size, topology used for migration and migration policies adopted.</td>
<td>Migration of highly fitted individuals across subpopulation increase the diversity of the local subpopulations</td>
<td>Some sort of distance in Hamming space</td>
<td>Recombination was done between colony and core groups. The colony may apply full or partial sharing of its member (selected randomly according to the fitness) with core by</td>
</tr>
<tr>
<td>Shifting Balance Theory in Dynamic Environment</td>
<td>Core group, Colonies, diversity measure, distance measure, Containment function, Migration from colony to core and migration interval</td>
<td>A parameter $Diversity(P)$ was suggested to see the diversity of the population. First populate the core group from the member of old core through reproduction. After that form colony and find out up to what extent member’s of colony lies in core group. Divide the colony in two groups called as inside and outside. Compute $Containment(A, B)$ and apply immigration of population</td>
<td>Hamming distance was used. Three measure was identified namely, diversity measurement, distance (distance between a single chromosome and a</td>
<td></td>
</tr>
</tbody>
</table>
Random Offspring Generation Approach  
Random operators, hybrid operators, 
- Blind offspring
- Random offspring

Concept of Random offspring generation was applied before the recombination. Test the similarity of individual, if individuals are not similar, simply apply crossover and mutation otherwise generate offspring to code a random solution.

Chaos Operator Based Approach  
Chaos operator, logistic equation, 
- Chaotic operator and Logistic equation

Chaotic operator ensures that there do not exist two identical populations, even if the two initial states are very close. Such population maintains the diversity and generates optimal solution.

Self-Adaptive Selection Pressure Steering Approach (SASEGASA)  
Simulated annealing, virtual population, 
- Success pressure
- Selection pressure

In this method, GA with simulated annealing was applied. During running GA if one cannot find \((\text{SuccRatio} \cdot \text{POP})\) then children outperform their own parent and \((\text{MaxSelPress} \cdot \text{POP})\) both shows premature convergence whereas \text{MaxSelPress} represent the upper limit of selection pressure. This information might be useful in preserving diversity and predicting the premature convergence.

Multi-Combinative Strategy  
Multi crossover, fuzzy rule crossover, 
- Multi-crossover, fuzzy rule crossover

Multi-parent recombination, MCMP and MCPC were applied with real/binary coded GA to maintain the diversity and keeping the search process alive.

GA using Self-Organizing Maps  
Self organizing map, Best matching unit (BML), Population distribution table (PDT), Self organizing table (SHT), Novelty of chromosome, Reseeding operator, Reseeding pool, and Activated unit.

SOM was used to visualize various aspects of search. It maintains the search history using two tables such as PDT and SHT. PDT was used to examine the diversity in the current population, while SHT gives an insight into the course of evolution. In addition, a reseeding operator and a control mechanism were applied to balance exploration and exploitation.

Age-Layered Population Structure (ALPS) Approach  
Individual’s age, Layered structure of populations and Age gap.

Age of individual’s was defined and populations were structured into multiple layers with maximum allowable age for an individual and selection, breeding and replacement is restricted to adjacent layers. Using odd-even number side by side along the chromosome. A procedure “Procedure Readjustment (P)” was used. This procedure consists of simple repair instruction to readjustments back the value inside the gene. It also ensures the sequence of odd-even remains at the respective gene after iterations.

Hybrid Particle Swarm Optimization (PSO) and GA  
PSO-GA (Type-1), PSO-GA (Type-2) and PSO-GA (Type-3).

Position update of the global best particles was changed. The position update was done through some hybrid mechanism of GA. Three different hybrid approaches were suggested: PSO-GA (Type-1), PSO-GA (Type-2) and PSO-GA (Type-3).

Selective Mutation Based Approach  
Selective mutation and individual rank.

The selective mutation technique was applied. Each individual was first ranked and then additionally mutated one bit in a part of their population in gene-space measure and containment function. Genotype or phenotype distance

Recombination was done after checking the similarity of individual using various crossover operators such as blind operators (UOPX, EDGX, MPX and SCHX) and hybrid operators (GX and HGX). Four different categories of mutation operators (adjacent swap, non-adjacent swap, sub list scramble and partial inversion) was introduced “virtual population” and “separation of population” when applied recombination operations.
strings which was selected to correspond to their rank. The selective mutation helps in diversity management and help GA to reach to global optimum and to quickly escape from local optima. Mating pool was created which maintains the diversity by generating the elite members more capable of satisfying the multiple objectives. Crossover and mutation operator was applied to elite members of mating pool and replaces the weaker individual.

Dynamic application of crossovers and mutation operators was applied. Probability value was assigned for each reproduction operator combinations. Resultant Effectivity (RE) was used which was set to zero initially and updated based on the fitness of a pair of parents and created offspring. RE was also used to update the probability of reproduction operator combination. The value of RE helps maintain the diversity of the population and selection of reproduction operator combination.

To know the premature convergence, FITT_DIFF was compared with MIN_DIFF. If the FITT_DIFF is greater than or equal to FITT_DIFF then there is no premature convergence and move to next generation otherwise it shows the situation of premature convergence. As this approach uses the merits of EMP and DARO therefore to decide which one will be applied to handle the situation of premature convergence the condition $G\%\text{GEN\_STEP} = 0$ was applied if it shows the true result then apply DARO otherwise EMP was applied to maintain the diversity.

The standard crossover operator was applied to elite members of mating pool and replaces weak individual with newly created offspring. Now, mutation operator was applied on newly created offspring and again replaces weak individual with newly generated offspring. Finally, apply selection mechanism. Three crossovers (Two point crossover, Two point genotypic or phenotypic) and four mutations (inverse mutation, block copier with fixed length, block copier with random length and stochastic operator) operators were used. Equal probability was assigned to each reproduction operator combination which gets updated in each generation. It is inspired by the approach suggest by Nicoară, Elena Simona (2009) [31].

Applied the combined mechanism of EMP [49] and DARO [49] approaches

<table>
<thead>
<tr>
<th>Approach</th>
<th>Competition</th>
<th>Selection Technique</th>
<th>Applied On</th>
<th>Key Concern</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crowding Method</td>
<td>Offspring compete against one of the parents that produce it.</td>
<td>Restricted tournament selection (RTS) adapts standard tournament selection (TS)</td>
<td>Multimodal function optimization</td>
<td>● Stochastic error by low crowding factor (CF),&lt;br&gt;● Maintaining bitwise diversity&lt;br&gt;● Replacement error is the major issue.</td>
</tr>
<tr>
<td>Incest Algorithm</td>
<td>Prevention</td>
<td>Based on Hamming distance if above a threshold. If in a generation, no pair can be chosen, then decrease the threshold.</td>
<td>Multimodal function optimization</td>
<td>● Additional computation cost to compute individual’s similarity which varies the execution time&lt;br&gt;● Difficult to maintain diversity if the member of the new population has common ancestors and consequently due to a finite fixed population size which causes a loss of genetic diversity.</td>
</tr>
<tr>
<td>Scheduled Sharing Approach</td>
<td>Organisms presents in a niche compete with their neighbors</td>
<td>Phenotypic fitness by a sharing factor (a number between zero and one)</td>
<td>Rugged two-dimensional landscape</td>
<td>● Finding the appropriate schedule for sharing&lt;br&gt;● Deciding the sharing functions&lt;br&gt;● It is one of the expensive schemes, and</td>
</tr>
<tr>
<td>Method/Approach</td>
<td>Population Interaction</td>
<td>Selection/Proportionate</td>
<td>Representation/Selection</td>
<td>Problem Solved</td>
</tr>
<tr>
<td>-------------------------------------------------------------------------------</td>
<td>-----------------------</td>
<td>-------------------------</td>
<td>--------------------------</td>
<td>-----------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Migration Model and nCUBE Based Approach</td>
<td>No correlation between neighboring individuals</td>
<td>Proportionate fitness and ranking selection</td>
<td>Using genotype's fitness an individual was selected if their fitness is better.</td>
<td>Kookie problem</td>
</tr>
<tr>
<td>Cooperation Based Approach</td>
<td>No correlation between neighboring individual</td>
<td>Fitness proportion selections</td>
<td>Flat function, DeJong’s five functions (F1 through F5), and One max problem.</td>
<td></td>
</tr>
<tr>
<td>Syntactic Analysis of Convergence</td>
<td>No correlation between neighboring individual</td>
<td>Fitness proportion selections</td>
<td>Multi-objective function optimization, problem of evolving minimal network sorters.</td>
<td></td>
</tr>
<tr>
<td>Pygmy Algorithm</td>
<td>No attempt was made to measure this.</td>
<td>Selection of parents for breeding was analogous to differing genders. One parent is selected from each list (Pygmy list and Civil Servant list)</td>
<td>Multimodal function optimization, De Jong’s functions known as spiky function (f5, f6, f7), Order-3 Deceptive problem.</td>
<td>Travelling Salesman Problem.</td>
</tr>
<tr>
<td>Adaptive Probability Based Approach</td>
<td>No correlation between neighboring individual</td>
<td>Fitness proportion selections</td>
<td>Multimodal function optimization, DeJong’s functions known as spiky function (f5, f6, f7), Order-3 Deceptive problem.</td>
<td>Travelling Salesman Problem.</td>
</tr>
<tr>
<td>Social Disaster Technique</td>
<td>Checked using a distance matrix calculation</td>
<td>Roulette wheel selection</td>
<td>Travelling Salesman Problem</td>
<td></td>
</tr>
<tr>
<td>The Island Model Genetic Algorithm</td>
<td>No correlation between neighboring individual</td>
<td>Biased rank based selection and fitness proportionate selection</td>
<td>Two linearly separable problems (fully deceptive order-4 function and Rastigin’s function) and two nonseparable problems (Powell’s Singular function and Rana’s function)</td>
<td></td>
</tr>
<tr>
<td>Shifting Balance Theory in Dynamic Environment</td>
<td>Competition among the individual based on the fitness both in the core and the colony.</td>
<td>Based on distance to the core (for inside section), and Fitness proportionate Selection (outside section)</td>
<td>An experiment was done on F8F2 function. F2 function of the De Jong test suite composed with a 1-D version of the Griewangk Function (F8)</td>
<td>Travelling Salesman Problem (instance taken from [23])</td>
</tr>
<tr>
<td>Random Offspring Generation Approach</td>
<td>No correlation between neighboring individual</td>
<td>Roulette-Wheel scheme</td>
<td>Mono-modal function optimization and Multimodal function optimization (performed better compared to mono-modal).</td>
<td>Travelling salesman problem and multimodal combinatorial problems.</td>
</tr>
<tr>
<td>Chaos Operator Based Approach</td>
<td>No correlation between neighboring individual.</td>
<td>Fitness proportionate selection</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Self-Adaptive Selection Pressure Steering Approach</td>
<td>No correlation between neighboring individual.</td>
<td>Standard selection mechanism such as a roulette wheel, linear rank or some kind of tournament selection with virtual population</td>
<td>Two linearly separable problems (fully deceptive order-4 function and Rastigin’s function) and two nonseparable problems (Powell’s Singular function and Rana’s function)</td>
<td></td>
</tr>
</tbody>
</table>

- It requires prior knowledge to write sharing function.
- Setting migration interval is an important factor.
- Taking the decision to migrate an individual to migrate, and
- Number of independent populations.
- Selection of appropriate representation techniques during the GA search.
- Remapping of chromosome dynamically, and
- Dealing with more than representation is quite difficult.
- Mutation rate affects the Hamming convergence.
- Finding the upper bound and lower to compute time to converge is difficult and time consuming and
- Qualitative prediction may not be possible with pure syntactic information.
- Not fit for multi-objective function in some situation because individuals in the same group cannot mate.
- The issue was addressed by applying Racial Preference factor for individuals.
- Selection of the Crossover and Mutation Probability
- Protecting the best solution explicitly, and
- Handling new parameters $k_1$ and $k_2$ for controlling the adaptive nature of $P_m$ and $P_C$.
- The modified greedy crossover approach was used.
- Decrease in crossover greediness, increases the variation in the population.
- Efficient usage of the population pool,
- No measure was provided for the decrement in crossover greediness’
- Tuning the probability of bad solutions results in extra computational cost.
- Maintains diversity by sharing information with different island,
- Migration interval and migration size play an important role.
- No information is available to decide the migration interval and migration size.
- Computation cost increases in dividing the colony and computing the fitness of individual population.
- This approach also suffers with the problem of migration interval and
- It was accepted that no dynamic model available for the dynamic environment.
- Computational overhead disregarded compared to Crowding [38], Sharing [16] and Scheduled Sharing [15] approaches.
- It helps in faster convergence
- Not capable to resolve the premature convergence completely and
- The tremendous amount of computation is required.
- Separation of population increases the size of the population;
- The computational time for this algorithm is huge, and
- Parallel computer architecture will be suitable to apply this approach.
Multi-Combinative Competition among the individual population based on their fitness when applying selection mechanism.

GA using Self-Organizing Maps Competition among the individual based on the self-organizing property. Based on the age of the SOM approach.

Age-Layered Population Structure (ALPS) Approach Competition among the individual population based on the novelty. Based on the age of individual’s structure.

Number Structuring Approach Competition among the individual based on the particle fitness. Based on the fitness value.

Hybrid Particle Swarm Optimization (PSO) and GA Competition among the particle based on the fitness value. Based on the particle fitness.

Selective Mutation Based Approach Competition among the individual based on the fitness value. Based on the fitness value.

Elite Mating Pool Approach Competition among the elite members of the mating pool based on their fitness. Based on the fitness value.

Dynamic Application of Reproduction Operator Approach Competition among the individual based on the fitness value. Based on the fitness value.

Hybrid Strategy using EMPA and DARO Competition among the individual based on the fitness value. Based on the fitness value.

Frequency with Nine Different Mutations Competition among the individual largely depends on their fitness and frequency. Based on the fitness value and frequency.

(SASEGASA) Based on the success ratio (see the algorithm section 4.14)

Randomly generate a real value \( R \) in the interval \([0, 1]\). Compute \( S \), where \( S \) sum of fitness values of the individual population. Evaluate \( RS \). Now, starting with the best individual of the population, the fitness values are summed till the result is higher than \( RS \). The last added individual is assumed as a potential parent. Binary tournament selection

Applied on 3D surfaces of type \( z = f(x, y) \). Three different surfaces of different complexities were used, namely Sinusoid Surface, Spherical Surface and Hyper Tangent Surface.

Royal Road function, the Deceptive F9 function from and the generalized Rosenbrock’s function. Antenna design optimization problem

Timetabling problem

Shaffer’s function, Rosenbrock’s function, Rastrigin’s function, Ackley’s function

Simple Function, a Mexican Hat Function, Shaffer Function 2 and De Jong Function 2.

Collections of Context Free Language (CFL) as well as Regular Languages (RL)

Collections of Context Free Language (CFL) as well as Regular Languages (RL)

Applied to different languages set used by Huijsen [51], Keller and Lutz [52], Tomita [53], and Dupont [54]. In addition, odd binary numbers \( \overline{0+1}^n \), even binary numbers \( (0+1)^n \) and \( 0^n \overline{2^n} \overline{(0+1)^n} \). Travelling Salesman Problem

to enhance the performance

• Increase in the complexity when creating new offspring using different techniques.

• Although, creating new offspring from the same pair of parents’ gene was found an effective way to improve the diversity, but at the latter stage of the evolution, the presence of too many look-alike individuals is still a problem.

• Maintaining the size of the SOM and

• The effect of variable length neurons on the performance of GASOM

• EA with deterministic crowding showed the best results for several while ALPS showed better results.

• Suitable only for small sized population.

• Not suitable for very complex problems.

• Sometimes diversify the particle position without reaching to global optimum or produces a worse solution.

• Showed fast convergence when the population size was large whereas slow convergence for small population size.

• Extra processing time require in applying mutation twice.

• Consume extra processing time in maintaining the separate pool to get the elite members during offspring creation.

• Slower than SGA.

• No fixed rule given to assign the initial probability to reproduction operator combination as suggested by Srinivas and Patnaik (1994) [19] and Jung, Sung Hoon (2009) [35].

• Not suitable for breaking the local optimum convergence.

• Again problem remains the same when applying EMP approach since this approach uses the features of EMP.

• Orthogonal array method was used for the selection of parameters which reduces the issues of selection of parameters as discussed with DARO.

• Extra processing time requires in computing the frequency and then applying nine mutations approaches.

• For the experimentation order based representation was used, the effect of other representation had not been discussed.
7. References


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Graphical Abstract:

- **Introduction to Genetic Algorithm**
  - Factors Affecting GA
    - Population Diversity
    - Selection Pressure
  - Theoretical Framework
    - Statistical Mechanics
    - Markov Chain Theory
    - Schema Theory
  - Approaches to Prevent Premature Convergence
    - Crowding Method
    - Incest Prevention Algorithm
    - Scheduled Sharing Approach
    - Syntactic Analysis of Convergence
    - Adaptive Probability Based Approach
    - The Island Model Genetic Algorithm
    - Shifting Balance Theory in Dynamic Environment
    - Self-Adaptive Selection Pressure Steering Approach (SASEGASA)
  - Strengths and Weaknesses of each approach
    - GA using Self-Organizing Maps
    - Age-Layered Population Structure (ALPS) Approach
    - Hybrid Particle Swarm Optimization (PSO) and GA
    - Dynamic Application of Reproduction Operator (DARO)
    - Frequency Crossover with Nine Different Mutations

- **Preventing Premature Convergence in Genetic Algorithm**
  - Summary and Analyses of Reviewed Literatures
  - Conclusions
Highlights of the Paper

- Detailed discussion on various approaches for handling premature convergence in GA.
- Theoretical framework is presented for convergence analysis of GA.
- Strengths and weaknesses of each approach are provided.
- Summary and comparison of the approaches is given for quick review.