

# Enhancing the Performance of Genetic Algorithms in Combinatorial Optimization of Large and Difficult Problems

## Using Variable Adaptive Mutation Rates Controlled by 'Inbreeding'

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### ABSTRACT

Blah blah blah, here is our abstract.

### Keywords

Genetic Algorithms, Combinatorial Optimization, N Queens Problem, Variable Mutation

### 1. INTRODUCTION

Genetic algorithms (GA) serve an important role in various applications, but particularly useful areas are those where they can perform stochastic generation of solutions for combinatorial problems [Using Genetic algorithms to solve np-complete problems][Solving the n-Queens Problem using Genetic Algorithms]. One notable example of this is the N-Queens problem. One approach to solving the N-Queens problem using a genetic algorithm is to implement chromosomes which model the position of each queen on the chessboard. The permuted values stored in the chromosome represent the sequential row positions of each queen, encoded into binary values. Each column value is the index, since there can never be more than one queen per column. Thus, the column positions increase by one for each queen and the intersections (referred to as collisions) are calculated per row or on the diagonal to determine whether a solution has been found. As with any genetic algorithm, evaluating the fitness of the chromosomes per generation is required to determine if a usable solution has been generated [Genetic Algorithms: a survey].

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By tuning the parameters of the genetic algorithm, performance can increase or decrease based on the solution landscape of the problem encountered [Some improvements on adaptive genetic algorithms for reliability-related applications][Genetic Algorithms and Directed Adaptation][Adaptive probabilities of crossover and mutation in genetic algorithms]. We have found that by replacing a static mutation value with an adaptive value controlled by chromosome similarity, better performance can be achieved than using static values, particularly when there is no *a priori* knowledge about the ideal static mutation rate. This is compounded when the solution landscape is large, such as in higher-order N-Queens problems.

### 1.1 Background and Problem Domain

Like many other complex optimization problems, the N-Queens problem becomes orders of magnitude more complex as the number of queens and the size of the chessboard increases [The N-queens problem and genetic algorithms]. Finding all solutions is a simple but non-trivial problem. When two or more queens share a row or diagonal, a collision occurs for each pair of queens, meaning that the board state is not a solution.

The Eight-Queens problem is the classical version of N-Queens. Even with this simplistic configuration, the problem is computationally expensive. There are  $\binom{64}{8}$  arrangements of the queens on a standard  $8 \times 8$  board. Exhaustive deterministic evaluation has shown that there are only 92 distinct solutions. Even including optimizations such as imposing constraints on placing only one queen per row, there are still  $8^8$  possible distinct arrangements of queens. In higher-order versions of the problem, combinatorial explosion occurs. The Nine-Queens problem has 352 distinct solutions. The Ten-Queens problem has 724. Table 2 in section 2 shows the growth of the number of solutions toward intractability.

While the number of solutions is known for problems involving up to 26 queens, there is currently no known formula to determine the exact number of distinct solutions for more complex problems. That is to say, higher order N-

Queens problems cannot be solved for all solutions exhaustively. Deterministic methods for approaching these problems are largely useless[CITE].

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- Cite me![2, 4, 1, 6, 7, 5, 3]

### 1.1.1 The N-Queens Puzzle

The natural question raised when approaching simple problems is why an exhaustive deterministic strategy is not used. Unfortunately, these approaches cannot work within human time-frames except in smaller problem landscapes. The issue is that a direct linear traversal of the solution landscape does not yield fruitful results easily in the case of the N-Queens problem. This stems from the fact that the relative fitness of each sequential board state doesn't matter unless it is a valid solution. The problem cannot be solved by producing a solution that has queens which will attack each other. Therefore, only solutions with the absolute maximum, 100% fitness, are useful solutions [Solving the n-Queens Problem using Genetic Algorithms]. What is more, as the problem landscape increases in size, the number of distinct solutions does not increase proportionally. A full brute force traversal of the landscape would be required to find all the distinct solutions, which is increasingly expensive in higher order problems.

Other stochastic solutions are impractical with this type of problem. This is because finding solutions among the landscape of possible board states yields a tremendous amount of duplicate solutions. Some method is required to direct the random traversal of the landscape toward distinct solutions and away from those which have already been found. The problem with this statement is that the nature of randomness cannot be constrained in this way - it will no longer be random. Therefore, methods such as genetic algorithms become an attractive solution approach because they represent a 'smarter' traversal of the problem landscape than what is offered by a linear brute force traversal, or a uniformly random traversal of the landscape[CITE]. Less time is spent on areas of the landscape which are not likely to be solutions. This leads us toward the core motivation of our solution's behavior.

### 1.1.2 Motivation

Deterministic modeling of the randomness of the solution landscape as seen in the N-Queens problem is impossible with known mathematical methods. To produce a geometric solution landscape would require that the dependent axis of the coordinate plane maps each sequential value to a uniformly random value in the actual solution landscape. Conceptually this is very difficult to imagine, but the reason why this conceptual strategy is useful will become clearer - first, we will discuss why a new approach is necessary in comparison to traditional GA approaches.

There are several optimizations to generating solutions which are available when approaching the N-Queens problem because of the symmetrical nature of the chess board. Since it is always square, one solution can be used to yield many other unique solutions using symmetry operations (reflection and rotation of the board). This means that up to 8 solutions can be generated by finding one distinct solution. However, symmetry operations cannot be used to find all distinct solutions because many symmetrical configurations of the board are equivalent. In the example of the Eight-

Queens problem, there are 11 board configurations which yield 8 solutions, and 1 which yields only 4. The result is still 92 unique solutions, with the majority of them calculable through symmetry operations alone.

Using symmetry operations, our initial exploration into solving the N-Queens problem yielded solutions quickly, but not quickly enough to beat deterministic solution generation methods. This was especially apparent in lower order problems where the solution landscape is small. We began to explore biological nuances of genetics in relation to genetic algorithms, and sought inspiration from the fact that nature has an apparent system for maintaining genetic variation within species. For example, pure-bred dogs often tend to have significant health problems which directly result from inbreeding [CITE]. Royal families historically used inbreeding to keep their line 'pure', which also resulted in genetic abnormalities[CITE].

These facts led us toward a potential approach to apply the effects of inbreeding towards the optimization of genetic algorithms. Our approach would make the mutation rate variable based on chromosome similarity, evoking the increased mutations seen inbreeding as a result of a too-similar biological chromosome. Our initial solution worked exceptionally well, providing all 92 solutions nearly instantaneously in comparison to our previous static mutation rate GA which required several seconds. This motivated us to explore the implications of our newfound strategy further.

## 1.2 Related Work

Classical genetic algorithms have been used extensively as a means to perform optimization in engineering for many years [Genetic and evolutionary algorithms come of age][Genetic Algorithms: a survey][Using Genetic algorithms to solve np-complete problems]. A popular problem to benchmark optimization algorithms in general is to use the N-Queens problem [The N-Queens Problem and Genetic Algorithms][Solving the n-Queens Problem using Genetic Algorithms]. However, as stated in [AUTHOR Solving the N-Queens Problem using Genetic Algorithms], the N-Queens problem is very different from most NP-Complete problems. This stems from the fact that in spite of the fitness functions used in the algorithm, solutions can only evaluate to a binary value. This prevents a directed search through the solution landscape since there is no feedback besides the board state found being absolutely correct or incorrect.

One major concern with using classical GA is determining the best parameters for genetic operators. That is to say, the probability of mutation and crossover. These operators' values are defined prior to the GA execution and remain static for the entire execution. Hesser and MÄd'ner's investigations define an algorithmic solution providing suitable, optimized values for genetic operators [Towards an Optimal Mutation Probability for Genetic Algorithms]. Their approach negates the need for determining the best parameters, but adds to the complexity of the problem. This was an important consideration for our approach.

Other unconventional attempts to improve the selection of GA operations have been made over the last few decades. Since GA are inspired by real life biological behaviors, adjustments tend to be based on adding augmentations describing natural phenomena to improve the GA performance. One major example of this is Adaptive Genetic Algorithms (AGA). AGA attempts to adjust the genetic operations ac-

tively throughout the evaluation of the GA to escape local optima and reach a maximum (or minimum) faster. Typical examples of AGA use a defined measurement of some performance metric of the population, such as the population's mean fitness, to determine when to adjust genetic operators [Genetic Algorithms and Directed Adaptation] [Adaptive probabilities of crossover and mutation in genetic algorithms]. Because of this adaptability, the use of AGA requires less *a priori* knowledge of the search space. Adaptation means that in some examples, as fitness increases, mutation rates decrease, or *vice-versa* [Genetic Algorithms and Directed Adaptation] [Adaptive probabilities of crossover and mutation in genetic algorithms].

In one specific example, Coyne and Paton only changed the mutation rate between two values; a 'hypermutable' (high) rate, and normal (typical) mutation rate [Genetic Algorithms and Directed Adaptation]. Use of the population fitness at large provides the impetus to switch between the two values. Srinivas and Patnaik provide a similar case for adaptive mutation rates. However, their case study's genetic operators are not restricted to two values. Rather, they are calculated to suit the situation [Adaptive probabilities of crossover and mutation in genetic algorithms]. A different approach to AGA is described by Tuson and Ross, in which each individual chromosome in a population has its own defined genetic operators. The genetic operators will be adapted and changed during the creation of the next generation [Adapting Operator Settings in Genetic Algorithms].

Ye, Li and Xie give a fundamental insight into adaptive genetic algorithms by comparing two major, contradictory processes: mutation-first, and crossover-first. Mutation-first adaptation relies on a high mutation rate and a low crossover rate initially. Over time, it will progress to the opposite - a low mutation rate and a high crossover rate. Crossover-first is the exact opposite from the beginning, with a low mutation and a high crossover rate initially. It was found that the mutation-first method outperformed the crossover-first [Some improvements on adaptive genetic algorithms for reliability-related applications]. This can be attributed to the need to explore the search space early in the execution (through mutation) and to make smaller adjustments (through increased crossover operations) later on [Some improvements on adaptive genetic algorithms for reliability-related applications]. An attempt was also made to provide a plausible methodology toward solving the local optimum convergence problem by introducing randomly generated individuals to the population. The intention was to increase diversity [Some improvements on adaptive genetic algorithms for reliability-related applications] and the approach with some success.

Our work seeks to provide an explanation for the behavior of our approach by utilizing these works in conjunction with an empirical study and a novel conceptual model for understanding the behavior of the algorithm.

## 2. OUR SOLUTION APPROACH

Ordinary genetic algorithms use three fundamental parameters to govern their evaluation. The first is the population size. We have not adjusted this parameter in our experiments, and kept them at a fixed value of 64 chromosomes as suggested by [CITE TEXTBOOK]. Second is the chromosome crossover rate, which governs the likelihood and percentage of a chromosome to splice with another. This

was also kept static, at a fixed rate of 70% [CITE TEXTBOOK]. The third parameter is mutation rate, which governs the likelihood that a chromosome will have a random gene modified. This is achieved using a randomization operation on the selected chromosome's encoded values.

To support the 'in-breeding' variant of the algorithm, we have selected to replace the fixed mutation rate parameter with a variable rate, governed by a separate parameter which was kept at a fixed value for the sake of minimizing variability in our experimental methodology. The new parameter, chromosome similarity threshold, is expressed as a percentage. We have selected 15% as an arbitrary threshold which seemed to work well during initial experimentation. Future studies into the nature of this parameter may yet yield improved results. The offset by which the mutation rate is adjusted (based on the chromosome similarity parameter) is set at a fixed value of 1%.

## 2.1 Applying Adaptive Variable Mutation Rates

In order to apply our adaptive variable mutation rate, a two step process is required. For each generation of chromosomes, the population's chromosome similarity must be evaluated. The similarity algorithm is detailed in Algorithm 1.

When the chromosome similarity is less than the specified threshold, the population is dissimilar, and inbreeding does not come into play. This means the mutation rate must decrease, because the population is not inbred. For the next generation, the mutation rate is decreased by the fixed value of 1%. The next generation is created and their chromosome similarity is calculated. If the chromosome similarity has increased again, and it is above the specified 15% threshold, the population is inbred. The mutation rate must increase.

Due to the stochastic nature of the algorithm, variations in chromosome similarity may result regardless of the mutation rate. Since the selected crossover rate of the genes is 70%, the resultant permutations of the genome are likely to be approximately 30% similar (due to cloning) unless mutation occurs on cloned chromosomes. The result of this adaptive change in mutation rate is that the chromosome similarity will approach an equilibrium close to the specified inbreeding threshold over generations.

### 2.1.1 Chromosome Similarity Algorithm

Chromosome similarity is calculated using a greedy algorithm seen in Algorithm 1. First, the chromosomes' genes are decoded and re-encoded into integer values, concatenated into one large integer value. next, the array of chromosomes' concatenated integer representations is sorted. Our implementation uses the quick-sort algorithm provided by Java for minimal computational overhead. This gives the algorithm its' characteristic asymptotic behavior, as the complexity of the actual sorting function itself is of higher complexity than the main similarity calculation algorithm. Quick-sort runs on average in  $O(n \log n)$  complexity, although it should be noted that in the theoretical worst case, quick-sort works in  $O(n^2)$  which is highly unlikely. The main algorithm runs in  $O(n)$  linear complexity independent of the sorting.

## 2.2 Implementation Specific Considerations

In order to detail how our specific implementation might compare to alternatives, we will elaborate upon the special implementation considerations we made. Our production

```

Function Similarity(chromosomes)
  input : An array of chromosomes
  output: Fraction of chromosomes that are similar

  similar  $\leftarrow$  0
  matched  $\leftarrow$  false
  length  $\leftarrow$  length of chromosomes

  // Sort using an arbitrary sorting algorithm
  sorted  $\leftarrow$  Sort(chromosomes)

  for  $i \leftarrow 0$  to length - 1 do
    if sorted[ $i$ ] == sorted[ $i + 1$ ] then
      similar  $\leftarrow$  similar + 1
      matched  $\leftarrow$  true
    else if matched then
      similar  $\leftarrow$  similar + 1
      matched  $\leftarrow$  false
    end

    // Case where the last item is a match
    if matched and ( $i + 1$  == length - 1) then
      similar  $\leftarrow$  similar + 1
    end
  end

  return similar / length
end

```

**Algorithm 1:** Chromosome similarity function

experiments were performed our development code, which was written in Java. Concurrent agile development was used as our main software development methodology, and GitHub was used extensively for version management and concurrent development of the source, documentation, and this report.

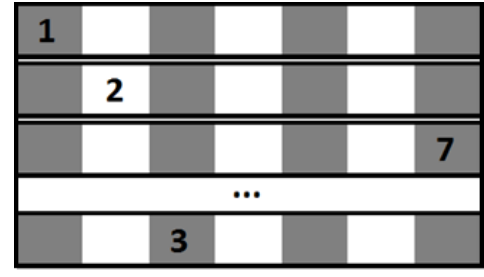
The project source code is available at <http://www.github.com/gnu-user/genetic-algorithms-research>

### 2.2.1 Chromosome Design

Chromosomes are designed with an optimization that allows the solution landscape to be minimized significantly, as detailed in section 1.1.2. The result is that the dimensionality of the problem decreases. Our encoding allows for an extendable chromosome with up to  $N$  genes. Figure 1 shows this chromosome pictorially for a  $7 \times 7$  chessboard. In the figure, the 4th, 5th and 6th rows are not shown, but are filled with a queen at the 4th, 5th and 6th column positions respectively. As a result, the encoded chromosome is represented by 7 Java Integer values in sequence. To encode this into one mutable array, the integers are simply concatenated together. The array specified in Figure 1 would thus be translated to 1274563.

### 2.2.2 Fitness Function

Fitness is evaluated by determining the number of collisions between queens on the chessboard - when at least two queens can attack each other, a collision occurs for each queen. This means that in a board state where one queen can attack another, two collisions result. Because it is impossible for there to be more than one queen per column given the chromosome's encoded data structure, only collisions across rows and diagonals must be calculated to determine fitness. The algorithm for determining the fitness of a given board state is given in Algorithm 2, showing the mechanisms we used to find collisions across rows and diagonals.



**Figure 1:** Illustrating the chromosome data structure.

```

Function Fitness(chromosome)
  input : A single chromosome
  output: A fitness value for the chromosome

  collisions  $\leftarrow$  0
  length  $\leftarrow$  length of the chromosome

  for  $i \leftarrow 0$  to length - 1 do
    // Check each gene against the current
     $j \leftarrow (i + 1) \bmod \text{length}$ 
    while  $j \neq i$  do
       $y_i \leftarrow \text{chromosome}[i]$ 
       $y_j \leftarrow \text{chromosome}[j]$ 

      // Check for vertical collision
      if  $y_i == y_j$  then
        collisions  $\leftarrow$  collisions + 1
      end

      // Check for diagonal collision
      if  $\text{abs}((i - j) / (y_i - y_j)) == 1$  then
        collisions  $\leftarrow$  collisions + 1
      end

       $j \leftarrow j + 1$ 
       $j \leftarrow j \bmod \text{length}$ 
    end
  end

  if collisions == 0 then
    return 1
  else
    return 1 / collisions
  end
end

```

**Algorithm 2:** Fitness function

The overall board state fitness is calculated as  $\text{fitness} = \frac{1}{C}$ , where  $\text{fitness} = 1$  if  $C = 0$ . The number of collisions is  $C$ . Maximum fitness is achieved when  $C = 0$  as there are no collisions, resulting in a unique board solution. The length of the chromosome corresponds to the number of queens and rows on the square chessboard,  $N$ . As a result, the fitness function must check across  $N - K$  rows and diagonals for each queen, where  $K$  is the column index of the current queen.

### 2.2.3 Selection Method

As in traditional GA, roulette wheel selection was used[CITE]. This process works by filling the roulette wheel with fractional parts of a cumulative probability distribution. In our

implementation the range of values will lie between 0 and the sum of all the fitness values of each chromosome in a generation.

Each chromosome is given a fraction of the roulette wheel, whose size is equal to the chromosome's fitness divided by the total fitness of all values. A random floating point value is generated between zero and the sum of the total fitness values. This floating point value will lie within one of the intervals representing one of the chromosomes. Whichever interval it falls in is the selected chromosome. The order in which the chromosomes' intervals are placed doesn't matter, as each 'spin' of the roulette wheel is random - the floating point number generated is created based on a uniformly random distribution.

#### 2.2.4 Chromosome Operations

As mentioned in section 2, the crossover rate is set at a fixed 70%. This means that 30% of the time, the genes in the child chromosomes will be an exact clone of the parents. The remaining times, a random crossover occurs. This is done using the traditional GA approach by selecting a random index as the bounds for the portion of the chromosome to cross over. All genes to the right of this index (including the index value itself) are swapped between parents to generate offspring. Between the two parents, the first parent's values to the right of the index, and the second parent's values to the left of the index are combined to create one child (and *vice-versa* for the second child).

Background mutation affects chromosomes randomly after all crossover operations take place. Each next-generation chromosome has a percentage chance of being affected by mutation, and this likelihood changes every generation due to the adaptive variable mutation rate. When a chromosome is selected for mutation, one of the genes in the chromosome is selected randomly. It is then set to a random valid integer. Valid values are between 0 and  $N$  corresponding to the index of each row on the chessboard.

#### 2.2.5 Chromosome Evaluation

Each generation, there is a likelihood that some of the chromosomes in the current population are valid solutions - any of the current generation's chromosomes with a fitness of 1 is some sort of solution. Whether it is unique or not is unknown at this stage. When a solution is found, it must be compared to the list of previously found solutions to ensure it is not a duplicate. Due to the stochastic nature of the algorithm, there is always a strong possibility that many solutions generated are duplicates of previously found solutions. The symmetry operations applied to each solution found may further increase the likelihood that the new population's solutions that have already been found. Duplicate solutions are discarded without having symmetry operations applied as they would already be identical to symmetric configurations of previously found solutions.

Upon saving any newly found solutions, the new generation replaces the previous generation and the genetic algorithm process cycle continues again.

### 2.3 Experimental Methodology

Experiments were conducted using the high performance computing facilities provided by SHARCNET. Testing of our process was performed by using two different configurations for each  $N$  value of the  $N$ -Queens problem. Tests were per-

formed for  $N = 8, 9, 10$  and so on up to 16, and then at increasing increments of 2 from 18 to 20, 22, 24, 26 and 32. A fixed number of generations was allowed for each trial in order to ensure that the tests had a finite and equal running time. For all runs, 10 million generations were allowed, except for  $N = 32$ , where 50 million generations were allowed given the increased complexity of the problem.

The first configuration used static mutation rates set at  $m = 1\%, 5\%, 10\%$ , and every further increment of 5% up to 100%. These were used as a control. The second configuration used the adaptive variable mutation rate, and the results were compared.

For  $N = 8$  through 16, 30 tests were performed at each static mutation rate ( $21 \times 30 = 630$  tests total), and then 30 runs were performed using variable mutation rates.

For  $N = 18$  through 26, 15 tests were performed under both configurations. For  $N = 32$ , 10 tests were performed. The decrease in the number of tests performed for higher values of  $N$  was largely due to the running time required to generate meaningful results. SHARCNET provisioned us to execute a maximum of 256 jobs at one time, with 7 days CPU time each. The 32-Queens problem required this limit to be fully utilized in order to give results. Given the intractable size of the higher-order  $N$ -Queens problems beyond 32-Queens, an unsustainable amount of time and CPU power would be required to finish these problems within our time constraints and were not tested.

Given the volume of data generated by the number of trials, statistical attributes of the results collected (such as mean population fitness and chromosome similarity) were calculated using the mean of every 1000 generations in order to observe the central tendency. A total of roughly 6 gigabytes of data was generated as a result of this process. This was manageable given our allocation of storage space on SHARCNET. The requirements for collecting the data at every generation would have been 1000 times that volume (approximately 6 terabytes) which far exceeded our allocation.

### 3. RESULTS AND ANALYSIS

It became clear very quickly when we plotted the results of our tests that the variable mutation method was not suitable in its current configuration for the simpler N-Queens problems. In fact, we found that given sufficient knowledge about the problem landscape for the lower-order problems, the right static mutation value would likely beat variable mutation every time. However, for ordinary non-benchmark usage of genetic algorithms it is useless to use a static mutation rate value unless you already know the best mutation rate for the problem. This generally speaks to the point that having an adaptive mutation rate approach will likely find this value. The smaller N-Queens problems are too small to merit the usage of a stochastic approach regardless, as deterministic methods can yield solutions quickly with guaranteed success.

#### 3.1 'Traversing' the Problem Landscape

Moving into more complex and larger problems is where the adaptive mutation approach becomes increasingly successful. Our reasoning is that this may have something to do with the similarity threshold value that was chosen (15%). Further tuning of this parameter may yield better results for larger or smaller problems. We have created a conceptual model for our findings which may yet be validated in future studies.

Mapping the stochastic solution landscape for the N-Queens problem means plotting a random non-deterministic traversal of the solution space to a linear axis. Each sequential value on this axis corresponds to each integer sequence of solutions encoded in the chromosomes. A linear traversal of this landscape implies a uniformly random traversal of the randomly generated solutions. This plot, in contrast to the deterministic solution space yielded by consecutively plotting the values of the integer encoded chromosome, shows very few similarities. There is no way to plot a random traversal except by noting that as time approaches infinity, all solutions will have been traversed and the full contour of the landscape will be visible. This could just as easily be yielded by a consecutive linear traversal, but the point of using GA rather than a deterministic traversal is that GA makes the traversal 'smarter' by searching for values which have better fitness, attracted toward fitness peaks[CITE]. The mutation operator helps to prevent the current position in the traversal from being stuck in a local optimum[CITE]. It provides an extra impulse to shift position further towards what a potential solution, and away from previously traversed areas of the landscape.

Given this knowledge, assuming a linear traversal of this random axis is possible in one given test, we can evaluate the characteristics of each GA's mutation parameter configuration and map them to this conceptual model. The starting point on the plot is a random location on the landscape. From there, the next location is likely to be within a radius of probable new chromosomes surrounding this area. We have come to refer to this radius of probable chromosomes as the 'step size', and we believe there is some relationship between the step size and the fixed mutation rate. The fixed mutation rate governs the maximum distance from the current position in the traversal to the furthest step away. What this means for a solution landscape like that of the N-Queens problem is that all solutions which are close to the intersections of these steps are easier to find because there

is a greater total probability of landing within their vicinity. Solutions which are clustered very close together on this random landscape, however, will likely be stepped around for several generations before convergence on the actual solution occurs.

The variable mutation rate's adaptive nature allows for this fixed step size to be ignored because it introduces the ability to vary the step size when convergence upon a solution occurs. The result is that the probability distribution over the entire solution landscape is more uniformly smaller away from solutions, and larger within the vicinity of solutions. Our reasoning is that if the static mutation rates have a fixed step size, the variable mutation rate does not, and adaptation will vary the step size so as to maintain the chromosome similarity. Further, this decreases the likelihood of overshooting a solution and having to step back around again, because the variability of the mutation rate allows for smaller or larger steps when needed.

In general, this conceptual model will require extensive further study to validate it's correctness. Meanwhile, the empirical evidence presented in our plotted results shows the validity of our adaptive mutation approach.

#### 3.2 Plotted Results and Observations

- cite specific results and why we think they are important, what is the significance of them and how they support our results/hypothesis that in the case of N Queens var. mutation is better than fixed.

There appears to be a relationship

- cite the result showing the best fixed mutation vs. the variable mutation for each N-queens problem, try and use both the figure and the table, the table has some additional interesting information which cannot be conveyed in the image alone.

An interesting result was observed when viewing the box plots showing the range of chromosome similarity value for each static mutation rate. We found that whenever the similarity has the greatest inter-quartile range, the corresponding static mutation rate performed best. This contributed heavily to the step size concept we believe is exhibited by the system. We also found that as the problem landscape grows larger in higher order N-Queens problems, the inter-quartile range decreases in size. There is a possibility that the convergence of these values may occur in even higher order problems, which suggests that there may be a pattern to the convergence which can be approximated and leveraged for predicting optimal static mutation rates. Again, since this requires exhaustive experimentation in order to definitively find the optimal static mutation rate for a given problem, this further supports the notion that a variable mutation rate is a more practical approach.

- Try and incorporate one of the scatter plots as well that show very interesting results and see if you can use it to compare/contrast the results of the following plots

- variable mutation rate scatter plot - variable mutation rate similarity scatter plot - best fixed mutation rate similarity scatter plot

#### 3.3 Why Not Fixed Mutation?

Results aside, fixed mutation gives a distinct advantage over traditional genetic algorithms because of the optimization of a specific parameter [CITE HERE]. By using a variable mutation rate, changing that parameter with a separate

simpler one, better results can be gleaned while knowing less about the problem landscape. Our results show that our variable adaptive mutation scheme can perform better than fixed static mutation values for large problems. As the problem becomes too large and unmanageable for traditional methods, our methodology can still produce useful results. Our results show that even having knowledge about the best static mutation value to use *a priori* still does not yield results as good as those generated from the adaptive variable mutation rate.

## 4. CONCLUSIONS AND FURTHER STUDY

- Further research into adjusting the inbreeding threshold, for the purpose of the research a constant inbreeding threshold of 15% was used, however further research could be done in testing different thresholds.
- Comparing the results of variable mutation with fixed mutation using other types of combinatorial/optimization problems such as TSP, constraint satisfaction problem (CSP), etc.
- Variable population size based on the amount of inbreeding (if you have a lot of inbreeding in nature the organisms will have higher mutation rate and deformities, the population will shrink)
- Having the mutation operator affect more than one gene if it goes > 100%?
- Cite [Some improvements on adaptive genetic algorithms for reliability-related applications] and discuss their findings compared to ours.

## 5. ACKNOWLEDGMENTS

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## APPENDIX

### A. RESULTS

The rules about hierarchical headings discussed above for the body of the article are different in the appendices. In the **appendix** environment, the command **section** is used to indicate the start of each Appendix, with alphabetic order designation (i.e. the first is A, the second B, etc.) and a title (if you include one). So, if you need hierarchical structure *within* an Appendix, start with **subsection** as the highest level. Here is an outline of the body of this document in Appendix-appropriate form:

#### A.1 Tables

**Table 1: Number of unique solutions to the N-Queens Problem**

N	Distinct Solutions
1	1
2	0
3	0
4	2
5	10
6	4
7	40
8	92
9	352
10	724
11	2680
12	14,200
13	73,712
14	365,596
15	2,279,184
16	14,772,512
17	95,815,104
18	666,090,624
19	4,968,057,848
20	39,029,188,884
21	314,666,222,712
22	2,691,008,701,644
23	24,233,937,684,440
24	227,514,171,973,736
25	2,207,893,435,808,352
26	22,317,699,616,364,044
27	Unknown
...	Unknown

#### A.2 Figures

- Additional figures can go here

**Table 2: Best Solution Comparison Between Fixed and Variable Mutation Rates**

Queens	Mutation Rate	Unique Solutions	N	Min Generations	Mean Generations	Max Generations
8	0.95 variable	92	30	1,793	4,746	10,815
		92	30	51,23	15,293	42,002
9	0.95 variable	352	30	48,811	144,456	372,281
		352	30	229,242	1,246,215	3,038,980
10	0.9 variable	724	30	188,372	408,402	771,208
		724	30	531,588	1,629,893	6,195,099
11	0.85 variable	2,680	30	2,400,188	3,799,795	5,933,213
		2,680	5	6,900,512	7,909,269	9,449,545
12	0.85 variable	13,690	1	9,890,455	9,890,455	9,890,455
		11,986	1	9,973,787	9,973,787	9,973,787
13	0.8 variable	32,128	1	9,998,058	9,998,058	9,998,058
		26,308	1	9,978,215	9,978,215	9,978,215
14	0.8 variable	41,520	1	9,996,747	9,996,747	9,996,747
		29,520	1	9,999,583	9,999,583	9,999,583
15	0.8 variable	30,356	1	9,982,638	9,982,638	9,982,638
		30,324	1	9,996,584	9,996,584	9,996,584
16	0.8 variable	15,016	1	9,987,788	9,987,788	9,987,788
		30,132	1	9,992,266	9,992,266	9,992,266
18	0.75 variable	16,392	1	9,997,986	9,997,986	9,997,986
		27,120	1	9,998,858	9,998,858	9,998,858
20	0.75 variable	7,872	1	9,972,915	9,972,915	9,972,915
		25,608	1	9,931,631	9,931,631	9,931,631
22	0.7 variable	6,008	1	9,959,712	9,959,712	9,959,712
		25,376	1	9,991,089	9,991,089	9,991,089
24	0.7 variable	6,440	1	9,929,626	9,929,626	9,929,626
		24,560	1	9,998,508	9,998,508	9,998,508
26	0.7 variable	6,280	1	9,969,840	9,969,840	9,969,840
		23,008	1	9,983,128	9,983,128	9,983,128
32	0.65 variable	15,216	1	49,732,320	49,732,320	49,732,320
		104,080	1	49,996,701	49,996,701	49,996,701



### **A.3 Source Code**

- Could link to github and possibly the github wiki, or explanation of the R analysis source code.

### **A.4 Raw Data**

- Link to the dropbox URL containing the actual data from SHARCNET used for the research