

# An Approach To The Traveling Salesman Problem Using A Hybrid Genetic Algorithm

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This paper explores the application and performance testing of a hybrid genetic algorithm implemented in Python. Variations of crossover and mutation are compared in a battle for fastest time to (near) convergence. The results lead us to infer that there is a need for reinforcement learning to exit deep local minima.

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## 1. INTRODUCTION

Mathematically described as the shortest Hamiltonian circuit, the Traveling Salesman Problem describes the problem of determining the shortest path length of a weighted graph traversal of  $n$  nodes with the constraint that the start and end node are the same. The problem was tested on a data set consisting of three real-world instances of the problem, namely graphs derived from maps of the Western Sahara, Uruguay, and Canada. These instances consist of 29, 734, and 4663 nodes respectively.

Since each path is a ordered set of nodes, the chosen representation is a sequence whose non-repeating elements consist of each node visited, in the path order. This sequential set is hereafter referred to as the *chromosome* to make the relationship to genetic crossover and mutation more apparent.

Representing the path this way is very compact, and can represent any solution path. A tree could be an alternative representation, however

this would introduce an extra layer of complexity into the problem. One key aspect of a representation is that it must be able to express our target solution, which is an evident characteristic.

There exists a positive correlation between the number of nodes and the computational complexity of the problem. The search space of this representation grows as an unbounded monotonic factorial function. For the Western Sahara data set, there are  $3.05 \times 10^{29}$  solutions alone. Uruguay has only twenty-five times as many cities, however the search space has  $4.12 \times 10^{1783}$  solutions. The situation is even bleaker for the Canada set, which has a whopping  $2.73 \times 10^{15080}$  solutions.

## 2. METHODOLOGY

A genetic algorithm has a number of elements that make variants unique from each other. Each algorithm has an initialization procedure, evaluation function, parent selection methodology, recombination operator(s), mutation operator(s), and survivor selection methodology. Each of these will be described in the following sections.

### A. Initialization: Clustering

As previously discussed, the search space for this problem is enormously large. Even with the smallest data set, brute-force search on the Western Sahara data set would require

$3.05 \times 10^{29}$  evaluations to find the optimal solution. Assuming that we are able to evaluate one solution every microsecond, it would still take 100,000,000,000,000 *centuries* to exhaustively evaluate every solution.

There is a strong need to remove unlikely candidates from the solution space. One method to reduce the number of solutions is to group nearby nodes into individual *clusters*.

Being one of the most popular clustering algorithms, *k-means clustering* was chosen to organize the nodes. Given a number  $k$ , all the nodes are separated into  $k$  clusters. Each cluster has a node that acts as the centre of the cluster. Then, the algorithm iterates over each node and assigns it to the cluster with the centre node that it is closest to. Conceptually, it can be thought of as dividing the TSP problem into smaller TSP sub-problems.[1][2] The algorithm can be concisely expressed as the following series of steps:

1. Given a number  $k$ , randomly pick  $k$  cities that will act as cluster centres.
2. For every city  $C$ :
  - (a) Calculate the distance between  $C$  and every cluster centre city.
  - (b) Pick the cluster with the centre that is closest to  $C$ .
  - (c) Add the city  $C$  to that cluster.
3. For every cluster\*:
  - (a) For every city  $C_1$ , calculate the cumulative distance between itself and every other city in the cluster.
  - (b) Pick the city with the smallest cumulative distance to every other city and assign it to be the new centre of the current cluster.
  - (c) Repeat **Step 3**  $n$  times (where  $n$  is specified in advance).
4. Now that the clusters have been established, order all the clusters by their centres, so that the closest clusters stay close to each other.

\*Step 3 is usually performed until the clusters become static. Due to computational limitations, we decided that it is better to constrain the number of iterations for **Step 3** to  $n$  (where  $n$  is specified in advance in the configuration).

## B. Evaluation

As discussed previously, our population is modeled as instances of orderings. This representation can be evaluated simply by adding together the weights of every consecutive pair of edges of the node pair. The calculation will yield a sum, which is to be minimized. The weights between all pairs of nodes have been added to a two-dimensional array, ensuring that this addition is a constant-time operation.

## C. Parent Selection

Tournament selection was the method of choice for parent selection. It was chosen due to the fact that it adds an extra element of randomness to the algorithm to try to combat the problem of reaching local minima. It involves the following steps:

1. Pick  $k$  individuals randomly, without replacement.
2. Out of these  $k$  individuals, select the individual with the best fitness and add it to the mating pool.
3. Repeat until a mating pool of the desired size is obtained.

## D. Recombination

To perform crossover between chromosomes, two operators were implemented and compared. The first of which, called "cut-and-crossfill" is implemented, which is then compared to a newer method called "best-order".

**Cut-And-Crossfill:** As described in Eiben's book, cut-and-crossfill is a relatively simple algorithm for recombining two chromosomes into a pair of chromosomes, using information from both parent chromosomes[3].

The algorithm can be concisely expressed as the following series of steps:

1. Define a new empty arrays  $o_1$  and  $o_2$

2. Choose a random point  $j$  on the interval  $[1, k - 1]$  where  $k$  is the length of the chromosome.
3. Copy elements  $[0, j]$  from the first parent into  $o_1$
4. Append to  $o_1$  the element  $[j + 1, k]$  of the first parent
5. Copy elements  $[0, j]$  from the second parent into  $o_2$
6. Append to  $o_2$  the elements  $[j + 1, k]$  of the second parent

The algorithm's ease-of-implementation and computational speed make it the algorithm of choice to compare to newer, more complicated operators.

**Best Order Crossover:** Best-order crossover is predicated on the assumption that the order of the alleles is more important than their positions. This operator incorporates genetic material from the current best chromosome in the population along with the genetic material from the two parent chromosomes participating. Thus, the order of particular genes in the offspring can be informed by the order of those genes in the best individual chromosome present in the population[4].

In the current problem context, the information describing the order of alleles with respect to each other is more important than a particular allele being present in a specific location, since there is no restriction on the starting position.

The Best Order Crossover operator involves the following steps:

1. Pick  $n$  random crossover points, resulting in  $n + 1$  sub-sequences.
2. For each resulting sub-sequence:
  - (a) Generate a random integer between  $r$ , where  $1 \leq r \leq 3$ .
  - (b) If  $r == 1$ , then the alleles corresponding to this subsequence will be taken from Parent 1 in the order that they are in Parent 1.
  - (c) If  $r == 2$ , then the alleles corresponding to this subsequence will be taken from Parent 1, but in the order that they appear in Parent 2.
  - (d) If  $r == 3$ , then the alleles corresponding to this subsequence will be taken from Parent 1, but in the order that they appear in the best individual in the current generation.

## E. Mutation

For mutation, we used permutation swap, insertion mutation, inversion mutation, scramble mutation, two-opt mutation and a cyclic "random" mutation operator.

**Permutation Swap:** This operator picks two random alleles in the chromosome and swaps their positions.

**Insertion Mutation:** This operator involves the following steps:

1. Pick two random alleles  $n_1$  and  $n_2$
2. Insert  $n_1$  adjacent to  $n_2$

**Inversion Mutation:** This mutation operator picks two random alleles in the chromosome and inverts (reverses) the sub-sequence of alleles between them. The length of the sub-sequence to be inverted/reversed can be specified in advance.

**Scramble Mutation** For this mutation operator, two random alleles in the chromosome are chosen and the sub-sequence of alleles between them are scrambled. The length of the sub-sequence to be scrambled can be specified in advance.

**2-Opt Swap Mutation** This operator was inspired by the 2-opt local search algorithm, which is often employed to solve the Travelling Salesman Problem (TSP). This mutation operator picks two adjacent pairs of alleles and swaps their respective elements[5].

This operator involves the following steps:

1. Pick a random alleles  $n_1$  and  $n_3$
2. Choose  $n_2$  such that it is an arbitrarily adjacent node to  $n_1$ , and likewise choose a  $n_4$  adjacent to  $n_3$ .

3. Treat the four alleles as adjacent pairs, where the first pair is  $(n_1, n_2)$  and the second pair is  $(n_3, n_4)$ .
4. Swap the elements the pairs with each other.

**Cyclic Random Mutation** An operator was devised that randomly picks a mutation operator out the aforementioned operators. The idea was that different operators are helpful for getting out of different types of local minima.

### F. Survivor Selection

We used  $(\mu + \lambda)$  selection for survivor selection, which involves the following steps:

1. Pool all  $\mu$  current generation individuals and  $\lambda$  offspring together.
2. Rank them by fitness
3. Pick the top  $\mu$  individual to enter the next generation

## 3. DISCUSSION

For all results, the Uruguay data set was used as the basis of comparison, with all other parameters of the algorithm held constant, other than the parameter under test. These parameters are expressed in Table 1. Uruguay was chosen due to the fact that the solution space is very large and the we can perform a run of this algorithm in a manageable amount of time.

### A. Smart Initialization

While the k-means clustering algorithm reduces the search space substantially, it does so by yielding a deep local minima. This local minima can reach as close to 113% of the optimal solution, which is very close to the optimum result. After the clusters of nodes are defined, there is a need to order them in their own Hamiltonian cycle, which is basically a smaller version of the TSP problem. Augmenting the k-means algorithm to properly order the initial clusters will yield even better results. The hybrid algorithm results in good quality solutions in a minimal number

**Table 1. Parameters Of The Hybrid Genetic Algorithm**

Parameter	Value
Data Source	TSPUruguay734.txt
Pop. Size	20
Init. Method	$k - means$
Mating Pool Size	10
Tournament Size	5
Crossover Rate	90%
Mutation Rate	80%
Generations	1000
BOX Cutting Points	240
K-Means Clusters	220
K-Means Iterations	5

of generations. Without the clustering, the algorithm starts at a much worse fitness value, as evidenced by Figure 1. The difference is substantial, in that random initialization resulted in a mean starting fitness of 1,579,829, whereas k-means was 121,647, an order of magnitude better.

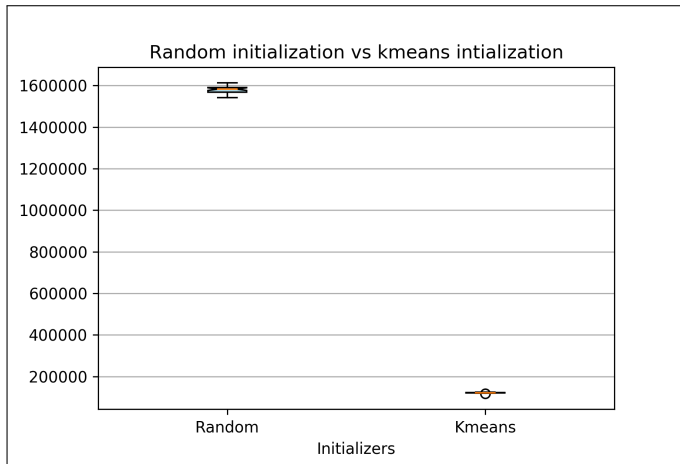
### B. Crossover

Figure 2 alludes to the idea that the mean fitness values are lower (better) for best-order crossover compared to cut-and-crossfill.

Plotting a notched boxplot in Figure 3, it is evident that the confidence intervals are nearly stacked end to end, in favor of best-order crossover. It is noteworthy that there are no outliers in each set, and the range is similar in each case.

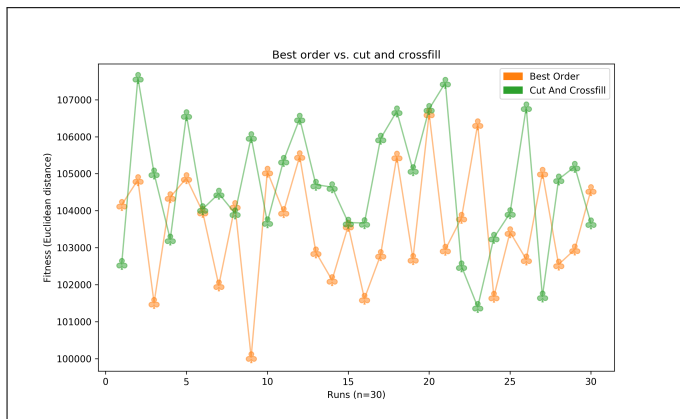
Figure 4 plots the histogram of each run. It is not apparent that the samples are normally distributed, so we have chosen the Mann-Whitney-U test as the statistical test of choice, since it does not require the assumption of normality.

For each crossover algorithm, thirty runs were conducted with all other parameters remaining



**Fig. 1.** Boxplots indicating the distribution of initial fitnesses using both kmeans and random initialization.

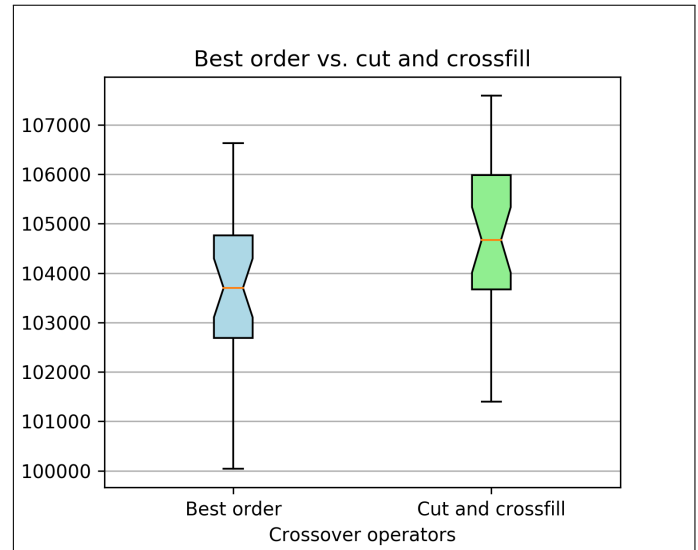
constant. A one-tailed Mann-Whitney-U test to test the null hypothesis that cut-and-crossfill has a mean best fitness equal to or smaller than best-order crossover. Obtaining a p-value of 0.0099, we reject the aforementioned null hypothesis. The alternative hypothesis is accepted, favoring the best-order crossover with 95% confidence.



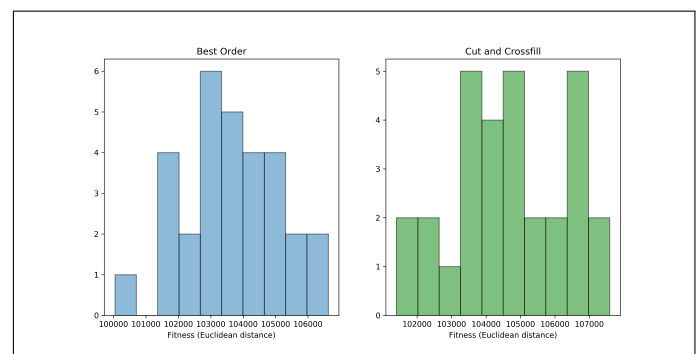
**Fig. 2.** Comparison of the mean best fitness of best-order vs. cut-and-crossfill crossover consisting of thirty runs each.

### C. Mutation

Figure 5 gives a clear indication of the performance of each mutation operator. There is a clear gap in the performance as evidenced by the non-overlapping range of samples. It is evident that the best performing mutation operators are

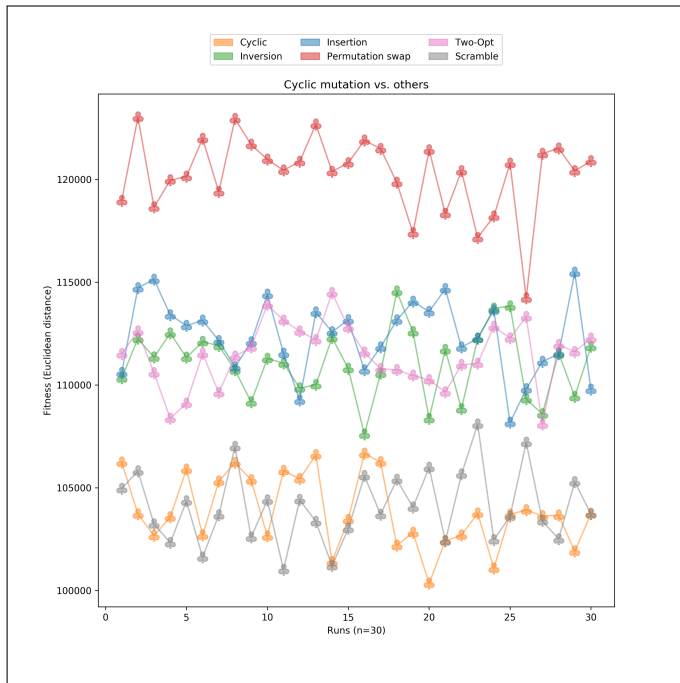


**Fig. 3.** Notched boxplots of the best fitnesses of best-order vs cut-and-crossfill crossover.

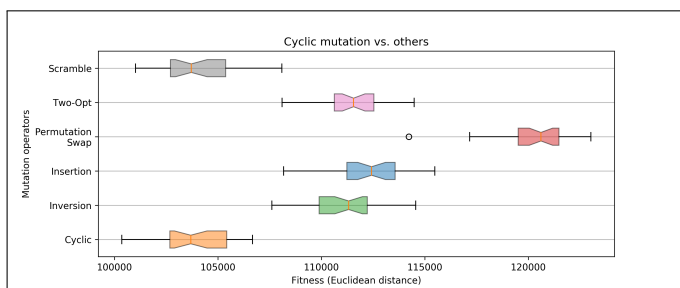


**Fig. 4.** Illustration of the distribution of samples of the best individual achieved in samples of thirty runs of each crossover type.





**Fig. 5.** Comparison of the mean best fitnesses produced by various types of mutation operators.



**Fig. 6.** Notched boxplots of the distribution of best fitness for various types of mutation operators.

cyclic and scramble, however we cannot make any conclusion as to which is better. Looking at the notched boxplots in Figure 6 we notice that, between cyclic and scramble, the confidence intervals and median are nearly the same. The scramble mutation has a slightly larger range in the right tail, but there is not enough information to determine whether or not this is statistically significant.

It is important to note that the resultant mean fitness of both cyclic and scramble is about 104,000. This 131% of the optimum distance of 79,114. This result is acceptable to us, in that we have only calculated 1000 generations to achieve this result.

## 4. CONCLUSION

Given more time, considerable focus should be spent on determining a method to get a closer match to the optimal ordering of clusters. There is potential to use a cluster-boundary informed sub-segment swap to as a mutation operator to further optimize the algorithm. Raven's *Biology*[6] describes the idea of *transposons*, sub-segments of a chromosome that can move to a different neighborhood.

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