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Complexity Analysis and Stochastic Convergence of Some Well-known Evolutionary Operators for Solving Graph Coloring Problem

Raja Marappan * and Gopalakrishnan Sethumadhavan

Department of Computer Applications, School of Computing, SASTRA Deemed University, Tirumalaisamudram 613401, India; sgk@mca.sastra.edu
* Correspondence: raja_csmath@cse.sastra.edu

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Abstract: The graph coloring problem is an NP-hard combinatorial optimization problem and can be applied to various engineering applications. The chromatic number of a graph $G$ is defined as the minimum number of colors required to color the vertex set $V(G)$ so that no two adjacent vertices are of the same color, and different approximations and evolutionary methods can find it. The present paper focused on the asymptotic analysis of some well-known and recent evolutionary operators for finding the chromatic number. The asymptotic analysis of different crossover and mutation operators helps in choosing the better evolutionary operator to minimize the problem search space and computational complexity. The choice of the right genetic operators facilitates an evolutionary algorithm to achieve faster convergence with lesser population size $N$ through an adequate distribution of promising genes. The selection of an evolutionary operator plays an essential role in reducing the bounds for minimum color obtained so far for some of the benchmark graphs. This research also focuses on the necessary and sufficient conditions for the global convergence of evolutionary algorithms. The stochastic convergence of recent evolutionary operators for solving graph coloring is newly analyzed.

Keywords: approximation methods; chromatic number; combinatorial optimization; complexity analysis; evolutionary approach; genetic algorithm; graph coloring; NP-hard; stochastic convergence

1. Introduction

A simple graph $G = (V, E)$ consists of a set of vertices $V = \{v_1, v_2, v_3 \ldots v_n\}$ and a set of edges $E = \{e_1, e_2, e_3 \ldots e_m\}$ such that each $e_i$ $(1 \leq i \leq m)$ is uniquely associated with an unordered pair of vertices $(v_j, v_k)$ $(1 \leq j, k \leq n)$ and $j \neq k$ [1,2]. Its adjacency matrix is denoted by $A(G)$, an $n \times n$ symmetric matrix where $A(j, k) = 1$ $(1 \leq j, k \leq n)$ if $(v_j, v_k) \in E(G)$; and $A(j, k) = 0$ $(1 \leq j, k \leq n)$ otherwise [2].

$\chi(G)$, the chromatic number of $G$, is defined as the minimum number of colors required for $V(G)$ so that no two adjacent vertices are of the same color [2]. The graph coloring problem (GCP) finds the value for $\chi(G)$ and applies it to register allocation, channel assignment, image segmentation, resource utilization, and scheduling [3–10]. With the increasing values of $n$, the complexity of determining $\chi(G)$ also increases. GCP is an NP-hard combinatorial optimization problem. Hence, a fast evolutionary and approximation method is expected to reduce the problem search space by maximizing the number of successful runs [1,11–16]. Tabu search [17], backtracking [18], branch and bound [11], evolutionary algorithm [19–21], branch and cut [22], particle swarm optimization (PSO) [23–26], ant colony optimization (ACO) [27], local and cuckoo search [28,29] are some existing methods for finding $\chi(G)$. Some recent applications of GCP are selective graph coloring [30,31], signed graphs coloring [32,33], scheduling, and resource allocation [7–10,34–38]. In comparison to other methods, GA is useful for solving multi-objective optimization problems with vast search space. Hence, it is
expected to design new evolutionary operators to achieve faster stochastic convergence with minimal
generations. Some recent evolutionary operators have achieved near-optimal solutions for some
benchmark graphs [4,39–41]. Moreover, some recent genetic operators have increased the number of
successful runs and reduced expected generations \( \bar{g} \), expected crossovers \( \bar{c} \), and the expected
mutations \( \bar{m} \). The notations \( \bar{g} \), \( \bar{c} \), and \( \bar{m} \) indicate respectively the average number of generations, the average
number of crossovers, and the average number of mutations performed to obtain the near-optimal
solution for the specified number of total runs and generations during the execution of the program.
This paper focused on the asymptotic analysis and stochastic convergence of some well-known and
current evolutionary operators.

Some of the recent recombination operators are similar to mutation and are executed in place of
recombination operation [2]. Some of the well-known and recent recombination operators operate on
multiple parent gene sequences [2]. The order-based uniform crossover operator [42] uniformly selects
genes from various parents. The penalty based crossover operator [43] generates offspring, which
results in a minimum penalty. The one-point and two-point crossover operators generate random
numbers and inherit genes from their parents. The graph adapted crossover generates genes based on
conflicting and neighboring vertices. The union independent set (UIS) crossover [44] operator unifies
the pairs of independent color sets from the selected parents. The greedy partition crossover (GPX) [45]
operator divides \( V(G) \) into \( k \)-sets and generates offspring. The multi-parent crossover [46] operator
is the extension of GPX, which creates offspring from multiple parents. The well-informed partition
crossover (WIPX) [47] operator selects the color classes from the randomly chosen parents based on
a scoring function. The penalty-based color partitioning crossover (PCPX) [48] operator generates
offspring based on the partition of \( V(G) \) and its penalty. The degree based crossover (DBX) [48] operator
applies the heuristics of the largest degree ordering (LDO) to order-based crossover or permutation
one point crossover (POP) [40]. POP crossover performs an order-based permutation operation. The
multi-point guided crossover (MPGX) [48] operator incorporates problem-specific knowledge. The
merging crossover (MOX) [40] operator merges two-parent gene sequences and generates order-based
offspring. The merging independent sets (MIS) [40] crossover groups the color sets of parents and
generates offspring.

A recent crossover operator, the single parent conflict gene crossover (SPCGX), similar to mutation,
is applied to the single parent to identify the conflicting genes and generate better offspring [49]. SPCGX
is combined with the conflict gene removal (CGR) procedure to identify and remove some conflicting
genes [50]. SPCGX is also connected with the advanced local guided search (ALGS) operator [2] to
fine-tune the offspring further and also to reduce the computational complexity. The single parent
conflict gene extended crossover (SPCGEX) operator is applied to the selected single parent for fixed
iterations. Extended SPCGEX (ESPCGEX) extends SPCGEX with CGR for fixed iterations to produce
better offspring. The multipoint SPCGX (MSPCGX) performs several crossovers for conflicting genes.
Some of the recent genetic operators have achieved the better solution for some large DIMACS
benchmark graphs in the reasonable expected generations [2].

The mutation operators operate on offspring generated by crossover operations. The random
mutation operator [43] randomly swaps the color of each vertex with low mutation probability \( p_m \). The
polynomial mutation operator rounds off the floating-point color values. The swap mutation is applied
to offspring generated by the DBX operator [51]. Problem-specific mutation [51] operators operate on
the offspring generated by the MPGX operator. The order mutation (OM) [52] operator generates a
random number \( r \) and performs \( r \) interchanges between vertices. The block mutation (BM) [52] performs
the translation of blocks of successive vertices. The color spread mutation (CSM) [52] randomly selects
the conflicting edges and moves the genes to new positions. The bad edge stretch mutation (BESM) [52]
reduces the number of conflicting edges. The conflict gene mutation (CGM) [2] assigns the conflict-free
integers to the offspring and performs fixed iterations. The CGM-CGR operator further reduces the
number of conflicts in the offspring. The extended CGM (ECGM) operator fine-tunes the offspring by
extending CGM for finite iterations. The multipoint CGM (MCGM) performs multipoint mutations at the conflicting vertices to reduce the number of conflicts.

Section 2 explains the need for complexity analysis and stochastic convergence for evolutionary operators. Section 3 focuses on the general structure of the genetic algorithm for solving GCP. Sections 4 and 5 explain some of the well-known and recent recombination and mutation operators. The necessary and sufficient condition for the global convergence of evolutionary algorithms and the stochastic convergence of recent genetic operators are newly analyzed in Section 6. Conclusions and future research directions are delineated in Section 7.

2. The Need for Complexity Analysis and Stochastic Convergence of Evolutionary Operators

GCP is very challenging for genetic algorithms because of its vast solution space. Hence, designing and choosing the right genetic operators in population-based methods are important for the following reasons [53–56]:

1. Solution space of GCP consists of \(n^n\) candidate solutions, and \(n^n\) rapidly grows with \(n\). The computational complexity of finding \(\chi(G)\) is proportional to \(m\) and \(n\). If the solutions are represented using \(\chi(G) < n\), then the solution space contains \(\chi(G)^n\) assignments. Therefore, the total number of different assignments in the solution space is \(\chi(G)!\)
2. The computational complexity of GCP is reduced only when the values of \(c\), \(m\), and \(g\) are minimized.
3. The evolutionary operators are expected to improve the performance of the genetic algorithm by reducing the search space as well as by increasing the successful runs.
4. The evolutionary operators should improve the near-optimal solution obtained from the existing methods [11,22,25–27,57–59].
5. The evolutionary operators should avoid additional memory resources and complex computational operations.
6. The performance of operators depends on how many parent gene sequences are considered for crossover and mutation. It also depends on the update of gene sequences [4,39–41].
7. A smaller population size (\(N \leq 15\)) is expected to reduce the complexity and to achieve a sufficient convergence rate. The crossover and mutation operations can be embedded with the right search operators to reduce the value of \(\frac{g}{N}\) further.
8. The evolutionary operators should effectively distribute the promising genes from one generation to the next generation in order to reduce the fitness function values.
9. The graph characteristics, graph instances such as \(n\), \(m\), and graph density, can be considered for choosing the right genetic operators [60].
10. The evolutionary operators should quickly converge with the reduced computational complexity.

3. The General Structure of Genetic Algorithm for Solving GCP

Notations

1. Degree, \(\delta(G)\) and \(\Delta(G)\) The number of edges incident on \(v_i\) (1 \(\leq i \leq n\)) is the degree, \(d(v_i)\) in \(G\). Let \(\Delta(G) = \max_{1 \leq i \leq n} |d(v_i)|\) if \(v_i \in V(G)\) and \(\delta(G) = \min_{1 \leq i \leq n} |d(v_i)|\) if \(v_i \in V(G)\) be the maximum and minimum degrees of \(G\) respectively.
2. Encoding of Gene Sequence The color values of \(V(G)\) are encoded as \((g_1, g_2, g_3 \ldots g_n)\), where \(g_j\) (1 \(\leq j \leq m\)) is a non-zero integer.
3. Density The density of \(G\) is \(\frac{2m}{n(n-1)}\).
4. Population Size, Crossover, Mutation Probability \(p_c\) is the crossover probability, and the mutation probability is denoted as \(p_m\). \(P_g\) is the population, which represents the collection of gene sequences for generation \(g\). \(N = |P_g|\) defines the population size for \(g\).
5. Fitness Function $f(G)$, the general fitness function of $G$, is the number of distinct genes in the gene sequence [2]. The main objective of GCP is to minimize $f(G)$, hence, $\chi(G) - f(G) = 0$.

6. Conflicting Genes When $g_a = g_b$ and $(v_a, v_b) \in E(G)$, the genes $g_a$ ($1 \leq a \leq n$) and $g_b$ ($1 \leq b \leq n$) corresponding to vertices $v_a$ and $v_b$ conflict with the gene sequence $(g_1, g_2, g_3 \ldots g_n)$.

The general structure of the genetic algorithm for solving GCP is presented below.

Step 1: Initialization
First, initialize $g = 0$. The initialization of $P_g$ randomly or heuristically generates values in $[1, \text{minimum color}]$.

Step 2: Evaluation of Fitness Function and Selection of Individuals
The fitness of individuals in the population is evaluated, and one or more parent gene sequence(s) are selected for the generation of the subsequent gene sequences. Different selection methods can be applied to select gene sequences for crossover operation.

Step 3: Recombination or Crossover Operation
The recombination operation is performed on the selected gene sequence(s) with probability $p_c$ (high value of $p_c$, for example, $p_c = 0.8$) to generate offspring.

Step 4: Performing Mutation
The crossover gene sequences or offspring are mutated with the chosen mutation probability $p_m$ (low value of $p_m$, for example, $p_m = 0.2$) to generate new offspring.

Step 5: Termination
Find $F_g$ (offspring) for the updated gene sequence. If $F_g$ (offspring) = 0, the generation of offspring stops; otherwise, $P_g$ is updated ($g = g + 1$). If $F_g$ (offspring) $< F_{g-1}$ (worst), $P_g$ is updated by replacing the worst gene with offspring. Then transfer the control to Step 2 for the next generation.

The pseudo-code of the general genetic algorithm is presented below.

// Pseudo code of the Genetic Algorithm
1: Initialization of individuals;
2: While termination conditions are not satisfied, do
3: Evaluate the fitness function and select better individuals;
4: Perform the recombination operation;
5: Perform the mutation operation;
6: Apply elitism operation;
7: Print the near-optimal results;

4. Asymptotic Analysis of Some Existing Well-known and Recent Crossover Operators

Some of the existing well-known and recent crossover or recombination operators for solving GCP are presented in this section. The asymptotic complexity of these operators is also analyzed in Table 1. The simple control constructs if and if-else statements that can be implemented in a constant complexity, $\Theta(c)$. The control constructs for, while, do-while, and repeat-until can be implemented in a linear complexity, $\Theta(n)$. The quadratic complexity is represented in $\Theta(n^2)$. For example, color assignments of $V(G)$ can be verified in $\Theta(n^2)$ complexity.

1. Order-based Uniform Crossover

The order-based uniform crossover operator uniformly selects the genes from multiple parents. The procedure can be described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.
b. Generate the offspring by selecting the first $n/2$ length of genes from $p_1$ and the next $n/2$ length of genes from $p_2$.
c. Randomly select any two vertices, $v_i$ and $v_j$ in the offspring, and scramble all vertices between $v_i$ and $v_j$. 
Analysis: Steps (a), (b) and (c) take $\Theta(n)$ complexity. Thus the order-based uniform crossover takes $\Theta(n)$ complexity.

2. Penalty Based Crossover

The penalty based crossover operator generates the offspring, which results in a minimum penalty. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.
b. Generate the offspring by determining the color for each vertex in every parent by assigning colors, which results in a minimum penalty.

Analysis: Step (a) takes $\Theta(n)$ complexity. Step (b) takes $\Theta(n^2 \chi(G))$ complexity. The computational complexity of penalty based crossover is $\Theta(n^2 \chi(G))$.

3. One-point Crossover

The one-point crossover operator generates random numbers and inherits genes from their parents. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.
b. Choose a random point $r$ in between $[1, n]$.
c. Generate the offspring based on $r$ by inheriting genes from both parents.

Analysis: Step (a) takes $\Theta(n)$ complexity. Step (b) can be implemented in constant running time. Step (c) requires $\Theta(n)$ complexity. Thus the one-point crossover takes $\Theta(n)$ complexity.

4. Two-point Crossover

The two-point crossover operator generates random numbers and inherits genes from their parents. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.
b. Choose two random points, $r$ and $s$ in between $[1, n]$.
c. Generate the offspring based on $r$ and $s$ by inheriting genes from both parents.

Analysis: Step (a) takes $\Theta(n)$ complexity. Step (b) can be implemented in constant running time. Step (c) requires $\Theta(n)$ complexity. Thus the two-point crossover takes $\Theta(n)$ complexity.

5. Graph Adapted Crossover

The graph adapted crossover generates genes based on the conflicting and neighboring vertices. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.
b. Choose a random vertex $v$ in $V(G)$.
c. Color the vertices with respect to the parents that have no conflicts for vertex $v$.
d. If $p_1$ and $p_2$ have conflicts with vertex $v$ then assign the least color to its neighboring vertices.

Analysis: Step (a) takes $\Theta(n)$ complexity. Step (b) requires constant running time. Step (c) requires $\Theta(n^2)$ complexity. Step (d) requires $\Theta(n^2)$ complexity. The complexity of graph adapted crossover is $\Theta(n^2)$. 
6. UIS Crossover

The UIS crossover operator unifies a pair of independent color sets from the selected parents. The procedure is described in the following steps.

a. Select two parent gene sequences, say \( p_1 \) and \( p_2 \).
b. Find the largest conflict-free subset of vertices having a color \( c \) in \( p_1 \).
c. Find the largest conflict-free subset in \( p_2 \) such that a common number of vertices in these subsets are maximal.
d. Repeat the steps (b) and (c) for every color.
e. Generate the offspring \( o_1 \) and \( o_2 \) by performing set union operation.

Analysis: Steps (a), (b) and (c) require \( \Theta(n) \) complexity. Step (d) requires \( \Theta(n \chi(G)) \) complexity. Step (e) requires \( \Theta(n) \) complexity. Thus the complexity of UIS crossover is \( \Theta(n \chi(G)) \).

7. Multi-Parent Crossover

Multi-parent crossover is an extension of GPX, which creates offspring from multiple parents. The procedure is described in the following steps.

a. Select the parent color classes from multiple parents.
b. Select the largest color class \( c \) in the population.
c. Add the color class \( c \) to the offspring.
d. Remove all vertices in \( c \) from all parent color classes.
e. Repeat the operation until the required number of color classes has reached.
f. Add the remaining vertices randomly to the color class.

Analysis: Steps (a), (b), (c) require \( \Theta(n) \) complexity. Steps (d) and (e) require \( \Theta(n^2) \) complexity. Step (f) requires \( \Theta(n) \) complexity. Thus the complexity of multi-parent crossover is \( \Theta(n^2) \).

8. WIPX Crossover

The WIPX operator selects the color classes from the randomly selected parents based on a scoring function. The procedure is described in the following steps.

a. Select the color classes from the randomly chosen parents based on a scoring function.
b. Determine the score of the color set as:

\[
\text{Number of conflicts} - \frac{1}{|V|} \left( \text{classSize} + \frac{\text{classDegree}}{n \times m} \right)
\]

where classSize is the cardinality of the color set, and classDegree is the sum of the degrees of the color class vertices.
c. Apply the multi-parent crossover operation.

Analysis: Step (a) requires \( \Theta(n) \) complexity. Step (b) requires \( \Theta(n^2) \) complexity. Step (c) requires \( \Theta(n^2) \) complexity. Thus the complexity of WIPX crossover is \( \Theta(n^2) \).

9. GPX Crossover

GPX operator partitions \( V(G) \) into \( k \) sets and generates the offspring. The procedure is described in the following steps.

a. Choose two parent configurations \( p_1 = (v_1^1, v_2^1, v_3^1 \ldots v_k^1) \) and \( p_2 = (v_1^2, v_2^2, v_3^2 \ldots v_k^2) \) where \( k \) is the number of color classes.
b. For each \( j \) (1 \( \leq \) \( j \) \( \leq \) \( k \)) do If \( j \) is odd then \( s = 1 \), else \( s = 2 \); Select \( i \) such that \( v_i^s \) has maximum vertices; \( v_1 = v_i^s \); Remove \( v_1 \) from both \( p_1 \) and \( p_2 \).

c. Choose a random color class and assign \( V - (v_1 \cup v_2 \cup v_3 \ldots \cup v_k) \).

d. Generate an offspring \( o_1 = (v_1, v_2, v_3 \ldots v_k) \).

Analysis: Step (a) requires \( \Theta(n) \) complexity. Step(b) requires \( \Theta(n^2) \) complexity. Steps (c) and (d) require \( \Theta(n) \) complexity. Thus GPX crossover takes \( \Theta(n^2) \) complexity.

10. PCPX Crossover

PCPX crossover operator generates offspring based on partitions of \( V(G) \) and its penalty. The procedure is described in the following steps.

a. Select two parent gene sequences, say \( p_1 \) and \( p_2 \).

b. Consider one parent at a time.

c. Extract a subpartition \( V'(G) \), which results in a minimum penalty among \( V(G) \) of the main partition.

d. Color \( V'(G) \) for the selected parent.

e. Remove \( V'(G) \) in both parents.

f. Find the largest color partition for the next parent.

g. Repeat the procedure until all vertices in the offspring are colored.

Analysis: Step (a) requires \( \Theta(n) \) complexity. Step(b) takes constant running time. Steps (c), (d), (e), (f) and (g) take \( \Theta(n^2) \) complexity. Thus PCPX crossover takes \( \Theta(n^2) \) complexity.

11. DBX Crossover

DBX crossover operator applies the heuristics of LDO in order based crossover or POP. The procedure is described in the following steps.

a. Apply the heuristics of LDO in order based crossover or POP operator.

b. Select a pair of random individuals, say \( P_1 \) and \( P_2 \).

c. Select a random crossover point \( r \) in \( P_1 \).

d. Copy the vertex colors starting at an initial point to \( r \) in \( P_1 \) into the offspring \( O_1 \).

e. Choose a vertex \( u \) in \( P_2 \) and \( v \) in \( O_1 \).

f. If \( u \) is not assigned an order in \( O_1 \) and if degree(\( v \)) < degree(\( u \)) then Find the vertex \( v \) having minimum order in \( O_1 \) with degree(\( v \)) < degree(\( u \)); Increment the vertex orders by 1 for all vertices having order \( \geq v \); Assign vertex \( u \) to the order of vertex \( v \); Else assign the least available order to vertex \( u \).

Analysis: Step (a) requires \( \Theta(n^2) \) complexity. The selection and copy operations in steps (b), (d), and (e) require \( \Theta(n) \) complexity. Step(c) takes a constant running time. Step (f) requires \( \Theta(n^2) \) complexity. Thus DBX takes \( \Theta(n^2) \) complexity.

12. MPGX Crossover

MPGX operator incorporates problem-specific knowledge. The procedure is described in the following steps.

a. Select a pair of random individuals, say \( P_1 \) and \( P_2 \).

b. Choose a random crossover point;

c. Assign count = \( Pc \times n \);

d. While count \( \neq 0 \) do Select a random vertex \( i \); \( c_1 = \text{color}[i] \) in \( P_1 \); \( c_2 = \text{color}[i] \) in \( P_2 \); If \( c_1 \neq c_2 \) and if penalty of \( P_1 \) reduces with \( c_2 \) then replace \( c_1 \) by \( c_2 \); If \( c_1 \neq c_2 \) and if penalty of \( P_2 \) reduces with \( c_1 \) then replace \( c_2 \) by \( c_1 \); count = count - 1
e. Generate offspring.

Analysis: Step (a) requires $\Theta(n)$ complexity. Steps (b), (c), and (e) take a constant running time. Step (d) takes $\Theta(n^2)$ complexity. Thus MPGX crossover takes $\Theta(n^2)$ complexity.

13. POP Crossover

POP crossover performs an order based permutation operation. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.

b. Choose a random point $r$ in between $[1, n]$.

c. Apply order based crossover.

d. Swap the first portion of strings and apply permutation on the remaining unused vertices copied in the sequence.

Analysis: Steps (a) and (c) require $\Theta(n)$ complexity. Step (b) requires a constant running time. Step (d) requires $\Theta(n^2)$ complexity. Thus the complexity of POP crossover is $\Theta(n^2)$.

14. MOX Crossover

The MOX operator performs merging of two-parent gene sequences and generates the order based offspring. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.

b. Merge $p_1$ and $p_2$ randomly into a single list, which consists of $2n$ elements.

c. Assign the first occurrence of each gene to offspring$_1$.

d. Assign the second occurrence of each gene to offspring$_2$.

Analysis: Steps (a) and (b) require $\Theta(n)$ complexity. Steps (c) and (d) take $\Theta(n^2)$ complexity. The complexity of MOX crossover is $\Theta(n^2)$.

15. MIS Crossover

The MIS crossover operator groups the color sets of parents and generates the offspring. The procedure is described in the following steps.

a. Select two parent gene sequences, say $p_1$ and $p_2$.

b. Group the color sets in both $p_1$ and $p_2$.

c. Copy the whole color sets from $p_1$ and $p_2$ into a single merged list.

d. Assign the first occurrence of each gene into offspring$_1$.

e. Assign the second occurrence of each gene into offspring$_2$.

Analysis: Steps (a), (b) and (c) require $\Theta(n)$ complexity. Steps (d) and (e) take $\Theta(n^2)$ complexity. The complexity of MIS crossover is $\Theta(n^2)$.

16. SPCGX Crossover

SPCGX operation is applied to the selected gene sequences, $i$ and $j$, to generate two new gene sequences as offspring. The offspring, $i'$ and $j'$ are generated by a chosen crossover probability $p_c$. SPCGX operator is defined as follows [50]:

a. Let $i = (i_1, i_2, i_3 \ldots i_q \ldots i_t \ldots i_n)$ and $j = (j_1, j_2, j_3 \ldots j_k \ldots j_l \ldots j_n)$ be the two selected sequences.

b. Generate random probability $p_{cr}$.

c. If $p_{cr} > p_c$ go to step (d).
d. Find all pairs of conflicting edges in \( i \) and \( j \).

e. Let \((q, r)\) and \((s, t)\) be such two pairs of conflicting edges (i.e., \( i_q = i_r \) and \( j_s = j_t \)) in \( i \) and \( j \), then generate two offspring: \((i_1, i_2, i_3 \ldots i_q \ldots i_{q+1} \ldots i_n)\) and \((j_1, j_2, j_3 \ldots j_s \ldots j_{s+1} \ldots j_n)\).

f. Repeat step (e) of crossover operation to all the identified conflicting pairs in step (d) of crossover operation and generate updated gene sequences \( i' = (i_1', i_2', i_3' \ldots i_q' \ldots i_n') \) and \( j' = (j_1', j_2', j_3' \ldots j_s' \ldots j_n') \).

Analysis: Step (a) requires \( \Theta(n) \) complexity. Steps (b), (c), and (e) take a constant running time. Steps (d) and (f) require \( \Theta(n^2) \) complexity. SPCGX takes \( \Theta(n^2) \) complexity.

17. SPCGX-CGR Crossover

SPCGX is applied with a CGR procedure with a chosen \( p_c \). SPCGX-CGR is given in the following steps [50]:

a. Apply the steps (a) to (e) of SPCGX crossover.

b. Check if the new genes \( r \) and \( t \) (i.e., \( i_r \) and \( j_t \)) corresponding to an edge \((r, t)\) are in conflict with its corresponding adjacent genes. If so, generate \( i_r = i_r + 1 \) and \( j_t = j_t + 1 \) such that \( i_r + 1 \leq \chi(G) \) and \( j_t + 1 \leq \chi(G) \).

c. Repeat step (f) of SPCGX with CGR procedure to all the conflicting pairs and generate updated gene sequences: \( i' = (i_1', i_2', i_3' \ldots i_n') \) and \( j' = (j_1', j_2', j_3' \ldots j_n') \).

Analysis: Steps (a) and (c) require \( \Theta(n^2) \) complexity. Step (b) takes a constant running time. SPCGX-CGR takes \( \Theta(n^2) \) complexity.

18. SPCGEX Crossover

The SPCGEX operator is applied on the selected gene sequences \( i = (i_1, i_2, i_3 \ldots i_n) \) and \( j = (j_1, j_2, j_3 \ldots j_n) \) as follows [2]:

a. Randomly select any two most conflicting vertices, \( r \) and \( t \) in gene sequences \( i \) and \( j \).

b. Generate offspring \((i_1, i_2, i_3 \ldots i_r+1 \ldots i_n)\) and \((j_1, j_2, j_3 \ldots j_t+1 \ldots j_n)\) such that \( i_r + 1 \leq \text{minimum color} \) and \( 1 + j_t \leq \text{minimum color} \).

c. Repeat the steps (a) and (b) for fixed iterations to obtain the updated offspring \( i' \) and \( j' \) as \((i_1', i_2', i_3' \ldots i_n')\) and \((j_1', j_2', j_3' \ldots j_n')\).

Analysis: Steps (a) and (c) require \( \Theta(n^2) \) complexity. Step (b) takes a constant running time. SPCGEX takes \( \Theta(n^2) \) complexity.

19. ESPCGEX Crossover

The steps in ESPCGEX crossover are defined as follows [2]:

a. Perform SPCGEX crossover steps (a) and (b).

b. If the new genes \( i_r \) and \( j_t \) are conflicting, then apply SPCGEX at the conflicting edges \((q, r)\) and \((s, t)\).

c. Repeat the steps (a) and (b) for fixed iterations to obtain the updated offspring \( i' \) and \( j' \) as \((i_1', i_2', i_3' \ldots i_n')\) and \((j_1', j_2', j_3' \ldots j_n')\).

Analysis: Steps (a) and (c) require \( \Theta(n^2) \) complexity. Hence ESPCGEX takes \( \Theta(n^2) \) complexity.

20. MSPCGX Crossover

The steps in MSPCGX crossover are defined as follows [2]:

a. Find the conflicting edges that count to every vertex in gene sequences \( j \) and \( i \).
b. Determine the conflicting vertices in \( j \) and \( i \).

c. Select arbitrarily conflicting vertices, \( r \) in \( i \) and \( t \) in \( j \).

d. Update \( i \) and \( j \) as \((i_1, i_2, i_3 \ldots i_r + 1 \ldots i_n)\) and \((j_1, j_2, j_3 \ldots j_t + 1 \ldots j_n)\) such that minimum color \( \geq i_r + 1 \) and minimum color \( \geq j_t + 1 \).

e. If the new genes \( i_r \) and \( j_t \) are conflicting, then generate \( i_r = i_r + 1 \) and \( j_t = j_t + 1 \) such that minimum color \( \geq i_r + 1 \) and minimum color \( \geq j_t + 1 \).

f. Repeat the steps (d)–(e) for remaining conflicting vertices and update \( i' \) and \( j' \) as \((i_1', i_2', i_3' \ldots i_n')\) and \((j_1', j_2', j_3' \ldots j_n')\).

Analysis: Steps (a), (b), (e) and (f) require \( \Theta(n^2) \) complexity. Step (c) takes \( \Theta(n) \) complexity. Step (d) takes a constant running time. MSPCGX takes \( \Theta(n^2) \) complexity.

### Table 1. Asymptotic Complexity of Well-known and Recent Crossover Operators.

| SNo | Crossover Operator                | Asymptotic Complexity |
|-----|----------------------------------|-----------------------|
| 1   | Order-based Uniform Crossover    | \( \Theta(n) \)       |
| 2   | Penalty Based Crossover          | \( \Theta(n^2 \chi(G)) \) |
| 3   | One-point Crossover              | \( \Theta(n) \)       |
| 4   | Two-point Crossover              | \( \Theta(n) \)       |
| 5   | Graph Adapted Crossover          | \( \Theta(n^2) \)      |
| 6   | UIS Crossover                    | \( \Theta(n \chi(G)) \) |
| 7   | Multi-Parent Crossover           | \( \Theta(n^2) \)      |
| 8   | WIPX Crossover                   | \( \Theta(n^2) \)      |
| 9   | GPX Crossover                    | \( \Theta(n^2) \)      |
| 10  | PCPX Crossover                   | \( \Theta(n^2) \)      |
| 11  | DBX Crossover                    | \( \Theta(n^2) \)      |
| 12  | MPGX Crossover                   | \( \Theta(n^2) \)      |
| 13  | POP Crossover                    | \( \Theta(n^2) \)      |
| 14  | MOX Crossover                    | \( \Theta(n^2) \)      |
| 15  | MIS Crossover                    | \( \Theta(n^2) \)      |
| 16  | SPCGX Crossover                  | \( \Theta(n^2) \)      |
| 17  | SPCGX-CGR Crossover              | \( \Theta(n^2) \)      |
| 18  | SPCGEX Crossover                 | \( \Theta(n^2) \)      |
| 19  | ESPCGEX Crossover                | \( \Theta(n^2) \)      |
| 20  | MSPCGX Crossover                 | \( \Theta(n^2) \)      |

5. Asymptotic Analysis of Some Existing Well-known and Recent Mutation Operators

Some of the existing well-known and recent mutation operators for solving GCP are presented in this section. The asymptotic complexity of these operators is also analyzed. The asymptotic complexity of these operators is also analyzed in Table 2.

1. Random Mutation

The random mutation operator randomly swaps the color of each vertex with a low mutation probability \( p_m \). The procedure is described in the following steps.

a. Select crossover offspring \( o_1 \).

b. Randomly swap the color class of each vertex with a low \( p_m \).

Analysis: Step (a) requires a constant running time. Step (b) takes \( \Theta(n) \) complexity. Thus the random mutation takes \( \Theta(n) \) complexity.

2. Polynomial Mutation

The polynomial mutation operator rounds off real values of colors to the nearest integer. The procedure is described in the following steps.
a. Select crossover offspring $o_1$.
b. Choose the random point and round off real values of colors to the nearest integer.

Analysis: Steps (a) and (b) require a constant running time. Thus the polynomial mutation takes $\Theta(c)$ complexity.

3. Problem-Specific Mutation 1

The problem-specific mutation operator is defined with problem-specific knowledge. It is operating on the offspring generated by MPGX operator. The procedure is described in the following steps.

a. Select crossover offspring $o_1$.
b. Find the missing color $c_1$ in the range of maximum color $[1, \text{maximum color}]$ used.
c. Identify all the vertices $V'(G)$ assigned with maximum color.
d. Replace colors of $V'(G)$ by $c_1$.

Analysis: Step (a) requires a constant running time. Steps (b) and (c) take $\Theta(n^2)$ complexity. Step (d) takes $\Theta(n)$ complexity. This mutation takes $\Theta(n^2)$ complexity.

4. Problem-Specific Mutation 2

The problem-specific mutation operator is defined with problem-specific knowledge. It operates on the offspring generated by MPGX operator. The procedure is described in the following steps.

a. Choose color $c_1$ randomly within one less than $[1, \text{maximum color}]$.
b. Identify all the vertices $V'(G)$ assigned to a maximum color.
c. Replace colors of $V'(G)$ by $c_1$.

Analysis: Step (a) requires a constant running time. Step (b) takes $\Theta(n^2)$ complexity. Step (c) takes $\Theta(n)$ complexity. This mutation takes $\Theta(n^2)$ complexity.

5. Problem-Specific Mutation 3

The problem-specific mutation operator is defined with problem-specific knowledge. It operates on the offspring generated by MPGX operator. The procedure is described in the following steps.

a. Choose a vertex $v$ randomly.
b. Choose a color $c$ randomly for $v$.
c. If the penalty is reduced to $v$ with color $c$, then set color[v] = c.

Analysis: Steps (a) and (b) require a constant running time. Step (c) takes $\Theta(n^2)$ complexity. This mutation takes $\Theta(n^2)$ complexity.

6. OM

The OM operator generates a random number $r$ and performs $r$ interchanges between vertices. The procedure is described in the following steps.

a. Let $(v_1, v_2, v_3 \ldots v_n)$ be the crossover offspring.
b. Generate a random number $r$;
c. Generate the offspring by performing $r$ interchanges between vertices.

Analysis: Steps (a) and (b) require a constant running time. Performing $r$ interchanges in Step(c) can also be done in linear time. The OM mutation takes $\Theta(n)$ complexity.
7. BM

The BM operation performs a translation of blocks of \( k > 0 \) successive vertices. The procedure is described in the following steps.

a. Let \((v_1, v_2, v_3 \ldots v_n)\) be the crossover offspring.

b. If \( k = 2 \) and \( i \in [1, n-1], j \in [1, n] \) are randomly generated then generate the offspring \((v_1, \ldots v_{i-1}, v_{i+2} \ldots v_j, v_i, v_{i+1}, v_{j+1} \ldots)\).

Analysis: Step (a) requires a constant running time. Step (b) can take \( \Theta(n) \) complexity. Thus BM operation takes \( \Theta(n) \) complexity.

8. CSM

The CSM randomly selects the conflicting edge and moves the genes to new positions. The procedure is described in the following steps.

a. Let \((v_1, v_2, v_3 \ldots v_n)\) be the crossover offspring.

b. Choose a randomly selected bad edge \( e = (v_i, v_j) \) and \( e' = (v_k, v_l) \) be the next bad edge such that \( i < j < k < l \).

c. Move the vertices from positions \( i + 1 \) to \( l - 1 \) to randomly chosen new positions.

Analysis: Step (a) requires a constant running time. Steps (b) and (c) need \( \Theta(n) \) complexity.

9. BESM

The BESM operation reduces the number of conflicting edges. The procedure is described in the following steps.

a. Let \((v_1, v_2, v_3 \ldots v_n)\) be the crossover offspring.

b. Choose a random bad edge \( e = (v_i, v_j) \).

c. Choose a direction either left-to-right or right-to-left randomly.

d. Let \( v_k \) be the farthest vertex adjacent to \( v_i \) so that \( k > j \).

e. Let \( e' = (v_l, v_m) \) be the first bad edge so that \( l > k \).

f. If there is no vertex \( v_k \) or edge \( e' \) with the specified properties, the mutation is void. Otherwise, the vertex \( v_i \) is moved to the position \( m + 1 \).

Analysis: Steps (a), (c) require a constant running time. Step (b) takes \( \Theta(n^2) \) complexity. Steps (d), (e) and (f) take \( \Theta(n) \) complexity. Hence BESM takes \( \Theta(n^2) \) complexity.

10. CGM

The CGM operates on offspring \( i' \) and \( j' \). The steps are given below [49]:

a. Select any two conflicting genes \( i'_{p} = i'_{g'} \) and \( j'_{k'} = j'_{l'} \) in \( i' \) and \( j' \).

b. Update \( i' \) and \( j' \) as \((i_{1'}, i_{2'}, i_{3'} \ldots i_{p-1} \ldots i_{n'}) \) and \((j_{1'}, j_{2'}, j_{3'} \ldots j_{k-1} \ldots j_{n'}) \) such that \( 1 \leq i_{p-1} \) and \( 1 \leq j_{k-1} \).

c. Update \( i'' \) and \( j'' \) as \((i_{1''}, i_{2''}, i_{3''} \ldots i_{n''}) \) and \((j_{1''}, j_{2''}, j_{3''} \ldots j_{n''}) \) by repeating steps (a)-(b) for fixed iterations.

Analysis: Steps (a) and (c) require \( \Theta(n^2) \) complexity. Step (b) requires a constant running time. Hence CGM mutation takes \( \Theta(n^2) \) complexity.
11. CGM-CGR Mutation

The CGM-CGR mutation is applied to the offspring $i'$ and $j'$ and its steps are given below [50]:

a. Perform CGM steps (a)–(b).

b. If the new genes $i_f'$ and $j_k'$ are conflicting then update $i_f' = i_f' - 1$ and $j_k' = j_k' - 1$ such that $1 \leq i_f' - 1$ and $1 \leq j_k' - 1$.

c. Update $i''$ and $j''$ as $(i_1'', i_2'', i_3'' \ldots i_n'')$ and $(j_1'', j_2'', j_3'' \ldots j_n'')$ by repeating steps (a)–(b) for fixed iterations.

Analysis: Steps (a), (b) and (c) require $\Theta(n^2)$ complexity. Hence CGM-CGR mutation takes $\Theta(n^2)$ complexity.

12. ECGM

The ECGM operation is defined as follows [2]:

a. Perform CGM steps (a) and (b).

b. If new genes $i_f'$ and $j_k'$ are conflicting, then perform CGM at $f'$ and $k'$.

c. Update $i''$ and $j''$ as $(i_1'', i_2'', i_3'' \ldots i_n'')$ and $(j_1'', j_2'', j_3'' \ldots j_n'')$ by repeating steps (a) and (b) for fixed iterations.

Analysis: Steps (a), (b) and (c) require $\Theta(n^2)$ complexity. Hence ECGM takes $\Theta(n^2)$ complexity.

13. MCGM

The MCGM operation is defined as follows [2]:

a. Find the conflict edges count to every vertex in $i'$ and $j'$.

b. Determine the conflict vertices in $i'$ and $j'$.

c. Select arbitrarily two conflicting vertices $f'$ in $i'$ and $k'$ in $j'$.

d. Update $i'$ and $j'$ as $(i_1', i_2', i_3' \ldots i_{f'} - 1 \ldots i_n')$ and $(j_1', j_2', j_3' \ldots j_{k'} - 1 \ldots j_n')$ such that $1 \leq i_{f'} - 1$ and $1 \leq j_{k'} - 1$.

e. If new genes $i_f'$ and $j_k'$ are conflicting then perform CGM at $f'$ and $k'$.

f. Update $i''$ and $j''$ as $(i_1'', i_2'', i_3'' \ldots i_n'')$ and $(j_1'', j_2'', j_3'' \ldots j_n'')$ by repeating steps (d) and (e) for fixed iterations.

Analysis: Steps (a), (b), (e) and (f) require $\Theta(n^2)$ complexity. Steps (c) and (d) take $\Theta(c)$ complexity. Hence MCGM takes $\Theta(n^2)$ complexity.

**Table 2.** Asymptotic Complexity of Well-known and Recent Mutation Operators.

| SNo | Mutation Operator             | Asymptotic Complexity |
|-----|-------------------------------|-----------------------|
| 1   | Random Mutation               | $\Theta(n)$           |
| 2   | Polynomial Mutation           | $\Theta(c)$           |
| 3   | Problem-Specific Mutation 1   | $\Theta(n^2)$         |
| 4   | Problem-Specific Mutation 2   | $\Theta(n^2)$         |
| 5   | Problem-Specific Mutation 3   | $\Theta(n^2)$         |
| 6   | OM                            | $\Theta(n)$           |
| 7   | BM                            | $\Theta(n)$           |
| 8   | CSM                           | $\Theta(n)$           |
| 9   | BESM                          | $\Theta(n^2)$         |
| 10  | CGM                           | $\Theta(n^2)$         |
| 11  | CGM-CGR                       | $\Theta(n^2)$         |
| 12  | ECGM                          | $\Theta(n^2)$         |
| 13  | MCGM                          | $\Theta(n^2)$         |
6. Global Convergence of Evolutionary Algorithms and Stochastic Convergence of Recent Genetic Operators

The necessary and sufficient condition for the global convergence of evolutionary algorithms is proved in this section \[61,62\]. The stochastic convergence of some recent genetic operators for solving GCP is also proved in the present section \[2\].

6.1. Markov Chain Model of Evolutionary Algorithms

Consider the following optimization problem:
Optimize \( f(x); x \in S \) where \( S \) is a measurable space, \( f(x) \) is the objective or fitness function where the absolute value of \( f(x) \) is \(< +\infty\).

Represent the optimal solution set by \( S^* = \{ x' \text{ such that } f(x^*) = \text{optimize } f(x), x \in S \} \).

Let \( \mu(S) \) be a measure to space \( S \) with \( \mu(S^*) > 0 \).

Consider the set \( S_0 = \{ x \text{ such that } f(x^*) - f(x) < \delta \} \) where \( \delta > 0 \) is a small number.

Choose an appropriate \( \delta \) to satisfy \( \mu(S_0) > 0 \).

The space \( S \) is called an individual space where each state \( s \in S \) is called an individual.

\( N \) represents population size.

\( X = S^N \) is the population space.

Denote the elements in \( x \) as \( x = [x_1, x_2, x_3 \ldots x_N] \).

Denote the optimal solution set \( X^* = \{ x' \text{ such that } \exists x_i \in x: x_i \in S^* \} \).

An evolutionary algorithm to solve this optimization problem is formulated as follows:

a. Initialization: Generate initial population \( P_0 \) at generation \( g = 0 \).

b. Crossover operation: Update the new population \( P_g \) by a crossover operator \( P_C(g) \).

c. Mutation operation: Update the new population \( P_g \) by a mutation operator \( P_M(g) \).

d. Selection operation: Select a new population \( P_S(g) \) by a selection operator \( P_S(g) \).

e. Termination: If termination conditions hold, then stop; otherwise \( g = g + 1 \) and update \( P_g \).

Since the state of \( P_{g+1} \) is only dependent on the state of \( P_g \), then \( \{ P_g; g = 0, 1, 2 \ldots \} \) can be modeled by a Markov chain with the following transition function \[61,62\]:

\[
P(g; x, dy) = P(P_{g+1} = dy \text{ such that } P_g = x) = \int_{u \in X} \int_{v \in X} P_C(g; x, du)P_M(g; u, dv)P_S(g; v, dy) \quad (1)
\]

Then \( \{ P_g; g = 0, 1, 2 \ldots \} \) converges to \( X^* \), if for any initial population \( P_0 \),

\[
\lim_{g \to +\infty} \mu_g(X^*) = 1 \text{ where } \mu_g(X^*) = \mu(P_g \in X^*).
\]

6.2. Convergence Conditions of Evolutionary Algorithms

The necessary and sufficient condition for the convergence of evolutionary algorithms is as follows \[61,62\]:

Let \( \{ P_g; g = 0, 1, 2 \ldots \} \) be the Markov chain given by (1). Define \( \alpha(g) \) as the difference between flow from the non-optimal solution set to the optimal solution set and vice versa.

\[
\alpha(g) = \int_{x \notin X^*} P(P_{g+1} \in X^* | P_g = x) \mu_g(dx) - \int_{x \in X^*} P(P_{g+1} \notin X^* | P_g = x) \mu_g(dx)
\]

Then \( \{ P_g; g = 0, 1, 2 \ldots \} \) converges to the optimal solution set \( X^* \) iff

\[
\sum_{g=0}^{+\infty} \alpha(g) = 1 - \mu_0(X^*)
\]
Hence the new gene takes one of the values in \((\text{incremental operators}) \text{SPCGX and SPCGEX produce a new gene as}\)

\[
\mu_{g+1}(X') = \int_{x \not\in X'} P(P_{g+1} \in X' | P_g = x) \, \mu_g(dx) + \int_{x \in X'} P(P_{g+1} \not\in X' | P_g = x) \, \mu_g(dx)
\]

\[
\mu_{g+1}(X') = \mu_g(X') - \int_{x \not\in X'} P(P_{g+1} \not\in X' | P_g = x) \, \mu_g(dx) + \int_{x \in X'} P(P_{g+1} \in X' | P_g = x) \, \mu_g(dx)
\]

\[
\mu_{g+1}(X') - \mu_g(X') = \alpha(g)
\]

\[
\mu_{g+1}(X') - \mu_0(X') = \sum_{k=0}^{\infty} \alpha(k)
\]

\[
\lim_{g \to \infty} \mu_{g+1}(X') = 1 \iff \sum_{g=0}^{\infty} \alpha(g) = 1 - \mu_0(X')
\]

The initial population \(P_0\) is not optimal, that is, \(\mu_0(X^*) = 0\). Then

\[
\sum_{g=0}^{\infty} \alpha(g) = 1
\]

6.3. Stochastic Convergence of Recent Evolutionary Operators

Some of the recent genetic operators like SPCGX, SPCGEX, ESPCGEX, ECGM, MSPCGX, and MCGM modify \(P_g\) through successive stochastic transformations, which can be analyzed by the Markovian model. The subsequent value of \(P_{g+1}\) is generated stochastically, and the evolutionary algorithm converges when the reachability and monotone conditions are fulfilled \([2,41,63]\). Some of the recent genetic operators converge stochastically.

6.3.1. Analyzing the Reachability Condition from \(j'\)

If \(j\) and \(j'\) be any two individuals in the finite search space \(S\), then \(j'\) is reachable from individual \(j\). That is, \(j' = (j_1', j_2', j_3' \ldots j_n')\) and \(j'' = (j_1'', j_2'', j_3'' \ldots j_n'')\) are the two gene sequences generated from the gene sequences \(j = (j_1, j_2, j_3 \ldots j_n)\) and \(j = (j_1, j_2, j_3 \ldots j_n)\) respectively. If \((0 < \text{Probability}\{ \forall j'' = (\text{crossover}(j) \& \text{mutation}(j'))\} < 1)\) then the individual \(j''\) is reachable from individual \(j'\) \([2]\).

First, consider the effect of crossover operation. Let \(S\) be the finite search space of the problem. Consider a gene sequence \(j \in S\) such that \(j = (j_1, j_2, j_3 \ldots j_n)\). The proposed crossover operators (incremental operators) SPCGX and SPCGEX produce a new gene as \(j_r = j_r + 1\) if \(\text{minimum color} \geq j_r + 1\). Hence the new gene takes one of the values in \((j_r, j_r+1)\). The CGR operator further checks the conflict at vertex \(r\). If there is a conflict, then the new gene at vertex \(r\) is generated as \(j_r = j_r + 1\) such that \(\text{minimum color} \geq j_r + 1\). Clearly, \(j_r\) assumes the value in \((j_r, j_r + 1, j_r + 2)\). Thus the required probability of assigning a new gene at the conflicting vertex is \(1/3\) for a single crossover operation. Assume that the crossover operation is performed for \(k\) times.

\[
\Rightarrow \text{Probability}\{ j' = \text{crossover}(j) \} = 1/3^k > 0 (1 \leq j, j' \leq N), k > 0
\]

Now consider the effect of mutation operation. The recent mutation operators set a value of the conflicting gene in \([1, f(G)]\). Thus the required probability of assigning a gene at the conflicting vertex is \(1/f(G)\) for a single mutation operation. Assume that the mutation operation is performed for \(l\) times.

\[
\Rightarrow \text{Probability}\{ j'' = \text{mutation}(j') \} = 1/f(G)^l > 0, l > 0.
\]
1. The values of 1000 runs each with 5,000,000 generations of the genetic algorithm, achieve a better near-optimal solution [2] after a finite number of generations. Hence, the successive probabilistic changes utilize the elitism operation.

2. For small graphs, some of the recent operators monotonically reduce $F$ during the iterations of an evolutionary algorithm, $P_g$ is modified by successive probabilistic transformations. The proportionate fitness selection selects one worst and two better gene sequences in every generation as follows:

   a. Evaluate $F_g(a)$ for each gene sequence $a (1 \leq a \leq N)$ in $P_g$.
   b. Determine $p(a)$ using
      \[
      p(a) = \frac{F_g(a)}{\sum_{a=1}^{N} F_g(a)}
      \]
   c. Compute $E(a) = N \cdot p(a)$ for all $a (1 \leq a \leq N)$.
   d. Select the better gene sequences $i$ and $j$ and the worst gene sequence $w$.

   The probability of selection of $i$ and $j$ is always greater than zero. $P_g$ is then updated by performing the elitism operation.

   The recombination and mutation operations are performed based on the values of $p_{cr}$ and $p_{mr}$. For small graphs, some of the recent operators monotonically reduce $F_g(i)$ and $F_g(j)$ during the modification of population by a finite number of probabilistic transformations [2]:

   1. The values of $F_g(i)$ and $F_g(j)$ monotonically decreases in subsequent generations and converge to a better near-optimal solution.

      That is, either $F_0(i) \geq F_1(i) \geq F_2(i) \geq \ldots \geq F_s(i) = 0$ or $F_0(j) \geq F_1(j) \geq F_2(j) \geq \ldots \geq F_s(j) = 0$.

   2. $F_g(i)$ and $F_g(j)$ monotonically decrease for a finite number of generations (say $t$).

      i.e., $F_0(i) \geq F_1(i) \geq F_2(i) \geq \ldots \geq F_t(i)$ and $F_0(j) \geq F_1(j) \geq F_2(j) \geq \ldots \geq F_t(j)$.

      Then, based on the values of $p_{cr}$ and $p_{mr}$, the fitness-function values again increase for a finite number of generations (say $s$).

      i.e., $F_t(i) \leq F_{t+1}(i) \leq \ldots \leq F_s(i)$ and $F_t(j) \leq F_{t+1}(j) \leq \ldots \leq F_s(j)$.

   For complex graphs, the successive probabilistic transformations find a better near-optimal solution after a finite number of generations. Hence, the successive probabilistic changes utilize $P_g$ as a monotone in some of the subsequent generations to achieve a better near-optimal solution. These successive probabilistic changes, applying for over a large number of generations, for example, 1000 runs each with 5,000,000 generations of the genetic algorithm, achieve a better near-optimal solution [2].

For the chosen crossover probability $p_c$ and mutation probability $p_m$, the probability of generating new gene sequence $j''$ is computed using the multiplication theorem of probability. The crossover and mutation operations are mutually independent. Hence the required probability of generating $j''$ is computed as follows:

   Probability of $\{j'' = \text{(crossover} (j) \text{& mutation} (j'))\} \geq$

   \[
   \text{(Crossover probability)} \times (\text{Probability of generating } j' \text{ from } j) \times (\text{Mutation probability}) \times (\text{Probability of generating } j'' \text{ from } j')
   \]

   That is

   \[
   \text{Probability of } \{j'' = \text{(crossover} (j) \text{& mutation} (j'))\} \geq p_c \cdot p_a \cdot p_m \cdot p_c^2 \cdot p_m \cdot \frac{1}{3} \cdot \frac{1}{f(G)} \geq 0
   \]

   Hence the gene sequence $j''$ is reachable from $j'$.

6.3.2. Analyzing whether $P_g$ is Monotone

During the iterations of an evolutionary algorithm, $P_g$ is modified by successive probabilistic transformations. The proportionate fitness selection selects one worst and two better gene sequences in every generation as follows:

   a. Evaluate $F_g(a)$ for each gene sequence $a (1 \leq a \leq N)$ in $P_g$.
   b. Determine $p(a)$ using
      \[
      p(a) = \frac{F_g(a)}{\sum_{a=1}^{N} F_g(a)}
      \]
   c. Compute $E(a) = N \cdot p(a)$ for all $a (1 \leq a \leq N)$.
   d. Select the better gene sequences $i$ and $j$ and the worst gene sequence $w$.

   The probability of selection of $i$ and $j$ is always greater than zero. $P_g$ is then updated by performing the elitism operation.

The recombination and mutation operations are performed based on the values of $p_{cr}$ and $p_{mr}$. For small graphs, some of the recent operators monotonically reduce $F_g(i)$ and $F_g(j)$ during the modification of population by a finite number of probabilistic transformations [2]:

1. The values of $F_g(i)$ and $F_g(j)$ monotonically decreases in subsequent generations and converge to a better near-optimal solution.

   That is, either $F_0(i) \geq F_1(i) \geq F_2(i) \geq \ldots \geq (F_s(i) = 0)$ or $F_0(j) \geq F_1(j) \geq F_2(j) \geq \ldots \geq (F_s(j) = 0)$.

2. $F_g(i)$ and $F_g(j)$ monotonically decrease for a finite number of generations (say $t$).

   i.e., $F_0(i) \geq F_1(i) \geq F_2(i) \geq \ldots \geq F_t(i)$ and $F_0(j) \geq F_1(j) \geq F_2(j) \geq \ldots \geq F_t(j)$.

   Then, based on the values of $p_{cr}$ and $p_{mr}$, the fitness-function values again increase for a finite number of generations (say $s$).

   i.e., $F_t(i) \leq F_{t+1}(i) \leq \ldots \leq F_s(i)$ and $F_t(j) \leq F_{t+1}(j) \leq \ldots \leq F_s(j)$.

For complex graphs, the successive probabilistic transformations find a better near-optimal solution after a finite number of generations. Hence, the successive probabilistic changes utilize $P_g$ as a monotone in some of the subsequent generations to achieve a better near-optimal solution. These successive probabilistic changes, applying for over a large number of generations, for example, 1000 runs each with 5,000,000 generations of the genetic algorithm, achieve a better near-optimal solution [2].
7. Conclusions and Future Work

GCP is an NP-hard combinatorial optimization problem and can be applied to innumerable engineering problems. $\chi(G)$ can be obtained using different approximations and evolutionary methods. In comparison to other methods, genetic algorithms are useful in solving multi-objective optimization problems with vast search space [2]. Hence, it is expected to design new evolutionary operators to achieve faster stochastic convergence with a minimal number of generations.

This paper explored the asymptotic analysis of some well-known and recent evolutionary operators for finding chromatic numbers. The asymptotic analysis of different crossover and mutation operators helps in minimizing problem search space and computational complexity. The choice of the right genetic operators facilitates an evolutionary algorithm to achieve faster convergence with lesser population size (N) through an effective distribution of promising genes. The selection of evolutionary operators also plays an essential role in reducing the bounds for minimum color obtained for the benchmark graphs [2]. The necessary and sufficient condition for global convergence of the evolutionary algorithm and the stochastic convergence of some recent evolutionary operators for solving GCP are analyzed in this paper.

Our current work enlightens some new research directions [64–66].

i. The recent genetic operators can be combined with better local search strategies in order to further reduce the computational complexity of GCP.

ii. The self-adaptive evolutionary operators with heuristics can be designed to find chromatic numbers as well as can be applied to channel allocation problem (CAP) and scheduling problems. CAP is an extension of GCP, which assigns channels to mobile stations.

iii. The performances of some recent operators can be evaluated under different parameter values, such as graph density, $N$, $p_c$, and $p_m$.

iv. Some of the recent operators can be combined to reduce $\bar{G}$.

v. The stochastic convergence of genetic algorithms can be extended to multi-objective optimization problems, and the necessary conditions for the global convergence of multi-objective optimization problems can be analyzed mathematically.

vi. An algebraic framework that may lead to a unification of different evolutionary operators for combinatorial problems can be designed and analyzed [64–66].

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