COMPARISON OF GENETIC OPERATORS ON A
GENERAL GENETIC ALGORITHM
PACKAGE

By

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PACKAGE

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Chapter I. Introduction to Genetic Algorithms

1.1 General Introduction

Genetic algorithms are now widely applied in science and engineering as stochastic algorithms for solving practical optimization problems. Genetic algorithms (GA) were first introduced by John H. Holland in his fundamental book “Adaptation in Natural and Artificial Systems” in 1975 [4]. Holland presented the algorithm as an abstraction of biological evolution and his schema theory laid a theoretical foundation for GA.

The idea behind GAs is to simulate what nature does. Simply speaking, GA is a simulation of Darwin’s Theory of Evolution – the fittest survive. A simple GA works as follows: First, a number of individuals (the population) are randomly initialized at the beginning of the algorithm. Individuals are then selected according to their fitness. Next, the genetic operators (crossover, mutation) are applied with certain probabilities on these selected individuals, the parents, to produce offspring. The original generation is then replaced by the new generation which consists in whole or in part of the newly created offspring. The above process is repeated if the termination criterion is not met; otherwise, the algorithm stops. [2]

Whereas in nature the "fitness" relates to the ability of the organism to survive and reproduce, that is, organisms with a better "fitness" score are more likely to be selected for reproduction, in genetic algorithms, the "fitness" is the evaluated result of a user-defined objective function. [1]
1.2 Biological Terminology

Let us introduce some of the basic biological terminology that is useful for a better understanding of GAs.

Each cell of a living creature consists of a certain set of chromosomes. Chromosomes are made of genes, which are blocks of DNA. Each gene encodes one or more characters (eye color, hair color, etc) that can be passed on to the next generation. Each gene can be in different states, called alleles. Genes are located at certain positions on the chromosome, which are called loci.

The cell of many creatures has more than one chromosome. The entire set of chromosomes in the cell is called the genome. If the chromosomes in each cell of the organism are unpaired, the organism is called haploid. If, on the other hand, the chromosomes are paired, the organism is called diploid. In nature, many living organisms are diploid, but almost all GAs employ haploid representations, since they are simple to construct.

In the natural reproduction process, pieces of gene material are exchanged between the two parents’ chromosomes to form new genes. This process is called recombination or crossover. Genes in the offspring are subject to mutation, in which a certain block of DNA in the gene undergoes a random change.

In genetic algorithms, a chromosome is used to represent a potential solution to a problem. Since we use single-chromosome individuals to represent the problem solution, the term individual and chromosome are often used interchangeably.
1.3 How GAs Work

Despite its amazing power, a genetic algorithm is actually quite elegant and so easy to understand that it can be expressed in just a few lines of computer pseudo-codes.

Following is the C version of the GA structure originally presented in [1]:

```c
Procedure Simple GA
{
    t=0;
    initialize P(t);
    evaluate P(t);
    while(! termination-condition)
    {
        select P(t+1) from P(t);
        genetic_operate P(t+1);
        evaluate P(t+1);
        t++;
    }
}
```

where, $P(t)$ is the population of individuals for iteration $t$.

Here is a more descriptive version of GA, slightly modified from [1]:

1. **[Initialize]** Generate a random population of $n$ chromosomes (suitable solutions for the problem).
2. **[Evaluate]** Evaluate the fitness $f(x)$ of each chromosome $x$ in the population.
3. **[Offspring]** Create a new population by executing the following steps.
   
   a. **[Selection]** Select $n$ parent chromosomes from the population according to their fitness (the better the fitness, the better the chance to be selected).
   
   b. **[Crossover]** Recombine the parents with a certain crossover probability to form new offspring.
   
   c. **[Mutation]** Mutate the new offspring with certain mutation probability at each locus (position in chromosome).
4. **[Replace]** Replace the current population with the newly generated population.
5. **[Test]** If the termination condition is satisfied, stop, and return the best chromosome found; otherwise, go to step 2.
The convergence of a genetic algorithm is originally based on Holland’s schema theory \cite{Holland}. However, his theory has raised criticism and controversy over the years. Since this thesis is focused on the real world application of GAs instead of theoretical deduction, interested readers may refer to \cite{5} for a precise mathematical description of genetic algorithms.

1.4 An Example

A good example speaks better than thousands of words. Let’s work on a simple example, One Max \cite{6}, to illustrate the basic steps of GA. As its name says, the goal of One Max is to maximize the number of 1’s in 8-bit-strings such as 10010011.

First, we create an initial population of individuals. For simplicity, we assume a population of size 4. The value of each individual is initially assigned randomly. Then the population is evaluated based on a fitness function to determine how well each individual does the required task. In our case, the fitness function is straightforward:

\[
f(v) = n, \text{ where } n \text{ is the number of } 1\text{’s in individual } v.
\]

For example, the fitness value of an individual with chromosome 10011001 would be 4. After initialization, a random population of four individuals and their fitness values are listed in Table 1.1.

<table>
<thead>
<tr>
<th>Individual</th>
<th>String Representation</th>
<th>Fitness</th>
</tr>
</thead>
<tbody>
<tr>
<td>V1</td>
<td>01101001</td>
<td>4</td>
</tr>
<tr>
<td>V2</td>
<td>00100100</td>
<td>2</td>
</tr>
<tr>
<td>V3</td>
<td>11110001</td>
<td>5</td>
</tr>
<tr>
<td>V4</td>
<td>00001110</td>
<td>3</td>
</tr>
</tbody>
</table>

Table 1.1 One max individual values
Next, select two parents to generate two offspring. In this example, we use the plain but widely adopted fitness-proportionate selection scheme, in which the probability of an individual being chosen to reproduce is proportional to its fitness value. In our case, let’s assume that the parents selected are [V3, V1] and [V3, V4], with V3 being selected twice because of its higher fitness. Note that V2 may also be selected, with a lower probability, in real world applications.

After the selection, we are ready to apply the crossover operator to the selected individuals. Suppose the crossover probability is 0.5, which means 50% of the population will undergo crossover (2 individuals in our case). Let’s say parent V3 and V1 undergo crossover after the fourth bit, the process is shown in the figure below:

![Figure 1.3 Single point crossover demonstration](image)

The offspring generated are V1’ = 01100001 and V3’ = 11111001. On the other hand, the offspring of V3 and V4, which do not go through crossover, are the exact copies of themselves. Suppose we utilize generational replacement model, i.e., replace the parents with the newly generated offspring, the new population now becomes:

<table>
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<th>Fitness</th>
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</thead>
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<tr>
<td>V1’</td>
<td>01100001</td>
<td>3</td>
</tr>
<tr>
<td>V2’ (V3)</td>
<td>11110001</td>
<td>5</td>
</tr>
<tr>
<td>V3’</td>
<td>11111001</td>
<td>6</td>
</tr>
<tr>
<td>V4</td>
<td>000001110</td>
<td>3</td>
</tr>
</tbody>
</table>

Table 1.2 Population after crossover
Next, the mutation operator is applied to individuals in the new population with a certain mutation probability. Suppose V4 is mutated at the 8th bit position, the resultant V4’ is 00001111 with a fitness value 4.

At this point, one cycle has been finished. Repeat the above steps. In the second round selection, V3’ and V4’ may be chosen as parents, and the offspring after crossover are V3’’=11111111, V4’’=00001001. Suppose the mutation happens to V2’ at the second position, which results in V2’’=10110001. After the second iteration, the population now is:

<table>
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<tr>
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<th>Fitness</th>
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</thead>
<tbody>
<tr>
<td>V1’</td>
<td>01100001</td>
<td>3</td>
</tr>
<tr>
<td>V2’’</td>
<td>10110001</td>
<td>4</td>
</tr>
<tr>
<td>V3’’</td>
<td>11111111</td>
<td>8</td>
</tr>
<tr>
<td>V4’’</td>
<td>00001001</td>
<td>2</td>
</tr>
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</table>

Table 1.3 Final population

We can see that individual V3’’ has fitness value 8, which is the maximum we are looking for; thus the GA can stop here if we have some way of knowing we have reached the optimal solution.

Interested readers can observe an animated java applet demonstration on [9] for a vivid picture of how a GA works.

The simple procedure just described is the basis for most applications of GAs. There are a number of details to fill in, such as the size of the population and various methods of crossover and mutation. The success of the algorithm often depends greatly on those parameters. We will talk more about them in the following chapters.
Chapter II. Literature Review

2.1 Brief Introduction to Optimization

The purpose of optimization is to find the minimum or maximum result of a mathematical function, called the objective function, by tuning the values of its variables. During the past decades, the significance of optimization has grown dramatically in engineering application, mathematical and computational models. Consequently, dozens of optimization methods have been developed over the years.

The exhaustive search method searches a sufficiently large sampling space of the objective function to find the global optimum. This brute force approach involves a huge number of function evaluations, which makes it only practical for a small number of parameters in a limited search space. \[7\]

Calculus optimization is a mathematical approach that can be applied if the objective function is smooth (continuously differentiable). First, the objective function is differentiated with respect to all the independent variables; next, the derivatives of function are set to zero and the equations are then solved to find the parameter values. This method is more computationally efficient than many of its peers. However, if the problem has too many parameters, then it becomes quite difficult or even impossible to solve the derivative equations, let alone that some practical problems can’t be expressed as mathematical functions \[7\]. Also, the nonlinear equations may have no solution, many solutions, or even infinity of solution points.
A hill-climbing method starts from a single point and selects a new point from the vicinity of the current point in each iteration. If the new point has a better value, the current point is replaced by the new point. Otherwise, the current point is kept and the process continues with a smaller step size or in another direction until the termination criterion is met. The drawback of this method is that it only guarantees a local optimum, which is determined by the position of the start point. Also, the objective function must be smooth (continuously differentiable) for most hill-climbing methods to succeed.

Two relatively new natural optimization methods, Simulated Annealing (SA) and Genetic Algorithm (GA) are successful in the area where the traditional methods fall short. Whereas GA models the natural selection and evolution process, SA simulates the annealing phenomenon. Compared with hill-climbing, SA introduces a probability of accepting the new point and the probability is controlled by an additional parameter $T$, the temperature. With a high temperature at the beginning, the algorithm is more likely to accept new points, and thus explore more regions in the solution space. As the temperature goes down, the probability of accepting new points becomes smaller and smaller. Since $T$ is lowered in small steps, the algorithm has an opportunity to discover the global optimum instead of converging to a local optimal point \[8\]. Some published SA algorithms fail on many practical problems \[24\]. Perhaps the best existing SA software package is the free C language program ASA, which was developed by Lester Ingber and is downloadable at [25].

Compared with traditional optimization methods, GA has advantages in several aspects\[7\]: it can optimize both continuous and discrete functions; it does not require
complicated differential equations or a smooth objective function; it can be highly parallelized to improve computational performance; it searches more of the solution space and thus increases the probability of finding the global optimum. The disadvantage is that GA is often much slower than calculus methods, when the latter can used.

2.2 GA Packages Review

Due to their inherent advantages, GAs have been widely applied in numerous scientific and engineering problems. A number of GA packages have been developed as implementations of genetic algorithms over the years. In this section, we will give a short review of existing GA packages.

2.2.1 GAlib

GAlib is a C++ genetic algorithm package developed at MIT (GAlib should not be confused with GALib, developed at the University of Tulsa). This genetic algorithm will create a population of solutions based on a sample data structure that you provide. In GAlib, the sample data structure is called a GAGenome, which functions as a chromosome. The library contains four types of genomes: GAListGenome, GATreeGenome, GAArrayGenome, and GABinaryStringGenome.

Each genome type has three operators: initialization, mutation, and crossover.

- Initialization operator: Initializes a genome when the genetic algorithm starts.
- Mutation operator: Mutates each genome based on its data type. For example, mutation on a binary string genome flips the bits in the string, but mutation on a tree would swap subtrees.
• Crossover operator: Generates children from two parent genomes based on data types. Users are allowed to develop new mating methods other than the default ones.

GAlib comes with these operators pre-defined for each genome type, but you can customize any of them so that it is specific not only to the data type, but also to the problem type.

The library contains four basic types of genetic algorithms.

• Simple genetic algorithm: The classical replacement method in which the entire generation is replaced by the newly produced individuals.

• Steady-state genetic algorithm: Only a fraction of the population is replaced by the offspring. The user can specify how large the fraction is.

• Incremental genetic algorithm: Only one or two children are created in each generation and the users have the choices to select the parents that are to be replaced. For example, the child could replace an individual randomly, or replace an individual that is most like it.

• Deme genetic algorithm: This algorithm “evolves multiple populations in parallel using a steady-state algorithm”. In each generation, some of the individuals are copied from one population to the other to maintain the population diversity.

2.2.2 Genesis\textsuperscript{[11]}

Genesis, which stands for GENEtic Search Implementation System, is a popular genetic algorithm package developed by John J. Grefenstette using C language.
Genesis has three levels of representation for its evolving structures. The lowest level
(Packed representation) is used to “maximize both space and time efficiency in
manipulating structures”. The middle level (String representation) represents
structures as arrays of characters. This level provides the users the freedom to create
their own genetic structures. The third level (Floating-point representation) represents
the genetic structures as vectors of real numbers. Numeric optimization problems are
solved by working on this level.

The user can specify the range, number of values, and output format of each
parameter. The system then automatically translates the structure among the three
representation levels.

Genesis consists of two main modules – initialization and generation. Initialization
module sets up the initial population using a file named "init.c". The generation
module (file "generate.c") comprises mainly the following procedures: selection,
mutation, crossover, and evaluation.

- Selection: The default selection procedure is Universal Sampling Selection
  implemented in file "select.c". Genesis also provides users the linear ranking
  selection method if the option ‘R’ is specified.
- Mutation: Genesis implements the classical mutation method – Uniform Mutation
  in file “mutate.c”.
- Crossover: Genesis implements the traditional two-point crossover method in file
  "cross.c".
Genesis requires the user to write an evaluation procedure to serve as the objective function. The procedure must be declared in the user's input file in a certain format.

Genesis allows a number of options to employ different strategies during the search. Each option is associated with a single character. For instance,

- **E** - Use the "elitist" selection strategy. The elitist strategy stipulates that the best performing structure always survives from one generation to the next.
- **G** - Use Gray code to encode integers.

### 2.2.3 Genetic Algorithm Driver [15]

The FORTRAN Genetic Algorithm Driver is a GA package created by D.L. Carroll. The latest version of this driver is dated 2001. The features of this package are listed as follows:

- **Encoding**: Bits string, which is the only option provided by this package.
- **Selection**: Tournament selection with “a shuffling technique for choosing random pairs for mating”.
- **Mutation**: The package implements jump mutation and creep mutation.
- **Crossover**: The package provides the option for single-point or uniform crossover.

In the latest version, the author added two features into the package. One is niching, the process of dividing individuals that are in some sense similar into groups; the other is micro-GA, the process of reinitializing a small subset of population.

For the use of his package, Carroll recommends the following settings:

- Binary coding (the only option of his GA)
- Tournament selection (the only option of his GA)
- Uniform crossover (iunifrm=1)
- Creep mutations (icreep=1)
- Niching or sharing (iniche=1)
- Elitism (ielite=1)

Carroll also suggests using the micro-GA technique (microga=1) with uniform crossover (iunifrm=1). Besides that, he advises using certain formulas to calculate the population size and creep mutation rate. Interested readers may refer to [15] for the detailed formulas.

### 2.2.4 Other GA Packages

Besides the GA packages introduced above, there are quite a number of other good ones such as GENETIC, GENOCOP, GAJIT, etc. GA-Archive\(^{[13]}\) is an excellent source to explore the existing GA packages. GA packages on that website are either archived as compressed files or maintained as links to the package authors’ web pages. There is an index of source code for implementations of genetic algorithms. The GA packages listed on the site are categorized according to the implementation language such as C/C++, Java, FORTRAN, Perl, Lisp, etc.
Chapter III. Blueprint of a General GA Package

In chapter 2, we reviewed several existing GA packages. Most packages use a single representation: either binary string or floating point; some packages are specially designed for certain types of problems. To handle the variety of optimization problems in practical applications, I will design and implement a general GA package that gives users much more options on the problem representation and genetic operators. In other words, this package offers the choice of representing the chromosome using bit-string, integer, or floating-point or a combination of the three. A preliminary version of this package, which only implements the simplest methods of crossover and mutation, was developed by Ting Zhu [26]. The package I propose will significantly extend the original package by providing users the choice of choosing from a rich set of selection, replacement, mutation, and crossover methods based on the specific problem type.

How to encode the chromosomes is the problem to solve when starting to work on a problem with GA. Encoding depends heavily on the problem. In this chapter, we will start with the introduction to some encodings that will be implemented in this package.

3.1 Encoding

The way in which candidate solutions are encoded is “a central factor in the success of a genetic algorithm” [1]. This package will employ three commonly used
encodings – bit-string, Gray-coded integer, and floating point, to represent candidate solutions.

3.1.1 Bit-string Encoding

Bit-string encoding, introduced by Holland in his original GA book, is the most widely used encoding method. Bit-string encoding has its own inherent advantages. First, it is easy to understand and implement; second, researchers have accumulated rather rich experiences on the parameter settings such as population size, crossover, and mutation rate in its context.

3.1.2 Integer Encoding

When the parameters of the problem are integers, it is natural to use a binary string to encode the variables. However, there is a problem with representing integers by ordinary binary strings. Consider the following example. Two integers 15 (01111) and 16 (10000) are selected as parents. Note that the integers are quite close to each other and their fitness values are also likely to be close. But, their binary string representations are quite different from each other. Moreover, let’s assume a single point crossover is applied after bit position 1. The resultant offspring are 11111 and 00000, whose corresponding integer values are 31 and 0, respectively. We can see that the offspring fall far away from their parents, which tends to interfere with the expected convergence [7].

One way to overcome this problem is to encode the binary string using Gray code. The procedures for converting a binary number $b=b_1b_2...b_m$ into a Gray code number $g=g_1g_2...g_m$ and vice versa are given in the following figure [2].
According to these conversion procedures, the Gray codes of the first 6 integers are listed in Table 3.1.

<table>
<thead>
<tr>
<th>Decimal value</th>
<th>Binary string</th>
<th>Gray code</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0000</td>
<td>0000</td>
</tr>
<tr>
<td>1</td>
<td>0001</td>
<td>0001</td>
</tr>
<tr>
<td>2</td>
<td>0010</td>
<td>0011</td>
</tr>
<tr>
<td>3</td>
<td>0011</td>
<td>0010</td>
</tr>
<tr>
<td>4</td>
<td>0100</td>
<td>0110</td>
</tr>
<tr>
<td>5</td>
<td>0101</td>
<td>0111</td>
</tr>
<tr>
<td>6</td>
<td>0110</td>
<td>0101</td>
</tr>
</tbody>
</table>

Table 3.1 Gray codes of first 6 integers

Note that the Gray coding representation has the property that any two consecutive points in the problem space differ by only one bit in the representation space, which is highly desired for GA convergence. Let's reconsider the previous example using Gray coding. The Gray codes of 15 and 16 are 01000 and 11000 respectively. After applying the same crossover procedure, the offspring are 11000 and 01000, whose integer values are 16 and 15 respectively. Thus we can observe that offspring are
much closer (actually identical in this extreme case) to their parents than in the previous case, which helps preserve the desired characteristics of the parents.

3.1.3 Floating-point Encoding

Despite the advantages it has, binary encoding has some drawbacks when applied to problems with continuous parameters. In such numerical problems that demand parameters of high precision, each variable requires a rather long binary string to fully represent it. This results in a huge search space which bogs down the GA performance severely\(^2\).

For applications with variables over continuous domains, it is more natural and logical to encode them using floating-point numbers instead of binary strings. In contrast to bit-string encodings, floating-point encoding represents the continuous parameters more accurately, requires less storage space, and makes it more straightforward to implement the genetic operators\(^7\).

3.2 Selection Methods

The next decision to make after encoding is selection – the process of selecting the individuals from the population as parents that will create offspring for the next generation. The aim of selection is to assign higher probabilities to individuals with greater fitness. There are two main issues associated with selection: population diversity and selective pressure\(^{16}\). A high selective pressure will result in only good individuals being selected, reducing the diversity needed for evolution toward the global optimum, thus tending to lead the GA to a premature convergence. On the other hand, a weak selective pressure will get too many not-so-good individuals
involved in the population and slow down the evolution process. Therefore, it is critical to seek a balance between these two factors.[16]

Numerous selection schemes have been developed over the years. Every method has its own strength and weakness; it is still an open question which method is superior to the others. In the following subsections, I will describe the selection methods that will be implemented in this package.

### 3.2.1 Improved Roulette Wheel Selection

Roulette Wheel Selection is the traditional selection method proposed in Holland’s original GA work: the expected value (the expected number of times of being selected) of an individual is proportional to its fitness. This method can be implemented using a roulette wheel in the following way. Each individual is assigned a slice of a circular roulette wheel, with the size of the slice being proportional to the individual’s fitness. The wheel is then spun \( N \) (population size) times. After each spin, the individual pointed by the wheel’s marker is selected. The procedures of this method are detailed in the following figure[2].

1. Calculate the fitness value \( f(i) \) for each chromosome \( v_i \)
2. Find the total fitness of the population
   \[
   F = \sum_{i=1}^{N} f(i)
   \]
3. Calculate each chromosome’s expected value \( E_i \)
   \[
   E_i = N \cdot f(i) / F
   \]
4. Calculate cumulative expected value for each chromosome
   \[
   S_j = \sum_{i=1}^{j} E_j
   \]
5. Generate a random number \( r \) of range \([0, N]\)
6. If \( r < S_{j-1} \), then select \( v_j \), otherwise select \( v_i \) such that \( S_{j-1} < r \leq S_j \)
7. Go to step 5 until \( N \) chromosomes have been selected.

**Figure 3.2 Roulette wheel selection procedures**
We can see that it takes $O(N)$ time in step 6 to locate the individual with the accumulative expected value that equals to $r$. If we notice that $S_i$ is a sorted array in ascending order, it is natural to come up with some ideas about binary search. Therefore, we replace step 6 with finding $S_i$ using binary search in the above algorithm. Thus the performance of the algorithm can be improved from $O(N)$ to $O(\log N)$.

In the case of a theoretically infinite population, the roulette wheel method will allocate the expected values of each chromosome in proportion to its fitness. However, if the real GA application has a relatively small population size, the actual number of an individual being chosen may be quite different from its expected value. In the worst case scenario, an extreme series of spin of the wheel can “allocate all the offspring to the worst individual in the population” [1]. Another restriction of roulette wheel method is that all fitness values of the objective function must be positive. The minimum value of the function is important: adding a constant to all the fitness values, a scaling technique that should be harmless, will change the expected values of individuals.

### 3.2.2 Stochastic Universal Sampling

To overcome the drawback of roulette wheel selection, James Baker proposed Stochastic Universal Sampling (SUS) [17]. SUS rebuilds the roulette wheel with $N$ equally spaced pointers. Thus unlike roulette wheel selection, SUS spins the wheel only once, and the individuals pointed by each of the $N$ markers are selected as parents. Below is an elegant C implementation of SUS [17].
Where \( S[i] \) is the accumulative probability of chromosome \( i \). Under this method, each individual is guaranteed to be selected at least \( \text{floor}(E_i) \) times but no more than \( \text{ceiling}(E_i) \) times, where \( \text{floor} \) and \( \text{ceiling} \) functions are the same as defined in C language.

SUS does a better job in sampling than roulette wheel, but does not solve the major problems with fitness proportionate selection. Under the fitness proportionate selection, a small number of highly fit individuals are much more likely to be selected and their descendants will quickly dominate the population. After a few generations, the super individuals may eliminate the desirable diversity and lead the GA to a premature convergence \(^1\).

### 3.2.3 Rank Selection

One way to address the problems associated with fitness proportionate selection methods is to introduce a mapping mechanism that maps the raw fitness values to intermediate parameters. These parameters are then used to calculate the expected values. Rank selection is a method in this category: the individuals are sorted according to their fitness values, and then the selection probabilities of individuals are calculated based on their ranks rather than on their actual fitness values. Doing so avoids allocating too much quota of the offspring to individuals with high fitness.
values. This also solves the problem associated with a scaling technique in roulette wheel and SUS selection.

Like everything else in the universe, the rank method also has drawbacks. In some cases, the GA takes a longer time to converge to an optimum point because of the lower selection pressure. Rank also requires sorting the entire population for each generation, which is a time consuming task in the case of a large population size [18].

There are many approaches to calculating the expected values based on ranking. Two ranking methods are employed in this GA package.

3.2.3.1 Baker Linear Rank Method [12]

The linear ranking method proposed by Baker is quite straightforward. In his method, each individual is ranked in ascendant order of fitness, from 1 to N. The expected value of each individual is set by the following formula,

\[
E(j) = Min + (Max - Min) \cdot \frac{Rank(j) - 1}{N - 1}
\]

where \( Min \) is the expected value of individual with rank 1, \( Max \) is the expected value of the individual with rank \( N \), and also is the upper bound of expected values defined by the user. Given the condition that \( \sum_j E(j) = N \), it is easy to show that \( Min = 2 \cdot Max \). Baker recommended the user to take \( Max = 1.1 \). Once the expected values are established, the SUS method is used to sample the population.
3.2.3.2 Reeves Rank Method \[18\]

Colin R. Reeves proposed another method of linear ranking, in which the probability of selecting the individual ranked \(k\) in the population is denoted by

\[ Pr[k] = a + bk \]

where \(a\) and \(b\) are positive scalars. The fact that \(Pr[k]\) must be a probability distribution gives us the following deduction:

\[
\sum_{k=1}^{N} (a + bk) = 1 \Rightarrow N(a + b \frac{N + 1}{2}) = 1
\]

In his work, Reeves gives a formal definition of the selection pressure:

\[
\phi = \frac{Pr[selecting fittest individual]}{Pr[selecting average individual]},\]

which is a parameter specified by the user.

If we interpret the average as the median individual, it is easy to get

\[
\phi = \frac{a + bN}{a + b(N + 1)/2}.
\]

In terms of \(\phi\) and \(N\), we are able to get

\[
b = \frac{2(\phi - 1)}{N(N - 1)} \quad \text{and} \quad a = \frac{2N - \phi(N + 1)}{N(N - 1)},
\]

which implies \(1 \leq \phi \leq 2\). It is easy to see that the cumulative probability of each individual can be expressed in terms of the sum of an arithmetic progression, so that finding the \(k\) for a given random number \(r\) is simply to solve the quadratic equation

\[
ak + b \frac{k(k + 1)}{2} = r \quad \text{for} \ k.
\]

Note that \(k\) is an integer, thus

\[
k = \text{ceiling}\left\{\frac{-(2a + b) + \sqrt{(2a + b)^2 + 8br}}{2b}\right\}.
\]

Note that the individual can be directly found by the above formula, which takes only \(O(1)\) time, whereas the traditional roulette wheel method takes \(O(N)\) time to find the
corresponding individual with a cumulative probability equal to $r$. For a large size of population, this will considerably improve the computational efficiency.

### 3.2.4 Tournament Selection

The tournament selection method is akin to the rank method, but is more time-efficient than rank selection because it works on a smaller set of individuals instead of working on the whole population. This method randomly picks a set of $m$ individuals from the population and then chooses the best one of them to be the parent. This process is repeated $N$ times to get the $N$ parents needed \[^2\]. Obviously, selection pressure rises as $m$ increases. The commonly used value of $m$ is 2.

Michalewicz proposed a tournament selection method which is blended with some simulated annealing flavor \[^2\]. Two individuals are randomly chosen, and then a random number $r$ in $[0, 1]$ is generated. The winner is determined in the following way: If \[ \frac{1}{1 + e^{-\frac{f(i) - f(j)}{T}}} < r, \] then the fitter individual is chosen; otherwise, the less fit individual is chosen. In the formula, $T$ is the temperature and $f(i)$ and $f(j)$ are the fitness values.

As in simulated annealing, the temperature starts at a high value, which indicates a low selection pressure. The temperature is gradually lowered according to a predefined scheme. The lower the temperature goes, the higher the selection pressure becomes, therefore allowing the GA to narrow down the search space on highly fit individuals.
3.3 Generation Replacement Model

The generation replacement model determines how the population proceeds from one generation to the next generation. In this package, several choices are provided as described in the following subsections.

3.3.1 Generational Replacement

This is the classical replacement method presented in Holland’s original GA work – at each generation, $N$ offspring are generated to make up the new generation. In other words, the entire generation is replaced; no individual from the last generation is kept.

3.3.2 Steady State Replacement

The potential problem of the generational replacement is that we are running the risk of discarding some good individuals in the old generation. To address this problem, De Jong introduced the idea of a ‘generation gap’ in his Ph.D. thesis [19]. At each generation, only a fraction of the population is replaced by the offspring. This fraction is called the generation gap. Therefore overlap is allowed between successive generations.

There are a number of mechanisms to select the parents that are to be replaced. In this package, I use the classical method: the least fit individuals in the old generation are chosen to be replaced by the newly generated offspring.

3.3.3 Evolution Strategy ($N+m$) Replacement [2]

Unlike the previous two replacement schemes, in which the children replace their parents, in this evolution scheme, the children compete with their parents for survival.
In a single generation, \( m \) offspring are generated and then are combined with the \( N \) parent to constitute an intermediate population of size \( N+m \), from which the best \( N \) individuals are chosen to form the next generation.

### 3.3.4 Elitism \(^2\)

It is possible that some individuals in the earlier generation are fitter than the best individual found in the last generation. Such individuals may be lost if they are not selected or are ruined by crossover or mutation. Thus it is intuitive to keep track of the best individuals in the entire evolution process and this is called elitism. Elitism ensures that the best \( n_{Elite} \) individuals of the population are passed on to the next generation without being altered by genetic operators. Elitism is essentially a special case of \((N+m)\) replacement.

### 3.4 Crossover

After parents have been selected through one of the methods introduced above, they are randomly paired. The genetic operators are then applied on these paired parents to produce offspring. There are two main categories of operators: crossover and mutation. Let us have a look at crossover in this section and move on to mutation in the next section.

The purpose of crossover is to vary the individual quality by combining the desired characteristics from two parents. Over the years, numerous variants of crossover have been developed in the GA literature, and comparisons also have been made among these methods \(^{20}\). However, most of these studies rely on a small set of test problems, and thus it is hard to draw a general conclusion on which method is better than others.
In this GA package, a number of commonly used crossover techniques are implemented. Some of them apply only to bit-string or Gray-coded integer genes on the chromosome.

### 3.4.1 Single Point Crossover

This is the traditional and the simplest way of crossover: a position is randomly chosen as the crossover point and the segment of the parents after the point are exchanged to form two offspring. The position can be anywhere between the first and the last allele. For example, let us suppose position 2 is the crossover point, the process is shown in the following table. Please note that \( x_i \) or \( y_i \) stands for a bit for bit-string chromosomes; whereas for floating point chromosome, it stands for a floating point variable.

<table>
<thead>
<tr>
<th>Parent</th>
<th>Offspring</th>
</tr>
</thead>
<tbody>
<tr>
<td>([x_1, x_2, x_3, x_4, x_5, x_6])</td>
<td>([x_1, x_2, y_3, y_4, y_5, y_6])</td>
</tr>
<tr>
<td>([y_1, y_2, y_3, y_4, y_5, y_6])</td>
<td>([y_1, y_2, x_3, x_4, x_5, x_6])</td>
</tr>
</tbody>
</table>

Table 3.2 Single point crossover

Single point crossover has several drawbacks. One of them is the so-called positional bias: the single point combination of this method can’t create certain schemas \([20]\). The other is the endpoint effect – the end point of the two parents is always exchanged, which is unfair to other points \([1]\).

### 3.4.2 Double Point Crossover

In this crossover method, two positions are randomly chosen and the segments between them are exchanged. For instance, assuming position 2 and 5 are chosen as crossover points, the process is shown in the following table.
Compared with single point crossover, double point crossover is able to combine more schemas and eliminate the ending point effect. But similar to single point crossover, there are some schemas that it can’t create.

3.4.3 Uniform Crossover

As a natural extension to single and double point crossover, uniform crossover allows each allele in the parents to be exchanged with a certain probability \( p \). For example, given \( p=0.6 \), and the series of random numbers drawn for alleles 1 to 6 are: 0.4, 0.7, 0.3, 0.8, 0.9, 0.5, the results will be:

<table>
<thead>
<tr>
<th>Parent</th>
<th>Offspring</th>
</tr>
</thead>
<tbody>
<tr>
<td>([x_1, x_2, x_3, x_4, x_5, x_6])</td>
<td>([x_1, y_2, x_3, y_4, x_5, x_6])</td>
</tr>
<tr>
<td>([y_1, y_2, y_3, y_4, y_5, y_6])</td>
<td>([y_1, y_2, x_3, x_4, y_5, y_6])</td>
</tr>
</tbody>
</table>

Since any point in the parent can be exchanged, uniform crossover can recombine any schemas. However, this advantage comes at cost. The random bit exchange may prevent the forming of valuable building blocks (short and highly fit segments that are able to form strings of potentially higher fitness through recombination) \(^3\), because the random exchange of uniform crossover can be highly disruptive to any evolution schemas \(^{21}\).

3.4.4 Arithmetic Crossover (floating point only)

The above crossover methods work fine for the bit-string representation. However, there is a problem when applying them to floating-point encoding: the crossed
parameter values in the parents are propagated to their offspring unchanged, only in different combinations \cite{7}. In other words, no new parameter values are introduced.

To address this problem, GA researchers introduced arithmetical crossover \cite{7}, the \textit{ith} parameter in offspring, $Z_i$, is a linear combination of the \textit{ith} parameters of two parents, $X_i$ and $Y_i$, i.e.,

$$Z_i = aX_i + (1-a)Y_i$$

where $a$ is a random value in the interval [0,1].

This linear formula guarantees that the value of the newly created offspring is always within the domain of the problem. For example, suppose the crossover happens at position 2 and 5, the resultant offspring are given in the table below:

<table>
<thead>
<tr>
<th>Parent</th>
<th>Offspring</th>
</tr>
</thead>
<tbody>
<tr>
<td>$[x_1, x_2, x_3, x_4, x_5, x_6]$</td>
<td>$[x_1, ax_2+(1-a)y_2, x_3, x_4, ax_5+(1-a)y_5, x_6]$</td>
</tr>
<tr>
<td>$[y_1, y_2, y_3, y_4, y_5, y_6]$</td>
<td>$[y_1, ay_2+(1-a)x_2, y_3, y_4, ay_5+(1-a)x_5, y_6]$</td>
</tr>
</tbody>
</table>

Table 3.5 Arithmetical crossover

3.4.5 Heuristic Crossover (floating point only)

Heuristic crossover \cite{22} uses the values of the fitness function to determine the direction of search according to the following formula:

$$Z = \begin{cases} 
    a(Y - X) + Y & \text{if } f(Y) \leq f(X) \\
    a(X - Y) + X & \text{otherwise} 
\end{cases}$$

where $a$ is also a random value in the interval [0,1]; $X$ and $Y$ are the parental parameter vectors; $f(X)$ and $f(Y)$ are the fitness function values.

Contrary to arithmetic crossover which always creates offspring within the range of their parents, heuristic crossover always generates offspring outside of the range of
the parents. Sometimes the offspring values are so far away from their parents that they get beyond the allowed range of that variable. In such a case, the offspring is discarded, and a new \( a \) is tried and a new offspring created. If after a specified number times no feasible offspring is generated, the operator quits with no offspring produced.

In this package, instead of being discarded, the infeasible offspring is assigned the maximum or minimum permissible value as follows:

\[
z_i = \begin{cases} 
\max(iMin, \min(a(y_i - x_i) + y_i, iMax)) & \text{if } f(Y) \leq f(X) \\
\max(iMin, \min(a(x_i - y_i) + x_i, iMax)) & \text{otherwise}
\end{cases}
\]

where \( iMin \) and \( iMax \) are the minimum and maximum allowable values for the \( ith \) parameter.

### 3.5 Mutation

Mutation is another essential operator of genetic algorithm. The idea behind mutation is to introduce diversity into the population and thus prevent the premature convergence. Like crossover, GA researchers have developed numerous variants of mutation method over the GA history. Since every method has its strength and weakness, it is still an open question as to claim which method is the best. This general GA package employs a number of widely used mutation procedures.

#### 3.5.1 Uniform Mutation

This is the conventional method of mutation, in which each allele has an equal opportunity to be mutated. For each allele, a random number \( r \) is generated and then
compared with the mutation rate $p_m$ (a user specified parameter), if $r > p_m$, then this allele is mutated; otherwise, stays unchanged.

For the binary encoding, either bit-string or Gray-coded integer, the mutation simply flips the bit from 0 to 1 or vice versa. For floating-point encoding, the allele is replaced with a random value within the domain of that variable. For example, if $x_2$ is selected for mutation in the chromosome $X = [3.221, 4.556, 2.341, 5.897]$, and the domain of $x_2$ is $[3.000, 5.500]$, then the chromosome after mutation may be $X = [3.221, 3.938, 2.341, 5.897]$.

### 3.5.2 Non-uniform Mutation (floating point only)

To increase the fine tuning of the chromosome and reduce the randomness of uniform mutation, Michalewicz presented a dynamic mutation operator called non-uniform mutation $^{[2]}$. The mechanism works as follows: if element $x_i$ was selected for mutation, the resultant offspring is determined by the following formula,

$$z_i = \begin{cases} x_i + \Delta(t, UB_i - x_i) & \text{if } a > 0.5 \\ x_i - \Delta(t, x_i - LB_i) & \text{if } a \leq 0.5 \end{cases}$$

where $a$ is a random number between 0 and 1, $LB_i$ and $UB_i$ represent lower and upper bounds of variable $x_i$. The function $\Delta(t, x)$ is defined by

$$\Delta(t, x) = x(1 - r^{(t/T)^b})$$

where $r$ is also a random number between 0 and 1, $T$ is the maximal generation number, and $b$ is a user-defined system parameter that determines the degree of dependency on iteration number $t$. This simulated annealing flavored formula returns a value in the range $[0, x]$ such that the probability of $\Delta(t, x)$ being close to 0 increases as $t$ increases. This property guarantees the newly created offspring is in the
feasible range. Moreover, it enables the operator to explore the space broadly with an initially large mutation step, but very locally at later stages; thus make the newly generated offspring closer to its predecessor than a randomly generated one \([2]\).

### 3.5.3 Boundary Mutation (floating point only)

Boundary Mutation is a variant of uniform mutation. In uniform mutation, the newly generated allele is a random value within the domain of that allele. In boundary mutation, however, the newly generated allele is either the upper bound or the lower bound of the domain, with equal probability \([2]\), i.e.,

\[
 z_i = \begin{cases} 
 UB & \text{if } a > 0.5 \\
 LB & \text{if } a \leq 0.5
\end{cases}
\]

where the parameters have the same meaning as in non-uniform mutation.

This mutation operator is extremely functional in optimizing problems where the optimal solution either lies on or near the boundary of the constrained search space. In addition, the conjunction of this operator with arithmetic crossover will help to counteract the later operator’s “contraction effect” on the population \([14]\). However, this operator has obvious disadvantages for many problems.

### 3.5.4 Creep Mutation (floating point only)

Davis introduced the creep mutation operator in his GA handbook \([23]\), in which the allele values which are to be mutated are incremented or decremented by the creep fraction – the fraction of the valid range for that variable’s value. The selected variable \(x_i\) is mutated according to the following formula:

\[
 z_i = \min( iMax, \max( iMin, \ x_i + rs(iMax- iMin)) )
\]
where \( iMax \) and \( iMin \) are the upper bound and lower bound of the variable’s domain respectively; \( r \) is a random number in the range \([-1, 1]\); \( s \) is the creep factor, a user-defined number between 0 and 1.

It is equally likely that the allele value will be incremented or decremented, and the value is adjusted to the boundary value if the newly generated allele falls out of the valid range of the allele.

### 3.6 Termination Condition

Unlike the traditional search methods that terminates when a local optimum is reached, GAs are stochastic methods that could run forever. The termination criterion plays an important role in the solution quality. Three commonly used approaches are adopted in this package.

#### 3.6.1 Maximum Number of Generations

The simplest and most common method is to check the current generation number; the genetic process will end when the specified number of generations has been reached. This method has several variants such as using a maximum number of fitness function evaluations, or using a maximum elapsed time, as the termination criterion\(^1\).

#### 3.6.2 Convergence Termination

The drawback of the above termination criterion and its variants is that they assume the user’s knowledge of the characteristics of the problems to be solved. In many occasions, however, it is quite difficult to specify the number of generations (or
evaluation or elapse time) without prior experience on the problem. Thus it is much better if the evolution process stalls when the chance for a significant improvement is fairly slim \cite{18}.

There are mainly two categories of termination criteria to tell the chances of improvement of the algorithm: chromosome structure diversity, and fitness improvement \cite{18}. Convergence termination belongs to the first category. It makes the termination decision by checking the number of converged alleles. If a predefined percentage of the chromosomes of the whole population have the same or similar values for this allele, the algorithm stops.

### 3.6.3 Progress in Fitness

As an instance of the second category mentioned above, this approach measures the progress made by the algorithm in a predefined number of generations. If there is no change, or if the improvement on the best chromosome is smaller than a user specified factor, the genetic process ends.
Chapter IV. Implementation of the GA Package

In this chapter, we’ll implement the GA package based on the population model and operators described in the previous chapter. The package is written in Fortran 77, a simple but powerful and highly efficient programming language. The program can mainly be divided into four function modules: Initialization, Genetic Operations, Output, and Utility. We begin with the introduction to the main modules.

4.1 Introduction to Modules

The package is designed with modular programming principles in mind. By partitioning associated functionalities into relatively independent modules, the package is made easier to understand, maintain and upgrade.

4.1.1 Initialization Module

This module reads from the input file the GA parameters such as population size, mutation rate, crossover rate, etc; and randomly populates the initial generation. The functionality of this module is implemented through two Fortran files: Inputf.for and Init.for.

Inputf.for contains the following subroutines:

- InputF: The main subroutine of the unit, which reads the GA configuration information from the input file by calling other auxiliary subroutines or functions.
- **rdLine**: This subroutine reads one line from the input file and extract a real number or a string from the line.
- **rdInt**: Reads an integer from the input file.
- **rdRgeF**: Reads the ranges of floating point variables.
- **rdRgeI**: Reads the ranges of integer variables.

**Init.for** is made up of the following subroutines:

- **Init**: The main subroutine of the unit, which calls subsidiary subroutines based on the type of encoding and the types of variables.
- **InitB**: This subroutine randomly initializes population in the form of bit strings.
- **InitI**: This subroutine randomly initializes the integer variables of the chromosomes.
- **InitF**: This subroutine randomly initializes the floating point variables of the chromosomes.

### 4.1.2 Genetic Operation Module

This module is the core algorithm of the package, which implements the key GA operations including selection, crossover, mutation, replacement, etc. This module comprises five Fortran source files, which are listed below.

**Eval.for** includes two subroutines:

- **Eval**: The main subroutine of the unit, which evaluates the fitness of each chromosome by calling user-defined function and records the best chromosome in the current generation.
• **usrFun**: The user-defined function, which is used to calculate the fitness value for each chromosome. Users may modify this function to optimize different problems.

Select.for comprises the following subroutines:

• **Select**: The main subroutine of the unit, which selects parents for the next generation based on various selection methods.

• **Roulet**: This subroutine performs the roulette wheel selection.

• **SUS**: This subroutine performs the stochastic uniform sampling selection.

• **Baker**: This subroutine calculates the expected value based on Baker linear method.

• **Reeves**: This subroutine performs the Reeves Rank linear selection.

• **Tourna**: This subroutine performs the tournament selection.

• **ToNewB**: This subroutine copies the selected chromosomes (bit-string encoding) into the new population.

• **ToNewD**: This subroutine copies the selected chromosomes (integer/floating-point variables) into the new population.

Cross.for contains the following subroutines:

• **Cross**: The main subroutine of the unit, which recombines the selected parent chromosomes to generate offspring based on different crossover methods.

• **ScrosB**: This subroutine performs single-point crossover on a pair of selected parent chromosomes (bit-string encoding).

• **ScrosD**: This subroutine performs single-point crossover on a pair of selected parent chromosomes (integer/floating-point encoding).

• **DcrosB**: This subroutine performs double-point crossover on a pair of selected parent chromosomes (bit-string encoding).
• DcrosD: This subroutine performs double-point crossover on a pair of selected parent chromosomes (integer/floating-point encoding).

• UcrosB: This subroutine performs uniform crossover on a pair of selected parent chromosomes (bit-string encoding).

• UcrosD: This subroutine performs uniform crossover on a pair of selected parent chromosomes (integer/floating-point encoding).

• Across: This subroutine performs arithmetic crossover on a pair of selected parent chromosomes (integer/floating-point encoding).

• Hcross: This subroutine performs heuristic crossover on a pair of selected parent chromosomes (integer/floating-point encoding).

Mutate.for contains the following subroutines:

• Mutate: The main subroutine of the unit, which mutates the offspring according to various mutation methods.

• MuteB: This subroutine performs uniform mutation on the selected chromosome (bit-string encoding).

• UmuteI: This subroutine performs uniform mutation on the integer variables of the selected chromosome.

• UmuteF: This subroutine performs uniform mutation on the floating point variables of the selected chromosome.

• DmuteI: This subroutine performs dynamic (non-uniform) mutation on the integer variables of the selected chromosome.

• DmuteF: This subroutine performs dynamic (non-uniform) mutation on the floating point variables of the selected chromosome.
• CmuteI: This subroutine performs creep mutation on the integer variables of the selected chromosome.

• CmuteF: This subroutine performs creep mutation on the floating point variables of the selected chromosome.

• BmuteI: This subroutine performs boundary mutation on the integer variables of the selected chromosome.

• BmuteF: This subroutine performs the boundary mutation on the floating point variables of the selected chromosome.

Replace.for includes the following subroutines:

• Replac: The main subroutine of the unit, which replaces parents with the newly generated offspring based on various replacement methods.

• GReplB: This subroutine replaces the whole generation with the newly generated offspring (bit-string encoding).

• GReplD: This subroutine replaces the whole generation with the newly generated offspring (integer/floating-point encoding).

• SReplB: This subroutine performs the steady state replacement on the generation (bit-string encoding).

• SReplD: This subroutine performs the steady state replacement on the generation (integer/floating-point encoding).

• EReplB: This subroutine performs the evolutionary strategy replacement on the generation (bit-string encoding).

• EReplD: This subroutine performs the evolutionary strategy replacement on the generation (integer/floating-point encoding).
Terminate.for comprises the following subroutines:

- **Termin**: The main subroutine of the unit, which stops the GA by different termination criteria.
- **tNGen**: This function terminates the GA by the number of generations the GA is supposed to run.
- **tAlleB**: This function terminates the GA by the percentage of converged alleles in the current generation (bit-string encoding).
- **tAlleD**: This function terminates the GA by the percentage of converged alleles in the current generation (integer/floating-point encoding).
- **tFitP**: This function terminates the GA by fitness progress made in a certain number of generations.

### 4.1.3 Output Module

The output module writes the results of the GA package, such as the optimal chromosome, the optimal fitness, the corresponding generation number, to the output file. The functionality of this module is implemented through a single Fortran file—Output.for, which is composed of one subroutine. The user may modify this subroutine to customize the output information.

### 4.1.4 Utility Module

This module groups the utility functions such as search, random, Gray/DeGray, bit-string operation, sort, etc. Doing so prevents the main GA program unit from being filled up with utility codes, and thus makes the main program clean and easy to follow.
The functionality of utility module is implemented in a single Fortran file—Util.for. This file contains quite a number of subroutines and functions. Some of the key units are listed below:

- binSch: This function does a binary search on a sorted array and returns the index of the element found.
- linSch: This function does a linear search on a regular array and returns the index of element found.
- sort: This subroutine sorts an array into descending order.
- schRnk: This function returns the rank of a chromosome for use in some rank selection methods.
- random: This function returns a randomly generated number between 0.0 and 1.0.
- delta: A simulated annealing flavored function used in dynamic mutation.
- bitLen: This function returns the length of the binary string required to represent an integer/floating-point variable.
- binToD: This subroutine converts the bit strings representation of the current generation into the integer/floating-point representation.
- btsToI: This function converts a single bit-string to an integer.
- Gray: This subroutine converts binary integers into the Gray-coded integers.
- deGray: This subroutine reverses the Gray-coded integers into binary integers.
- doXOR: This function does an exclusive OR on two bits.

4.2 Program Structure

The modules described in the above sections are linked together by the main program—GA, which performs the genetic operations by calling the subroutines implemented in separate files.
The flow chart below outlines how the program runs:

<table>
<thead>
<tr>
<th>program GA</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. <strong>InputF</strong> - Reads the GA configuration information from the input file.</td>
</tr>
<tr>
<td>2. <strong>Init</strong> - Randomly initialize the population within domains of the problem.</td>
</tr>
<tr>
<td>3. <strong>Eval</strong> - Evaluate the fitness of each chromosome in the 1\textsuperscript{st} generation.</td>
</tr>
<tr>
<td>4. <strong>while</strong> (termination condition not met)</td>
</tr>
<tr>
<td>Select - Select parents based on various selection methods.</td>
</tr>
<tr>
<td>Cross - Recombine the parents based on various crossover methods.</td>
</tr>
<tr>
<td>Mutate - Mutate the offspring based on various mutation methods.</td>
</tr>
<tr>
<td>Replace - Replace the generation based on various selection methods.</td>
</tr>
<tr>
<td>Eval - Evaluate the fitness of each chromosome of the current generation.</td>
</tr>
<tr>
<td>5. <strong>Output</strong> - Write the result to output file.</td>
</tr>
<tr>
<td><strong>End</strong></td>
</tr>
</tbody>
</table>

![Program flow chart](image)

**4.3 Data Structures**

The major population data are stored in the following data structures:

- `bitStr(i, j, k)`: This three-dimensional array stores the bit-string representation of the population, where `i` stands for the chromosome index, `j` stands for the string index, and `j` stands for the bit index.

- `ints(i, j)`: This two-dimensional array stores the integer variables in the chromosome, where `i` stands for the chromosome index, `j` stands for the index of an integer variable.
4.4 A Sample Run

In this section, we will go through an example to show how the package works. The function to be optimized is one of the Rosenbrock functions:

\[ f(x_1, x_2) = 100(x_1^2 - x_2)^2 + (1-x_1)^2, \quad -3 \leq x_1 \leq 3, \quad -2.0 \leq x_2 \leq 2.0, \]

where \( x_1 \) is an integer and \( x_2 \) is a floating point variable.
Step 1: Input file

First, we must set up the GA configuration parameters in an input file; the name of the file could be arbitrary, let’s name it ‘ga.inp’ in our case. Below is the format and content of this input file:

```
##############################Start of input file#################################
Population Size: 100
Crossover Rate: 0.4
Mutation Rate: 0.3
###
Seed for Random Function: 10000
###
Encoding: Mix
###
Selection Method: SUS
Crossover Method:
Crossover Method(Non-FP): Double Point
Crossover Method(FP): Double Point
Mutation Method(Non-FP): Uniform
Mutation Method(FP): Uniform
Replacement Method: Generational
###
Termination Criterion: Number of Generations
Number of Generations: 200
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
###Variables setting
Number of Bit-string Variables: 0
#String Var1 length: 12
Number of Floating-point Variables: 1
  FP Var1 Range: -2.0, 2.0
Number of Integer Variables: 1
  Int Var1 Range: -3, 3
###
##############################End of input file#################################
```

We can see that most of the settings in the input file are self-explanatory. Please note that lines starting with '#' are comments. The user may add his or her own comments to make the input file even more readable.

We used ‘Mix’ in the encoding type, which means the problem has both integers and floating point variables and they are encoded in integer/floating-point format. On the
other hand, if the encoding type is set to ‘Bit String’, then all the variables will be encoded in the form of bit strings.

The package provides plenty of options the user can choose. They are listed as follows:

- Selection methods include: Roulette Wheel, SUS, Baker Rank, Reeves Rank, and Tournament.
- Crossover methods include: Single-point, Double-point, Uniform, Arithmetic, and Heuristic.
- Mutation methods include: Uniform, Dynamic (Non-uniform), Boundary, and Creep.
- Replacement method include: Generational, Steady State, and Evolution Strategy.
- Termination criteria include: Number of Generations, Percentage of Converged Alleles, Fitness Progress, and Hybrid. Hybrid uses all the three termination criteria, and the GA stops if any of them is satisfied.

The function setting section specifies the number of integer and floating point variables and their ranges. If the problem has only one type of variables, then the number of variables of the unenclosed type should be set to zero.

**Step 2: User function**

After the setup of the input file, the next step is to specify the function to be optimized. To do this, we need to customize the ‘usrFun’ function in file ‘Eval.for’.
Step 3: Compile and run

After the specification of the user function, compile the package by issuing the following command on UNIX platform:

```
g77 GA.for -o GA.out
```

To run the program, type in

```
g77 GA.out
```

The system will prompt you for names of the input file and the output file. Enter them, and the program will then start running and the result will be written into the output file you specified.

Step 4: Output

The output file records the result of the program, which includes the optimal chromosome, its fitness value, and its generation number. The user may modify the
subroutine ‘Output’ to write other information into the output file. The output file for our sample run is shown below:

<table>
<thead>
<tr>
<th>Optimal fitness: 0.00000590</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optimal chromosome: 1, 0.99975700</td>
</tr>
<tr>
<td>Generation number: 197</td>
</tr>
</tbody>
</table>

Figure 4.4 Output file

Our sample Rosenbrock function has the minimum value 0.0 at (1, 1.0). Obviously, the package found a solution that is very close to the theoretical optimum. While working with other problems, if the user is not satisfied with the results, he/she can always adjust the GA parameters such as population size, number of generations, another crossover method, etc, to get a better result.
Chapter V. Tests and Results

In this chapter, we will test the GA package using various testing functions. The tests will be performed in two parts. In the first part, a variety of optimization problems will be input to the package and the output from the package will be compared with the theoretical optimal solution. In the second part, we will test certain problems by varying the selection, crossover, or mutation methods in order to compare the performances of different genetic operation methods. To save space and highlight the test results, the input files of all the test cases are not included here but are placed in the appendices of the thesis.

5.1 General Testing

In this section, experiments will be conducted to test the robustness of this GA package on a wide range of optimization problems. Let’s start with some classical testing problems – Rosenbrock functions.

5.1.1 Mixed Mode Rosenbrock Functions

Rosenbrock functions have several variants, one of which, the Standard Rosenbrock function, has been used as a sample in the previous chapter. In this subsection, we will test the other two forms: Flat-ground and Hollow-ground Rosenbrock functions.

Test 1. Flat-ground Rosenbrock Function

\[ f(x_1, x_2) = 100|x_1^2 - x_2| + (x_1 - 1)^2, \quad -3 \leq x_1 \leq 3, \quad -2.5 \leq x_2 \leq 2.5, \]
where $x_1$ is an integer and $x_2$ is a floating point number. This function has a minimum value 0 at $(1, 1.0)$. It is called flat-ground because a cross-section parallel to the $x_2$ axis shows a V shape, like a flat-ground knife blade.

The results of a single run are shown in the following table:

<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Best Fitness</th>
<th>Best Chromosome</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>0.07821111</td>
<td>1, 0.97203375</td>
</tr>
<tr>
<td>200</td>
<td>0.07821111</td>
<td>1, 0.97203375</td>
</tr>
<tr>
<td>400</td>
<td>0.00384010</td>
<td>1, 1.00619685</td>
</tr>
<tr>
<td>600</td>
<td>0.00384010</td>
<td>1, 1.00619685</td>
</tr>
<tr>
<td>800</td>
<td>0.00007044</td>
<td>1, 1.00083928</td>
</tr>
<tr>
<td>1000</td>
<td>0.00000016</td>
<td>1, 1.00003996</td>
</tr>
</tbody>
</table>

Table 5.1 Results of Flat-ground Rosenbrock function

It is easy to observe from the result table that, as the number generations grows, the GA converges to a point near the optimal point of the flat-ground Rosenbrock function. At generation 1000, the output result is already quite close to the theoretical optimal value.

However, it is worth mentioning that since the results come from a single run of the GA, there are some flat points existing between generations. For example, generation 100 and 200 have the same best fitness, so do generation 400 and 600. We will see in the next experiment that this phenomenon can be eliminated by using the average results of a number of runs.

Test 2. Hollow-ground Rosenbrock function

$$f(x_1, x_2) = 100 \sqrt{|x_2 - x_1^2| + (x_1 - 1)^2}, \quad -3 \leq x_1 \leq 3, \quad -2.5 \leq x_2 \leq 2.5,$$
where $x_1$ is an integer and $x_2$ is a floating point number. This function also has a minimum value 0 at (1, 1.0). It is called hollow-ground because a cross-section parallel to the $x_2$ axis shows a square root cusp, like a hollow-ground knife blade.

The following table shows the average results of 20 runs:

<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Average Best Fitness</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>1.35859689</td>
</tr>
<tr>
<td>200</td>
<td>1.06156089</td>
</tr>
<tr>
<td>400</td>
<td>0.74671261</td>
</tr>
<tr>
<td>600</td>
<td>0.64429165</td>
</tr>
<tr>
<td>800</td>
<td>0.45186317</td>
</tr>
<tr>
<td>1000</td>
<td>0.39576666</td>
</tr>
</tbody>
</table>

Table 5.2 Results of Hollow-ground Rosenbrock function

Mathematically speaking, the hollow-ground function is more difficult to optimize than flat-ground, because its partial derivatives have infinite discontinuities rather than finite discontinuities as in the flat-ground function. However, the GA package works remarkably well on this function. The GA converges as the number of generations grows; and at generation 1000, it presents us a solution close to the theoretical optimal value.

Moreover, we can see that the flat point phenomenon has been eliminated by using the average results of a number of runs instead of a single run. That is, the average fitness value decreases steadily as the number of generations increases.

### 5.1.2 0/1 Knapsack Problem

An important combinatorial optimization problem is the 0/1 knapsack problem, the task of this problem is to find a binary vector $v=b_1b_2…b_n$ such that
\[ P = \sum_{i=1}^{n} b_i P[i] \] is maximized while the constraint \( \sum_{i=1}^{n} b_j W[i] \leq C \) is satisfied,

where \( W[i], P[i] \) are the given sets of item weights and profits respectively; \( C \) is the capacity of the sack, and is defined as half of the sum of the item weights; \( n \) is the total number of items.

Various heuristic and exact methods have been devised to solve this problem. Among these, genetic algorithms have emerged as a powerful new search paradigms. The solution of this problem can be represented using bit string encoding: 1 indicates the corresponding item is selected, 0 not selected. For instance, the bit string 100101 means that item 1, 4, and 6 are selected into the sack.

**Experiment 1. Knapsack Problem with 10 Items**

In this experiment, we test the problem with 10 items. The item weights and profits are shown in the table below:

<table>
<thead>
<tr>
<th>Item</th>
<th>Weight</th>
<th>Profit</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>100</td>
<td>40</td>
</tr>
<tr>
<td>2</td>
<td>50</td>
<td>35</td>
</tr>
<tr>
<td>3</td>
<td>45</td>
<td>18</td>
</tr>
<tr>
<td>4</td>
<td>20</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>6</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>7</td>
<td>1</td>
<td>50</td>
</tr>
<tr>
<td>8</td>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td>9</td>
<td>10</td>
<td>49</td>
</tr>
<tr>
<td>10</td>
<td>8</td>
<td>19</td>
</tr>
</tbody>
</table>

Table 5.3 Weights and Profits of Knapsack problem

The results of this experiment are shown in the following table in next page:
<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Best Fitness</th>
<th>Best Chromosome</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>191</td>
<td>0 1 1 0 0 0 1 1 1 1</td>
</tr>
<tr>
<td>100</td>
<td>191</td>
<td>0 1 1 0 0 0 1 1 1 1</td>
</tr>
<tr>
<td>150</td>
<td>193</td>
<td>0 1 1 0 0 1 1 1 1 1</td>
</tr>
<tr>
<td>200</td>
<td>193</td>
<td>0 1 1 0 0 1 1 1 1 1</td>
</tr>
<tr>
<td>300</td>
<td>193</td>
<td>0 1 1 0 0 1 1 1 1 1</td>
</tr>
<tr>
<td>500</td>
<td>193</td>
<td>0 1 1 0 0 1 1 1 1 1</td>
</tr>
</tbody>
</table>

Table 5.4 Results of Knapsack problem with 10 items

It took the GA only 150 generations to catch the maximum profit value, 193, for this knapsack. In the next experiment, we are going to test a much harder knapsack problem with 100 items.

**Experiment 2. Knapsack Problem with 100 Items**

In this experiment, we test the problem with 100 items. We intentionally built the item weights and profits such that the first 50 items ought to be put into the sack. By this way, we are able to know what the correct solution is. GA reads the item weights and profits from an input file named knapsack.txt, which is too lengthy to be listed here and thus is attached in the appendices of this thesis.

The results of this experiment are shown in the following table:

<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Best Fitness</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>3325</td>
</tr>
<tr>
<td>1000</td>
<td>3556</td>
</tr>
<tr>
<td>1500</td>
<td>3608</td>
</tr>
<tr>
<td>2000</td>
<td>3701</td>
</tr>
<tr>
<td>2500</td>
<td>3701</td>
</tr>
<tr>
<td>3000</td>
<td>3701</td>
</tr>
<tr>
<td>3500</td>
<td>3759</td>
</tr>
</tbody>
</table>

Table 5.5 Results of Knapsack problem with 100 items

Even though the large number of items makes this problem much more difficult to optimize, the GA still successfully discovers the maximum profit value, 3759, which is the exact solution for this knapsack.
5.1.3 Floating-point Functions

In general, floating point problems are harder to optimize than integer problems. Thus, in this subsection we are going to test the more challenging problems with pure floating-point variables. It is not a surprise to observe that these problems usually require many more generations to obtain satisfactory results.

**Test1. Bohachevsky Function**

The Bohachevsky function \(^2\) is defined as follows:

\[
f(x_1, x_2) = x_1^2 + 2x_2^2 - 0.3 \cos(3\pi x_1) - 0.4 \cos(4\pi x_2) + 0.7
\]

where \(-5.0 \leq x_1 \leq 5.0\), \(-5.0 \leq x_2 \leq 5.0\), and \(x_1\) and \(x_2\) are both floating-point numbers. This function has a minimum value of 0 at (0.0, 0.0).

The average results of 20 runs are shown in the following table:

<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Average Best Fitness</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>0.10841466</td>
</tr>
<tr>
<td>1000</td>
<td>0.01410626</td>
</tr>
<tr>
<td>1500</td>
<td>0.00421070</td>
</tr>
<tr>
<td>2000</td>
<td>0.00072386</td>
</tr>
<tr>
<td>2500</td>
<td>0.00014686</td>
</tr>
<tr>
<td>3000</td>
<td>0.00006754</td>
</tr>
</tbody>
</table>

Table 5.6 Results of Bohachevsky function

The GA package performs well on this tough problem. After 3000 iterations, it found the solution value 0.00006754, which is very close to the optimal value 0.

**Test2. Schaffer Function F7**

The Schaffer function F7 \(^2\) is defined as follows:

\[
f(x_1, x_2) = (x_1^2 + x_2^2)^0.25 \sin^2(50(x_1^2 + x_2^2)^{0.1}) + 1.0
\]

where \(-5.0 \leq x_1 \leq 5.0\), \(-4.0 \leq x_2 \leq 4.0\), and \(x_1\) and \(x_2\) are both floating-point numbers. This function also has a minimum value of 0 at (0.0, 0.0).
The average results of 20 runs of this experiment are shown in the following table:

<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Average Best Fitness</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>0.13019798</td>
</tr>
<tr>
<td>1000</td>
<td>0.10833847</td>
</tr>
<tr>
<td>1500</td>
<td>0.10169734</td>
</tr>
<tr>
<td>2000</td>
<td>0.09560061</td>
</tr>
<tr>
<td>2500</td>
<td>0.09227524</td>
</tr>
<tr>
<td>3000</td>
<td>0.07749864</td>
</tr>
<tr>
<td>3500</td>
<td>0.07537527</td>
</tr>
</tbody>
</table>

Table 5.7 Results of Schaffer function F7

Clearly, this problem is tougher than Bohachevsky function, because after 3500 iterations, the package has not yet found a solution that is as good as the one found in the Bohachevsky function test. Nonetheless, the solution found is still rather satisfactory. Moreover, it is easy to see the trend from the table that the fitness value keeps getting better as the generation number increases.

5.2 Comparison Experiments

In the section, we will test certain problems by varying the encodings or genetic operators to compare their performances. We start with the comparison test on bit string encoding (Gray-coded integer stored as bit string) and floating-point encoding.

5.2.1 Comparison Experiments on Encodings

The function used to test the encodings is defined as follows:

\[ f(x_1, x_2, x_3, x_4) = 10x_1^2 - |x_2| + |x_3| + |x_4| , \]

where -5.0 ≤ x1 ≤ 5.0, -8.0 ≤ x2 ≤ 8.0, -10.0 ≤ x3, x4 ≤ 10.0. All are floating-point numbers.

This function has a minimum value of -8.0 at (0.0, -8.0, 0.0, 0.0). The reason we
choose such a relatively simple function is that both encodings will make observable progresses within a reasonable amount of generations.

The two tests differ only by representation and applicable crossover methods, and equivalent in all other aspects. Such an approach gives us a better basis for a relatively fair comparison.

Following is the average results of 20 runs given by both of the encodings:

<table>
<thead>
<tr>
<th>Number of Generations</th>
<th>Average Best Fitness (Bit string)</th>
<th>Average Best Fitness (Floating-point)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>-6.90264789</td>
<td>-7.85441872</td>
</tr>
<tr>
<td>200</td>
<td>-7.12828289</td>
<td>-7.91631431</td>
</tr>
<tr>
<td>300</td>
<td>-7.22735720</td>
<td>-7.95372628</td>
</tr>
<tr>
<td>400</td>
<td>-7.33161072</td>
<td>-7.97331920</td>
</tr>
<tr>
<td>500</td>
<td>-7.40571019</td>
<td>-7.98351918</td>
</tr>
<tr>
<td>600</td>
<td>-7.41846586</td>
<td>-7.98543751</td>
</tr>
<tr>
<td>700</td>
<td>-7.44085430</td>
<td>-7.98692431</td>
</tr>
<tr>
<td>800</td>
<td>-7.47598949</td>
<td>-7.98820886</td>
</tr>
<tr>
<td>900</td>
<td>-7.49707214</td>
<td>-7.99019063</td>
</tr>
<tr>
<td>1000</td>
<td>-7.49821985</td>
<td>-7.99132945</td>
</tr>
</tbody>
</table>

Table 5.8 Encoding comparison

To make it easier to compare the progress of the encodings, the following chart in the next page is drawn based on the results table.
It is easy to conclude from the results table and the comparison chart that floating-point encoding outperforms bit string encoding on all the 10 runs and converges much faster. The same conclusion was also reached in [22]. Another important standard not shown in the above figure but worth mentioning is the time performance. It took bit string encoding 60 seconds to finish a single run whereas floating-point only 7 seconds. That is, floating-point encoding is approximately 10 times faster than its contender. Needless to say, for optimization problems with pure floating-point variables, floating-point encoding is a better choice than bit string encoding.

5.2.2 Comparison Experiments on Selection Methods

In this subsection, we compare the performances of the different selection methods implemented in this package.
Experiment 1. Six-hump Camel-back Function

The first function used to test the encodings is defined as follows \cite{2}:

\[ f(x_1, x_2) = (4 - 2.1x_1^2 + \frac{1}{3}x_1^4)x_1^2 + x_1x_2 + (-4 + 4x_2^2)x_2^2. \]

where \(-3.0 \leq x_1 \leq 3.0\), \(-2.0 \leq x_2 \leq 2.0\), and they are both floating-point numbers. This function has a minimum value of \(-1.0316\) at \((-0.0898, 0.7126)\).

In this experiment, only the selection methods change from run to run, while all the other parameters remain the same. The average experiment results of 20 runs are given by all the selection methods are displayed in the table below:

<table>
<thead>
<tr>
<th>Number of Generation</th>
<th>Average Best Fitness from Selection Methods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Roulette Wheel</td>
</tr>
<tr>
<td>100</td>
<td>-1.03114500</td>
</tr>
<tr>
<td>200</td>
<td>-1.03140260</td>
</tr>
<tr>
<td>300</td>
<td>-1.03148664</td>
</tr>
<tr>
<td>400</td>
<td>-1.03151137</td>
</tr>
<tr>
<td>500</td>
<td>-1.03153135</td>
</tr>
<tr>
<td>600</td>
<td>-1.03154257</td>
</tr>
<tr>
<td>700</td>
<td>-1.03155473</td>
</tr>
<tr>
<td>800</td>
<td>-1.03157913</td>
</tr>
<tr>
<td>900</td>
<td>-1.03157961</td>
</tr>
<tr>
<td>1000</td>
<td>-1.03158692</td>
</tr>
</tbody>
</table>

Table 5.9 Selection methods comparison I

The following figure gives us a more direct portrait of the performance comparison:
The results indicate that all the selection methods converge to the optimal value with Roulette wheel the closest and SUS the furthest. Generally speaking, SUS has better performance than Roulette wheel\[^{[1]}\], but this is not observed in our case. This may be attributed to the relatively large population size used in our experiment, which helps Roulette wheel reduce the statistical error in the sampling process.

In the real world, we need a GA that not only converges to the global optimum, but also accomplishes the searching within a reasonable period of time. The following table presents the average time of the 20 runs of the selection methods.

<table>
<thead>
<tr>
<th>Selection Method</th>
<th>Time (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roulette Wheel</td>
<td>39</td>
</tr>
<tr>
<td>SUS</td>
<td>41</td>
</tr>
<tr>
<td>Baker Rank</td>
<td>76</td>
</tr>
<tr>
<td>Reeves Rank</td>
<td>79</td>
</tr>
<tr>
<td>Tournament</td>
<td>43</td>
</tr>
</tbody>
</table>

Table 5.10 Time complexity of selection methods I
Even though time performance depends on the machine on which the program is running, and thus may vary from machine to machine, the table still offers us a general view about the time performances of the experimented selection methods. We can see that the rank methods are slower than the non-rank methods. That is because the rank methods require the extra workload of sorting the entire population for each generation.

**Experiment 2. A Variant of Rosenbrock Function**

We do the experiment with a smaller population size using the following function:

\[ f(x_1, x_2, x_3, x_4) = 10x_1^4 + 5x_2^2 + |x_3| + |x_4|, \]

where \(-3.0 \leq x_1 \leq 3.0\), \(-2.0 \leq x_2 \leq 2.0\), \(-10.0 \leq x_3, x_4 \leq 10.0\), and they are all floating-point numbers. This function has a minimum value of 0.0 at (0.0, 0.0, 0.0, 0.0).

The average experiment results of 20 runs given by all the selection methods are demonstrated in the table below:

<table>
<thead>
<tr>
<th>Number of Generation</th>
<th>Average Best Fitness from Selection Methods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Roulette Wheel</td>
</tr>
<tr>
<td>200</td>
<td>0.62074692</td>
</tr>
<tr>
<td>400</td>
<td>0.24587467</td>
</tr>
<tr>
<td>600</td>
<td>0.12767855</td>
</tr>
<tr>
<td>800</td>
<td>0.06404160</td>
</tr>
<tr>
<td>1000</td>
<td>0.03956093</td>
</tr>
<tr>
<td>1200</td>
<td>0.02484145</td>
</tr>
<tr>
<td>1400</td>
<td>0.01926923</td>
</tr>
<tr>
<td>1600</td>
<td>0.01613904</td>
</tr>
<tr>
<td>1800</td>
<td>0.01147213</td>
</tr>
<tr>
<td>2000</td>
<td>0.01048350</td>
</tr>
</tbody>
</table>

Table 5.11 Selection methods comparison II

The average results are drawn in the following chart. Please note that the fitness value axis used a logarithmic scale ("semi-log") to make the small differences among selection methods more observable.
Similar to the previous experiment, the results indicate that all the selection methods are converging toward the optimal value. However, unlike experiment 1, Baker rank outperforms all the other competitors. Roulette wheel not winning the game may be attributed to the smaller population size employed in this experiment.

The average time performances are compared in the following table.

<table>
<thead>
<tr>
<th>Selection Method</th>
<th>Time (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roulette Wheel</td>
<td>52</td>
</tr>
<tr>
<td>SUS</td>
<td>58</td>
</tr>
<tr>
<td>Baker Rank</td>
<td>101</td>
</tr>
<tr>
<td>Reeves Rank</td>
<td>99</td>
</tr>
<tr>
<td>Tournament</td>
<td>60</td>
</tr>
</tbody>
</table>

Table 5.12 Time complexity of selection methods II

The same pattern is observed as in the previous experiment. That is, the non-rank methods are much faster than the rank methods.
There are other interesting phenomena worth mentioning. For example, SUS selection methods did not perform well in either experiment (second worst in test 2 and worst in the test 1). As for the two rank methods, Baker performs slightly better than Reeves in both the experiments.

5.2.3 Comparison Experiments on Crossover Methods

In this subsection, we conduct experiments to compare the performance of crossover methods implemented in this package.

Experiment 1. A Variant of Easom Function

The function selected for this experiment is defined as follows:\(^2\):

\[
f(x_1, x_2, x_3, x_4, x_5) = 1 - \cos(x_1) \cos(x_2) e^{-(x_1-\pi)^2-(x_2-\pi)^2} + 50x_3^4 + 10x_4^2 + 2|x_5|,
\]

where \(-3.0 \leq x_1, x_2 \leq 5.0\), \(-10.0 \leq x_3, x_4, x_5 \leq 10.0\), and they are all floating-point numbers. This function has a minimum value of 0.0 at \((\pi, \pi, 0.0, 0.0, 0.0)\).

The average experiment results of 20 runs given by all the crossover methods are demonstrated in the table below:

<table>
<thead>
<tr>
<th>Number of Generation</th>
<th>Single Point</th>
<th>Double Point</th>
<th>Uniform</th>
<th>Arithmetic</th>
<th>Heuristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>300</td>
<td>6.53963748</td>
<td>5.90342281</td>
<td>7.22753797</td>
<td>3.52300873</td>
<td>7.04832794</td>
</tr>
<tr>
<td>600</td>
<td>3.84814084</td>
<td>3.76738204</td>
<td>3.09755289</td>
<td>1.69161245</td>
<td>3.83451180</td>
</tr>
<tr>
<td>900</td>
<td>2.48216041</td>
<td>2.86849293</td>
<td>2.16271252</td>
<td>1.05478382</td>
<td>2.2649110</td>
</tr>
<tr>
<td>1200</td>
<td>1.91600364</td>
<td>1.82472980</td>
<td>1.77030236</td>
<td>0.76704362</td>
<td>1.93611046</td>
</tr>
<tr>
<td>1500</td>
<td>1.49884491</td>
<td>1.65801896</td>
<td>1.50038787</td>
<td>0.49035510</td>
<td>1.41774216</td>
</tr>
<tr>
<td>1800</td>
<td>1.1868305</td>
<td>1.48398141</td>
<td>1.19765466</td>
<td>0.33054074</td>
<td>1.23351500</td>
</tr>
<tr>
<td>2100</td>
<td>1.00443808</td>
<td>1.20197642</td>
<td>1.00505996</td>
<td>0.19385261</td>
<td>1.04875887</td>
</tr>
<tr>
<td>2400</td>
<td>0.83122766</td>
<td>1.00045785</td>
<td>0.85575393</td>
<td>0.09411635</td>
<td>0.90943283</td>
</tr>
<tr>
<td>2700</td>
<td>0.68335077</td>
<td>0.78665758</td>
<td>0.71635394</td>
<td>0.06433915</td>
<td>0.81680672</td>
</tr>
<tr>
<td>3000</td>
<td>0.50794689</td>
<td>0.65037215</td>
<td>0.54692489</td>
<td>0.04681860</td>
<td>0.69502116</td>
</tr>
<tr>
<td>3300</td>
<td>0.41780563</td>
<td>0.53414647</td>
<td>0.41821502</td>
<td>0.03354762</td>
<td>0.57154467</td>
</tr>
</tbody>
</table>

Table 5.13 Crossover methods comparison I
The results are drawn using logarithmic scale in the following chart:

![Comparison of Crossover Methods](image)

Figure 5.4 Crossover methods comparison I

Similar to the selection methods experiment, the results obtained from this experiment also indicate that all the crossover methods lead the GA to the convergence with Arithmetic the closest and Heuristic the furthest.

If we take a closer look, the performance of the contenders can roughly be divided into three groups: Arithmetic in the superior group, Uniform and Single point in the intermediate group, Heuristic and Double point in the inferior group. Arithmetic outperforming other competitors may be attributed to its unique capability of generating new offspring within the range of the parents and its ability of creating more genetic schemas than the single point and double point crossovers.
The time performances of the crossover methods are compared in the following table.

<table>
<thead>
<tr>
<th>Time (sec)</th>
<th>Single Point</th>
<th>Double Point</th>
<th>Uniform</th>
<th>Arithmetic</th>
<th>Heuristic</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>220</td>
<td>226</td>
<td>231</td>
<td>221</td>
<td>219</td>
</tr>
</tbody>
</table>

Table 5.14 Time complexity of crossover methods I

Unlike the time complexity variation observed in the selection methods test, the time performances of all the crossover methods are pretty close to each other.

**Experiment 2. A Variant of Squared Radius Function**

The experimented is repeated using the following function:

\[ y = \sum_{n}^{n} n x_n^2 \]

where \(-10.0 \leq x_n \leq 10.0\), for \(n = 1\) to \(5\), and are all floating-point numbers. This function has a minimum value of 0.0 at \((0.0, 0.0, 0.0, 0.0, 0.0)\).

The average experiment results of 20 runs given by all the crossover methods are demonstrated in the table below:

<table>
<thead>
<tr>
<th>Number of Generation</th>
<th>Single Point</th>
<th>Double Point</th>
<th>Uniform</th>
<th>Arithmetic</th>
<th>Heuristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>300</td>
<td>23.59322033</td>
<td>20.19726184</td>
<td>21.23134067</td>
<td>7.82278207</td>
<td>34.01789885</td>
</tr>
<tr>
<td>600</td>
<td>11.84879552</td>
<td>13.07524367</td>
<td>11.31882723</td>
<td>2.78603796</td>
<td>20.25611526</td>
</tr>
<tr>
<td>900</td>
<td>6.27316003</td>
<td>7.24433722</td>
<td>6.44028531</td>
<td>1.03588042</td>
<td>13.80272412</td>
</tr>
<tr>
<td>1200</td>
<td>4.17823117</td>
<td>4.32724269</td>
<td>3.24527675</td>
<td>0.53103251</td>
<td>10.13046719</td>
</tr>
<tr>
<td>1500</td>
<td>2.48152476</td>
<td>3.01946040</td>
<td>1.98943355</td>
<td>0.19708694</td>
<td>9.41284500</td>
</tr>
<tr>
<td>1800</td>
<td>1.88963018</td>
<td>2.34222704</td>
<td>1.40765973</td>
<td>0.08382558</td>
<td>7.07262043</td>
</tr>
<tr>
<td>2100</td>
<td>1.27979614</td>
<td>1.36751549</td>
<td>0.99008349</td>
<td>0.04833465</td>
<td>4.94206167</td>
</tr>
<tr>
<td>2400</td>
<td>1.03410710</td>
<td>0.96161507</td>
<td>0.80888099</td>
<td>0.02563513</td>
<td>4.05980383</td>
</tr>
<tr>
<td>2700</td>
<td>0.81998893</td>
<td>0.71481592</td>
<td>0.52794773</td>
<td>0.01396669</td>
<td>2.77400604</td>
</tr>
<tr>
<td>3000</td>
<td>0.56398378</td>
<td>0.54536879</td>
<td>0.38935216</td>
<td>0.00879699</td>
<td>2.28372226</td>
</tr>
<tr>
<td>3300</td>
<td>0.46887681</td>
<td>0.45644971</td>
<td>0.30927899</td>
<td>0.00636384</td>
<td>2.17092762</td>
</tr>
</tbody>
</table>

Table 5.15 Crossover methods comparison II

The results are drawn using logarithmic scale in the following chart:
The same convergence pattern is observed in this experiment. Moreover, the performance hierarchy is nearly preserved: Arithmetic, Uniform, Double Point, Single Point, and Heuristic. In both the competitions, Arithmetic is the leading performer whereas Heuristic is the worst player.

The time performances of the crossover methods are compared in the following table.

<table>
<thead>
<tr>
<th></th>
<th>Single Point</th>
<th>Double Point</th>
<th>Uniform</th>
<th>Arithmetic</th>
<th>Heuristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time (sec)</td>
<td>218</td>
<td>222</td>
<td>227</td>
<td>230</td>
<td>226</td>
</tr>
</tbody>
</table>

Table 5.16 Time complexity of crossover methods II

Like the time complexity observed in the experiment 1, the time performances of all the crossover methods are at the same level.
As we have mentioned in Chapter 3, the performance of crossover methods have complicated relationships with the fitness function, encoding, and other parameters of the GA. There is no crossover method that guarantees to work best in all the cases. With that said, however, the experiments of this section still offer us some guidance on the usage of crossover methods. In general, Arithmetic or Uniform crossover should be the first choice. Only when you are not satisfied with the results obtained, then consider trying other candidates.
Chapter VI. Conclusions and Future Work

6.1 Conclusions

Genetic Algorithm models the natural selection and evolution process and has been successful in areas where the traditional methods fall short. GA has quite a number of advantages over traditional optimization methods: it can optimize both continuous and discrete functions; it does not require complicated differential equations or a smooth objective function; it can be highly parallelized to improve computation performance; it searches more of the solution space and thus increases the probability of finding the global optimum.

Most of the existing GA packages either use single representation or offer limited choices of genetic operator. To handle the variety of optimization problems in practical applications, a more comprehensive and powerful general GA package is designed and implemented in this thesis. This package gives users a rich set of options on the problem representation and genetic operators based on the specific problem type. The following list outlines the feature methods that are implemented in the package:

- **Encodings:** Bit strings, Floating point, Integer (Gray code)
- **Selection Methods:** Improved Roulette Wheel, Stochastic Universal Sampling, Baker Linear Rank, Reeves Rank, Tournament
• **Generation Replacement Models:** Generational, Steady State, Evolution Strategy 
  \((N+m)\) Replacement, Elitism

• **Crossover Methods:** Single Point, Double Point, Uniform, Arithmetic, Heuristic

• **Mutation Methods:** Uniform, Dynamic(Non-uniform), Boundary, Creep

• **Termination Criteria:** Hybrid, Number of Generations, Convergence, Progress in Fitness

The package has been tested through two groups of experiments. The first group tests the general robustness by inputting a variety of optimization problems to the package. The second group compares the performances of different genetic operators by varying the encoding, selection, or crossover methods on a certain input function. The experiments results indicate that the GA package achieved convergence and found satisfactory solution.

Some other experiments are made but are not listed in Chapter 5. Based on all the experiments conducted, I summarized the following tips regarding the usage of this package:

• To optimize problems with pure floating-point variables, floating-point encoding outperforms bit string encoding in both the solution quality and time required.

• Baker rank SUS are both recommended for selection methods. If time is also considered, then SUS has the upper hand.

• Arithmetic and Uniform are recommended for crossover methods.

• Uniform and Dynamic are recommended for mutation methods.
• Generational replacement converges faster than Steady state, but has more risk of leading the GA to a local optimum point. In contrast, Steady state converges relatively slow, but usually leads the GA to the global optimum point.

• For termination criteria, Number of generations is recommended. The other two criteria have small probabilities to lead the GA into infinite loop in case the expected progress is not made. In this package, this case is handled by specifying an upper bound of loops.

Once again, these tips just serve as general suggestions. Please keep in mind that the performances of genetic operators are intricately related to the problem context and other GA parameters. No operator is guaranteed to work best in all the cases. When the results obtained are inferior to the expectation, the best way is to adjust the GA parameters and give it another try.

6.2 Future Work

More experiments can be done in effort to resolve the influence of various factors on the performance of genetic algorithms. In particular, the answer to question like how to select the best representation and operators for a specific problem will save the time people spend on trying different sets of GA configurations.

There is still room for improvement to make this package more powerful. The following features may be considered to be added into this package in the future:
The combination with other optimization methods: Needless to say, GA is a powerful optimization technique. On the other hand, there exists a rich set of optimization methods and each of them has its unique strength. The combination of GA with these methods will help yield better solution on some problems. For example, after a certain number of generations, GA is probably in the valley where the global optimum resides, and a hill-climbing method might be used to take over the task hereafter to locate the optimum point more rapidly and accurately, providing the function is smooth (continuously differentiable).

Parallelism: Genetic algorithms are very suitable for parallel implementation. The inclusion of parallelism will enable this GA package to search a wide range of sampling space simultaneously, and thus greatly reduce the time complexity.
REFERENCES


APPENDIX

INPUT FILES OF TESTS AND EXPERIMENTS

Input file for Flat-round Rosenbrock function

Population Size: 500
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 100589
###
Encoding: Mix
###
###Selection methods include: Roulette Wheel, SUS,Baker Rank,Reeves Rank,Tournament
###Crossover methods include: Single Point, Double Point,Uniform, Arithmetic, Heuristic
###Mutation methods include: Boundary, Creep
###Replacement Method include: Generational,Steady State,Evolution Strategy
###
Selection Method: SUS
Crossover Method(Non-FP): Uniform
Crossover Method(FP): Arithmetic
Mutation Method(Non-FP): Uniform
Mutation Method(FP): Uniform
Replacement Method: Steady State
###
Termination Criterion: Number of Generations
Number of Generations: 1000
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
###Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 1
FP Var1 Range: -2.5,2.5
###
Number of Integer Variables: 1
  Int Var1 Range: -3, 3
###

Input file for Hollow-round Rosenbrock function

Population Size: 500
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 100589
###
Encoding: Mix
###
### Selection methods include: Roulette Wheel, SUS, Baker Rank, Reeves Rank, Tournament  
### Crossover methods include: Single Point, Double Point, Uniform, Arithmetic, Heuristic  
### Mutation methods include: Boundary, Creep  
### Replacement Method include: Generational, Steady State, Evolution Strategy  
### Selection Method: SUS  
Crossover Method (Non-FP): Uniform  
Crossover Method (FP): Arithmetic  
Mutation Method (Non-FP): Uniform  
Mutation Method (FP): Dynamic  
Replacement Method: Generational  
### Termination Criterion: Number of Generations  
Number of Generations: 1000  
#Percentage of Converged Alleles: 0.9  
#Factor of Fitness Progress: 0.01  
### Variables setting  
Number of Bit-string Variables: 0  
Number of Floating-point Variables: 1  
  FP Var 1 Range: -2.5, 2.5  
###  
Number of Integer Variables: 1  
  Int Var 1 Range: -3, 3  
###  
Weights and profits of Knapsack problem with 100 items

| Item 1: 6, 98 | Item 2: 3, 67 | Item 3: 8, 86 | Item 4: 2, 75 | Item 5: 8, 71 | Item 6: 3, 76 | Item 7: 3, 60 | Item 8: 10, 83 | Item 9: 2, 63 | Item 10: 5, 58 | Item 11: 2, 58 | Item 12: 7, 81 | Item 13: 5, 52 | Item 14: 1, 91 | Item 15: 1, 83 | Item 16: 4, 98 | Item 17: 8, 93 | Item 18: 9, 99 | Item 19: 9, 51 | Item 20: 8, 82 | Item 21: 4, 89 | Item 22: 5, 53 | Item 23: 4, 76 | Item 24: 2, 62 | Item 25: 5, 71 |
|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|
Input file for Knapsack problem with 100 items

Population Size: 100
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 100589
###
Encoding: Bit String
###
Selection Method: SUS
Crossover Method(Non-FP): Uniform
Crossover Method(FP): Uniform
Mutation Method(Non-FP): Uniform
Mutation Method(FP): Uniform
Replacement Method: Steady State
###
Termination Criterion: Number of Generations
Number of Generations: 2000
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
###Variables setting
Number of Bit-string Variables: 1
String Var1 length: 100
Number of Floating-point Variables: 0
###
Number of Integer Variables: 0
###

Input file for Bohachevsky function

Population Size: 600
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 10058
###
Encoding: Mix
###
Selection Method: SUS
Crossover Method(Non-FP): Uniform
Crossover Method(FP): Arithmetic
Mutation Method(Non-FP): Uniform
Mutation Method(FP): Dynamic
Replacement Method: Steady State
###
Termination Criterion: Number of Generations
Number of Generations: 3000
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
###Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 2
FP Var1 Range: -5, 5
FP Var2 Range: -5, 5
###
Input file for Schaffer function F7

Population Size: 600  
Crossover Rate: 0.4  
Mutation Rate: 0.1  
###  
Seed for Random Function: 10058  
###  
Encoding: Mix  
###  
Selection Method: SUS  
Crossover Method(Non-FP): Uniform  
Crossover Method(FP): Arithmetic  
Mutation Method(Non-FP): Uniform  
Mutation Method(FP): Dynamic  
Replacement Method: Steady State  
###  
Termination Criterion: Number of Generations  
Number of Generations: 4000  
Percentage of Converged Alleles: 0.9  
Factor of Fitness Progress: 0.01  
###  
### Variables setting  
Number of Bit-string Variables: 0  
Number of Floating-point Variables: 2  
FP Var1 Range: -5, 5  
FP Var2 Range: -5, 5  
###  
Number of Integer Variables: 0  
###

Input file for encoding comparison (bit-string)

Population Size: 800  
Crossover Rate: 0.4  
Mutation Rate: 0.4  
###  
Seed for Random Function: 10059  
###  
Encoding: Bit String  
###  
Selection Method: SUS  
Crossover Method(Non-FP): Uniform  
Crossover Method(FP): Uniform  
Mutation Method(Non-FP): Uniform  
Mutation Method(FP): Uniform  
Replacement Method: Generational  
###  
Termination Criterion: Number of Generations  
Number of Generations: 1000  
Percentage of Converged Alleles: 0.9  
Factor of Fitness Progress: 0.01  
###  
### Variables setting  
Number of Bit-string Variables: 0  
Number of Floating-point Variables: 2  
FP Var1 Range: -5, 5  
FP Var2 Range: -5, 5  
###  
Number of Integer Variables: 0  
###
Number of Bit-string Variables: 0
Number of Floating-point Variables: 4
FP Var1 Range: -5, 5
FP Var2 Range: -8, 8
FP Var3 Range: -10,10
FP Var4 Range: -10,10
###
Number of Integer Variables: 0
###

###Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 4
FP Var1 Range: -5, 5
FP Var2 Range: -8, 8
FP Var3 Range: -10,10
FP Var4 Range: -10,10
###
Number of Integer Variables: 0
###

Input file for encoding comparison (floating point)

Population Size: 800
Crossover Rate: 0.4
Mutation Rate: 0.4
###
Seed for Random Function: 10059
###
Encoding: Bit String
###
Selection Method: SUS
Crossover Method(Non-FP): Uniform
Crossover Method(FP): Heuristic
Mutation Method(Non-FP): Uniform
Mutation Method(FP): Uniform
Replacement Method: Generational
###
Termination Criterion: Number of Generations
Number of Generations: 1000
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
###Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 4
FP Var1 Range: -5, 5
FP Var2 Range: -8, 8
FP Var3 Range: -10,10
FP Var4 Range: -10,10
###
Number of Integer Variables: 0
###

Input file for selection methods comparison I

#Change the selection method(FP) for each run.
Population Size: 300
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 10058
###
Encoding: Mix
###
Selection Method: Roulette Wheel
Crossover Method(Non-FP): Uniform
Crossover Method(FP): Arithmetic
Mutation Method (Non-FP): Uniform
Mutation Method (FP): Dynamic
Replacement Method: Steady State
###
Termination Criterion: Number of Generations
Number of Generations: 1000
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
### Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 2
   FP Var1 Range: -3, 3
   FP Var2 Range: -2, 2
###
Number of Integer Variables: 0
###

Input file for selection methods comparison II

# Change the selection method (FP) for each run.
Population Size: 200
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 10058
###
Encoding: Mix
###
Selection Method: Roulette Wheel
Crossover Method (Non-FP): Uniform
Crossover Method (FP): Arithmetic
Mutation Method (Non-FP): Uniform
Mutation Method (FP): Dynamic
Replacement Method: Steady State
###
Termination Criterion: Number of Generations
Number of Generations: 1000
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
### Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 4
   FP Var1 Range: -3, 3
   FP Var2 Range: -2, 2
   FP Var3 Range: -10, 10
   FP Var4 Range: -10, 10
###
Number of Integer Variables: 0
###

Input file for crossover methods comparison I

# Change the crossover method (FP) for each run.
Population Size: 500
Crossover Rate: 0.4
Input file for crossover methods comparison II

#Change the crossover method(FP) for each run.
Population Size: 500
Crossover Rate: 0.4
Mutation Rate: 0.1
###
Seed for Random Function: 1005896
###
Encoding: Mix
###
Selection Method: SUS
Crossover Method(Non-FP): Uniform
Crossover Method(FP): Single Point
Mutation Method(Non-FP): Uniform
Mutation Method(FP): Dynamic
Replacement Method: Steady State
###
Termination Criterion: Number of Generations
Number of Generations: 3300
Percentage of Converged Alleles: 0.9
Factor of Fitness Progress: 0.01
###
###Variables setting
Number of Bit-string Variables: 0
Number of Floating-point Variables: 5
FP Var1 Range: -10, 10
FP Var2 Range: -10, 10
FP Var3 Range: -10, 10
FP Var4 Range: -10, 10
FP Var5 Range: -10, 10
###
Number of Integer Variables: 0
###
FP Var3 Range: -10,10
FP Var4 Range: -10,10
FP Var5 Range: -10,10
###
Number of Integer Variables: 0
###
VITA

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