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# Multiniche Crowding in Genetic Algorithms and Its Application to the Assembly of DNA Restriction-Fragments

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## Abstract

The determination of the sequence of all nucleotide base-pairs in a DNA molecule, from restriction-fragment data, is a complex task and can be posed as the problem of finding the optima of a multimodal function. A genetic algorithm that uses multiniche crowding permits us to do this. Performance of this algorithm is first tested using a standard suite of test functions. The algorithm is next tested using two data sets obtained from the Human Genome Project at the Lawrence Livermore National Laboratory. The new method holds promise in automating the sequencing computations.

## Keywords

Genetic algorithms, multimodal functions, DNA restriction-fragment assembly, human genome project

## 1. Introduction

This paper describes how the concept of multiniche crowding in a genetic algorithm permits one to find several extrema of a multimodal function simultaneously. Although a complete theoretical analysis supported by performance data on the method is still being worked out, initial results from experiments with a standard suite of test functions are persuasive about the utility of this method to solve problems of practical interest. The method is tested by applying it to solve a problem in molecular biology, believed to be NP-hard.

Section 2 of this paper describes multimodal function optimization using the new genetic algorithm (GA) model. The validity of the new method is demonstrated in Section 3 by applying it on several multimodal test functions. Section 4.1 gives a tutorial background on the Human Genome Project and the relevance of the restriction-fragment assembly

problem. Section 4.2 summarizes the experimental procedure used by one group of biologists to gather the data used in this paper. Section 4.3 summarizes some of the related work on restriction-fragment map assembly. Section 5 addresses the representational issues including a description of the genetic operators and the fitness function used to solve this problem. Section 6 includes results and discussion, and Section 7 summarizes ongoing work.

Because genetic algorithms are being used in this paper to solve a problem in genetics, a word of caution about the terminology used is in order. Words like “chromosomes” occur while describing the genetic algorithm as well as during the description of the DNA fragment assembly process. To the biologist, a chromosome is DNA. To the computational scientist, a chromosome is a data structure. The context will make the intent abundantly clear.

## 2. Genetic Algorithms for Multimodal Search

Searching for extrema in a multimodal search space is different from locating the extremum of a unimodal function. When a search technique proven to be useful for unimodal functions is applied to multimodal functions, the method tends to converge to an optimum in the local neighborhood of the first guess. One can use ideas like simulated annealing to escape from a local optimum. However, there are many applications where the location of “k best extrema” of a multimodal function is of interest. Searching for these locations goes by the name “multimodal optimization.”

There are many versions of genetic algorithms, one differing from another in some detail. For the purposes of this paper, it is sufficient to focus on two basic steps common to all genetic algorithms. During the *selection step*, a decision is made as to who is allowed to produce offspring, and during the *replacement step* another decision is made as to which of the members from generation  $i$  are forced to perish (or vacate a slot) to make room for an offspring.

The simple GA (or SGA) (Holland, 1975), which will be used as a point of departure for presenting our method, starts by randomly generating a population of  $N$  individuals, that is, individual solutions. These individuals are evaluated for their fitness. Individuals with higher fitness scores are selected, with replacement, to create a mating pool of size  $N$ . This method of selection is called *fitness proportionate reproduction* (FPR). The genetic operators of crossover and mutation are applied at this stage in a probabilistic manner, which results in some individuals from the mating pool reproducing. The assumption here is that each pair of parents produce only one pair of offspring through the crossover operation. That is, the second generation is comprised of those first-generation individuals who never got a chance to reproduce and the offspring of those who got a chance to reproduce. The procedure continues until a suitable termination condition is satisfied, say a specified number of generations. (For the purposes of this paper, a generation is every  $N/2$  mating operations, where  $N$  is the population size.) All other versions of GAs differ from this basic method in some detail or another.

The steady-state GA (SSGA) (Whitley, 1988; Syswerda, 1989), differs from SGA mainly in the replacement step and, to a lesser extent, on the way the genetic operators are applied. The SSGA selects two individuals using FPR and allows them to mate to produce two offspring. This selection step is identical to the corresponding step of SGA. However, in SSGA, the offspring are inserted into the population, thus replacing two individuals, soon after they are generated. The SGA, on the other hand, generates  $N$  offspring prior to replacing the entire population. In other words, SGA uses *simultaneous* replacement strategy whereas the SSGA uses *successive* replacement strategy. Thus, SGA and SSGA

are analogous, respectively, to the Jacobi and Gauss-Seidal methods of solving systems of algebraic equations.

Both SGA and SSGA suffer from the possibility of premature convergence to a local minimum, primarily due to the *selection pressure* exerted by the FPR rule. Simply assigning an exponential number of mating opportunities to those members of the population that exhibit above-average survival traits, as the FPR rule does, is not a good strategy for a thorough exploration of complex search spaces with multiple peaks. Due to this reason as well as the *deceptiveness* exhibited by multimodal search spaces (Goldberg, Deb, & Horn, 1992), SGA and SSGA are not suitable for multimodal search and optimization.

## 2.1 Multimodal Search: Crowding-based Methods

The GA model described in this paper, called *multiniche crowding* (MNC), has the ability to converge simultaneously to multiple solutions by encouraging competition between individuals within the same locally optimal group (Cedeño and Vemuri, 1992). In MNC, both the selection and replacement steps of the SGA are modified with the introduction of some form of *crowding* in order to render it suitable for searching spaces characterized by multiple peaks or niches. For completeness, the concept of crowding is briefly reviewed here.

Crowding (De Jong, 1975) is a generalization of *preselection* (Cavicchio, 1970). In *crowding*, selection and reproduction are the same as in the SGA; replacement is different. For concreteness, it is assumed that two parents are selected to produce two offspring. In order to make room for these offspring, it is necessary to identify two members of the population for replacement. The policy of replacing a member of the present generation by an offspring is carried out, as follows, in two steps. First, a group of  $C$  individuals is selected at random from the population.  $C$ , called the *crowding factor*, also indicates the size of the group. A value of  $C = 2$  or  $3$  appears to work well for De Jong. Second, the bit strings in the offspring chromosomes are compared with those of the  $C$  individuals in the group using Hamming distance as a measure of similarity. The group member that is most similar to the offspring is now replaced by the offspring. This procedure is repeated for the other offspring as well. This second offspring can conceivably replace its own sibling that just entered the population pool, although such a scenario is rather unlikely. In any event, crowding is essentially a successive replacement strategy. This strategy maintains the diversity in the population and postpones premature convergence. However, crowding cannot maintain stable subpopulations too long due to the selection pressure imparted by FPR. Summarizing, in crowding offspring replace similar individuals from the population. Crowding slows down premature convergence of the traditional GA and in most cases can find the global optimum in a multimodal search space. On the other hand, it does not facilitate convergence to multiple solutions, and after many generations one of the peaks takes over.

In *deterministic crowding* (Mahfoud, 1992) selection pressure is eliminated and *preselection* is introduced to obtain a very fast GA suitable for multimodal function optimization. In this method selection pressure is eliminated by allowing individuals to mate at random with any other individual in the population. Pressure is applied, however, during the replacement step using preselection. Toward this goal, each of the two offspring is first paired with one of the parents. This pairing is not done randomly; rather, the pairings are done in such a manner that the offspring is paired with the more similar parent. Then each offspring is compared with its paired parent; the individual with the higher *fitness* is allowed to stay in the population and the other is eliminated. It is not clear if multiple solutions can be maintained for many generations using this method, although it appears that multiple solutions are sustained for more generations than when crowding is used alone.

## 2.2 Multimodal Search: Multiniche Crowding

In *multiniche* crowding (MNC), both the selection and replacement steps are modified with some type of crowding. The idea is to eliminate the selection pressure caused by FPR while allowing the population to maintain some diversity. This objective is achieved, in part, by encouraging mating and replacement within members of the same niche while allowing for some competition for population slots among the niches. The result is an algorithm that (a) maintains stable subpopulations within different niches, (b) maintains diversity throughout the search, and (c) converges to different local optima.

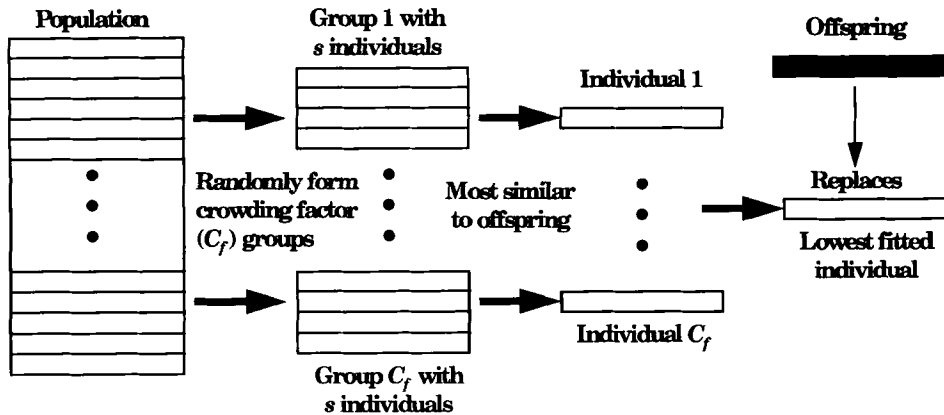
In MNC, the FPR selection is replaced by what we call *crowding selection*. In crowding selection, each individual in the population has the same chance for mating in every generation. Application of this selection rule takes place in two steps. First, an individual A is selected for mating. This selection can be either sequential or random. Second, its mate M is selected, not from the entire population but from a group of individuals of size  $C_s$ , picked at random (with replacement) from the population. The mate M thus chosen must be the one who is the most “similar” to A. The similarity metric used here is not a *genotypic metric* such as the Hamming distance, but a suitably defined *phenotypic distance metric*. Crowding selection promotes mating between individuals from the same niche while allowing matings between individuals from different niches.

During the replacement step, MNC uses a replacement policy called *worst among the most similar*. The goal of this step is to pick an individual from the population for replacement by an offspring. Implementation of this policy follows these steps. First,  $C_f$  groups are created by randomly picking  $s$  individuals (with replacement) per group from the population. These groups are called *crowding factor groups*. Second, one individual from each group that is most phenotypically similar to the offspring is identified. This gives  $C_f$  individuals that are candidates for replacement by virtue of their similarity to the offspring that will replace them. From this group of most similar individuals, we pick the one with the lowest *fitness* to die, and that slot is filled with the offspring. The offspring could possibly have a lower fitness than the individual being replaced.

Both selection and replacement steps in the MNC are primarily based on a similarity metric. Fitness is also considered during replacement to promote competition between members of the same niche. Competition between members of different niches occurs as well. The following pseudocode summarizes the salient features of the MNC:

1. Generate initial population of  $N$  individuals
2. For gen = 1 to MAX\_GEN
3.     For  $i = 1$  to  $N$
4.         Use *crowding selection* to find mate for parent  $i$
5.         Mate and mutate
6.         Insert offspring in the population using *worst among most similar* replacement.

If FPR is used in Step 4 above and if the individual with the lowest fitness in the population (Step 6) is replaced with the newly generated offspring, this model corresponds to a steady-state GA. In contrast with the most common generational GA, offspring are available for mating as soon as they are generated, and good individuals can survive for many generations. A similar technique called *enhanced crowding* (Goldberg, 1989) has been used



**Figure 1.** Schematic showing worst among most similar replacement policy.

before in classifier systems, but there the most similar individual out of a group of worst candidates is replaced. A pictorial interpretation of the MNC replacement policy is shown in Figure 1.

In summary, MNC differs from the other crowding-based methods in the following salient features:

1. Both selection and replacement use a form of crowding.
2. Mating among members of the same niche is encouraged.
3. Competition among members of the same niche is promoted.

These features appear to be responsible for the apparently superior performance of this method over other crowding-based methods insofar as maintaining solutions in multiple peaks. The next section shows some performance results when this method is applied to some standard test functions.

### 3. Application of MNC to Multimodal Test Functions

To evaluate the performance of the MNC model described above, we used five different test functions. The first three of these functions have been used earlier by other investigators. Functions  $F_1(x)$  and  $F_2(x)$ , defined by

$$F_1(x) = \sin^6(5.1\pi x + 0.5)$$

and

$$F_2(x) = \exp^{-4(\ln 2)(x-0.0667)^2/0.64} \sin^6(5.1\pi x + 0.5)$$

are shown in Figure 2. These functions correspond to the five-optima sine functions used by Goldberg and Richardson (1987) in their work on sharing. Both functions were maximized using binary chromosome encoding for numbers in the interval  $[0, 1]$ . In both cases, the

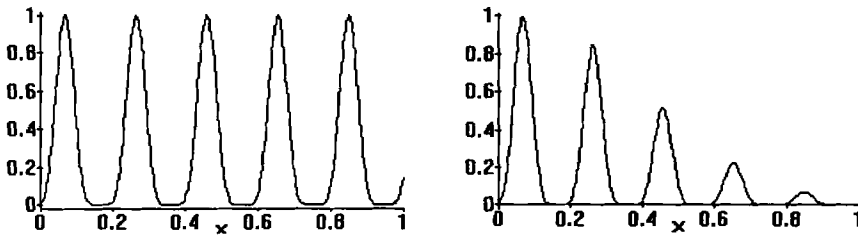


Figure 2. Test functions  $F_1(x)$  (left) and  $F_2(x)$  (right).

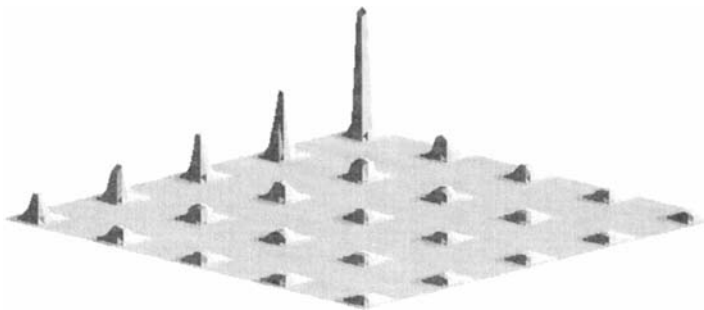


Figure 3. Test function  $F_3(x)$ , inverted Shekel's foxholes.

GA with sharing was able to maintain stable subpopulations and diversity in the population. Convergence was good, but not all the peaks in  $F_1$  had the individuals distributed close to the top. It is not clear that in  $F_2$  sharing will be able to maintain proportional numbers of individuals for larger numbers of generations.

Function  $F_3(x, y)$ , called “Shekel's foxholes” with 25 optima, was used by De Jong in his work on crowding. The inverted version of  $F_3$  is shown in Figure 3. The peaks are located on the grid intersections formed by the values  $x = [32, 16, 0, -16, -32]$  and  $y = [32, 16, 0, -16, -32]$ . All 25 peaks have the same base width. The height of peak  $i$  is given by the expression  $0.002 + 1/i$ , where the peaks are located at  $(32, 32), (16, 32), \dots, (-32, -32)$ . Using a value of  $C = 2$  a GA using crowding consistently found the global optima. Crowding allows the GA to escape local optima by maintaining diversity in the population during the early stages.

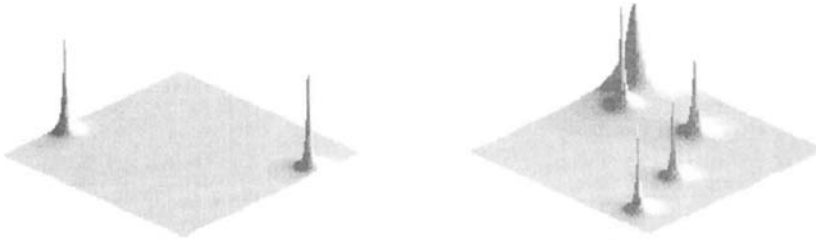
We also considered other functions not exhibiting the symmetry present in the above functions. Function  $F_4(x, y)$ , shown on the left in Figure 4, contains two global optima with the same height and width but located far apart. Function  $F_5(x, y)$ , the sample shown on the right in Figure 4, contains five optima with height, width, and location chosen at random in every run. Both of these functions are defined by the summation:

$$\sum_{i=1}^p \frac{A_i}{1 + W_i((x - X_i)^2 + (y - Y_i)^2)},$$

where  $p$  indicates the number of peaks in the function,  $(X_i, Y_i)$  the coordinates of peak  $i$ ,

**Table 1.** Parameters for functions  $F_4$  and  $F_5$ .

Function	Peak Location	Width	Height
$F_4$	(45k, 2k)	0.0004	100
	(15k, 62k)	0.0004	100
$F_5$	(17.1, 34.4)	1.9	50.5
	( 8.7, 4.1)	0.17	44.8
	(20.3, 11.0)	3.6	87.1
	(38.3, 47.9)	3.1	51.5
	(57.9, 54.1)	3.7	55.5

**Figure 4.** Test functions  $F_4$  (scaled version on left) and  $F_5$  (right).

$A_i$  is the height of peak  $i$ , and  $W_i$  determines how narrow or wide the base of peak  $i$  is. Table 1 summarizes the parameters for functions  $F_4$  and  $F_5$  shown in Figure 4.

The MNC method was applied on these test functions, and the simulations were done on a 486/33MHz PC. The variables  $x$  and  $y$  were encoded using two 32-bit chromosomes. The initial population was generated at random. *Interval crossover* was used for mating. In interval crossover only one offspring is generated. For each pair of parent chromosomes,  $x_1$  and  $x_2$  (assume without loss of generality that the cardinal value of binary string  $x_1$  is less than the cardinal value of binary string  $x_2$ ), the offspring's chromosome is selected at random from the interval  $[x_1 - \delta, x_2 + \delta]$ . The value for  $\delta$  is usually small. This allows the offspring to move outside the boundaries delineated by their parents. For this test  $\delta$  was set at the decimal value 1,000. We obtained better results using interval crossover over single-point crossover. This is due in part to the disruptive effect on niche preservation caused by single-point crossover. For example, consider the individuals 0100000000000 and 0011111111111. If these individuals are part of the same peak, a single-point crossover may conceivably generate the offspring pair 0000000000000 and 0111111111111, which may not be in the same peak as the parents. On the other hand, interval crossover will generate an offspring close to the parents and more likely to belong to the same niche. After mating bit mutation is applied to the offspring.

The crossover probability ( $p_c$ ) was set at 0.95. Normally one tends to choose a lower value for the crossover probability in order to allow individuals to survive from one generation to the other. In contrast, in the MNC,  $p_c$  is usually chosen in the interval  $[0.9, 1.0]$  for three reasons. First, the individuals with highest fitness will survive multiple generations. Second,

**Table 2.** Function-specific parameters used in the MNC method.

	$F_1$ & $F_2$	$F_3$	$F_4$	$F_5$
Population size ( $N$ ):	100	500	100	200
Number of chromosomes:	1	2	2	2
Crowding selection size ( $C_s$ ):	15	75	5	15
Crowding factor ( $C_f$ ):	2	2	3	3
Crowding factor group size ( $s$ ):	15	75	5	15

low  $p_c$  values tend to increase the number of duplicate chromosomes in the population. Third, offspring are likely to replace individuals with similar phenotypic features and thus alleviate the need to carry other individuals. The mutation probability ( $p_m$ ) was set at 0.001. Mutation may allow an individual to escape from a niche to other areas not explored by the algorithm. Furthermore, it is useful to note, in passing, that duplicates encountered after mating (mating pair do not undergo crossover and mutation) or during replacement (offspring same as individual in population) are not inserted in the population.

The phenotypical similarity between two individuals was determined by the Euclidean distance between them. Like Deb and Goldberg (1989), we also had better performance when the similarity was based on the phenotype rather than the genotype. The MNC method was executed for 100 generations in each run. Other parameters are summarized in Table 2.

These parameters were chosen after a number of trial runs. Although the optimal value of these parameters is unknown at this moment, a rule of thumb that worked fine on these functions was the following. The population size was selected to be  $N \geq 20p$ , where  $p$  is the number of peaks. This value of  $N$  allowed small peaks to survive for many generations, and the need to use a large population is avoided.

For the crowding selection size,  $C_s$ , and the crowding size,  $s$ , the values were chosen to be at least  $2p$ . The idea is to get a group size that will likely contain an individual from the same niche during selection or replacement. At the same time it must be small enough to allow competition between different niches. A large group size will restrict selection and replacement among members of the same niche only. For these test functions a value of  $3p$  for  $C_s$  and  $s$  produced good results. The crowding factor,  $C_f$ , was set at 2 or 3. Higher crowding factor values increased the chances of high-fitness individuals to survive for more generations. On the other hand, higher values can cause niches from small peaks to disappear from the population in fewer generations.

These parameter values seem to balance competition between individuals in the same niche and individuals from different niches. Consider for example a value of  $C_s = 1$ . This will reduce the selection step to basically random selection as is used in deterministic crowding. Increasing the value of  $C_s$  will localize a mate closer to the parent. For very large values (i.e.,  $C_s$  closer to  $N$ ) the mate chosen will most likely be the parent itself.

The same applies during replacement for the value of  $s$ . What is interesting is the effect of the combined values of  $s$  and  $C_f$  during replacement. A value of  $s > 1$  and  $C_f = 1$  reduces the replacement step to crowding, and fitness plays no role in the MNC method. These settings will reduce the method to some form of random search since fitness is not used during selection. Consider now  $s = 1$  and  $C_f > 1$  during replacement. In this case similarity



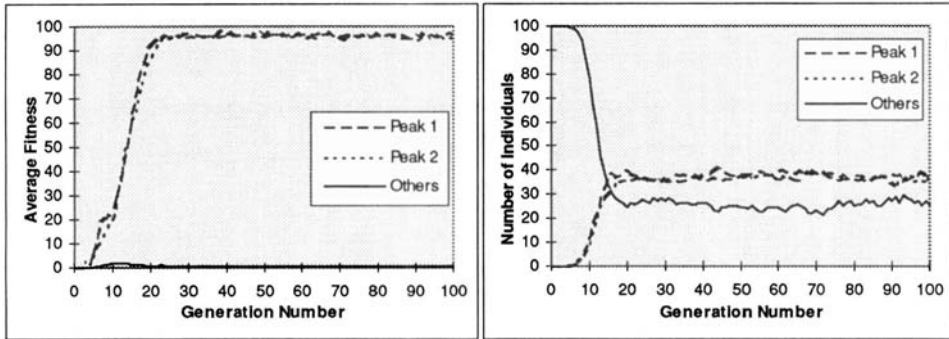


Figure 5. Average fitness (left) and number of individuals (right) for function  $F_4$ .

plays no role, and only fitness is used during replacement. Higher peaks will tend to take over the population with these settings. At a minimum both  $s$  and  $C_f$  must be greater than 1 to allow multiple niches to survive. Increasing  $s$  will promote more competition between members of the same niche. Increasing  $C_f$  will allow fitness to play a bigger role during replacement. An analysis of the MNC method and the effect of these parameters is ongoing and will be reported in another paper.

An interesting approach will be to allow the parameters  $C_s$ ,  $s$ , and  $C_f$  to dynamically change with the population. This will allow the algorithm to adapt itself to the idiosyncracies of the search space. A related idea, based on implicit fitness sharing (Smith, Forrest, & Perelson, 1993), effectively maintained diverse populations while modeling the immune system.

### 3.1 Results on Multimodal Test Functions

For all test cases the MNC method was able to maintain stable subpopulations in most of the peaks without converging prematurely to any one of them. Only very small peaks in  $F_3$  and  $F_5$  did not have any significant number of individuals in the last generation even though some were present during many earlier generations. All peaks in functions  $F_1$ ,  $F_2$ , and  $F_4$  were found. Even the local optima at  $x = 1$  was found in  $F_1$  and was kept until the last generation. Stable subpopulations were consistently maintained in all peaks until the last generation. The distance between the peaks in  $F_4$  did not cause any problem for the MNC method. Figure 5 shows for function  $F_4$  the average fitness of the individuals (on the left) in each peak and the number of individuals (on the right) in each peak. The results were averaged over 20 runs. An individual solution is considered part of a peak if it is located in the region above 10% of the height of the peak. The solid line represents individuals located in neither peak. As can be seen, the number of individuals stabilizes around the same time as the average fitness does. After that the number of individuals in the peaks does not fluctuate greatly for the last 70+ generations.

In general, the number of individuals at the lower peaks decreased as the fitness of the individuals in other peaks increased. During the initial generations all peaks had about the same number of individuals since their average fitness was comparable. As the average fitness increased in some peaks so did the number of individuals in those peaks. In other words, the number of slots in the population occupied by a peak depends on its average fitness relative

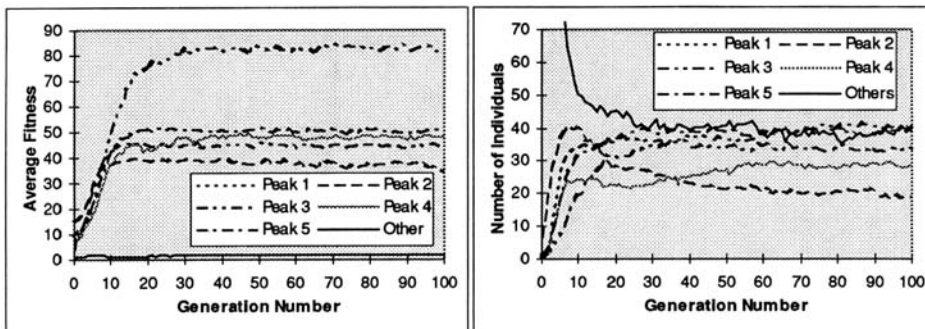


Figure 6. Average fitness (left) and number of individuals (right) for function  $F_5$ .

to that of other peaks. Of course, there are other factors affecting the peak count that need to be studied further. As the average fitness approaches a peak's optimum, the number of individuals in that peak stabilizes. This can be observed in Figure 6 for peak 2 located near  $(0, 0)$  and all other peaks in function  $F_5$ . The plot on the left has the average fitness of each peak for every generation. The plot on the right has the number of individuals in each peak for every generation. During the initial 10 generations the wider peak (peak 2) had more individuals in the population. As the fitness of the individuals in the other peaks improved the number of individuals increased. After about 20 generations the number of individuals in the other peaks was greater than those in the wider peak. There are, of course, other factors that contribute to this pattern that need to be investigated further. It shows the ability of the MNC method to avoid premature convergence.

We also observed that the number of individuals in a peak is related to more than just its average fitness. In function  $F_3$  where the optima are located on a  $5 \times 5$  grid, peaks along the same  $x$  and  $y$  axes as the global optima had more individuals than other peaks with higher average fitness. Some of the extra individuals can be attributed to mutation since a bit change in one of the chromosomes will cause an individual to move along the  $x$  or  $y$  axis. We ran the same test with mutation set at 0.0, and no major changes were observed. More analysis is needed to determine other factors affecting the number of individuals in a peak.

The properties exhibited by the MNC method are encouraging. The population converges to multiple optima, and stable subpopulations evolve in different niches. Three main factors appear to contribute to the success of the algorithm. First, using crowding selection and worst among most similar replacement allowed niches to form naturally and compete for population slots among them. Second, the mating operator used preserved parents' phenotype in the offspring and therefore allowed individuals from a niche to generate offspring in the same niche. Third, phenotypic similarity allowed individuals to form niches in the solution space. Although a more rigorous analysis is necessary before the merits of this method can be substantiated, this method is applied nevertheless to solve the problem of DNA restriction-fragment assembly, a problem widely believed to be NP-hard.

#### 4. Background on the DNA Restriction-Fragment Assembly Problem

The purpose of this section is to provide a simplified summary of the biological background necessary to understand some of the central issues in the DNA restriction-fragment assembly problem as it arises in the Human Genome Project.

The genetic material contained in all the chromosomes of a cell is collectively called the genome. Chromosomes are essentially DNA molecules, the ladder-shaped, double-helical structures. For the purpose of this paper, suffice it to say that the most important parts of this double-helix are the “steps” of the ladder. These steps, called base-pairs, denoted here by the letters A, C, G, T, can theoretically form 16 pairs. However, only AT, TA, CG, and GC pairings are allowed. That is, if the left half of the base-pair is known, the right half is uniquely determined and vice versa. It is estimated that the human genome, contained in all the 23 pairs of chromosomes, is comprised of about three billion base-pairs. A hypothetical sequence of these may look like AATCTTCGGGCCT. . . occupying three billion positions. Specific subsequences of this are called genes.

The monumental task of the Human Genome Project is to (a) associate with each gene all the properties controlled by that gene, (b) associate each gene with one of the 23 chromosomes in the body, (c) identify the exact positioning of a gene on a chromosome—known as the *mapping* problem or the *genetic-linkage* problem—and (d) decipher the exact sequence of base-pairs that constitute a given gene—known as the *sequencing problem*. Although each human is uniquely characterized by his/her genome, apparently one individual differs from another in only a small percentage (about 0.2%) of this material. For that matter, the human genome differs from the simian genome by only a few percentage points. Thus the Human Genome Project is only analyzing a sort of composite genome: 23 chromosome pairs donated by a few European and U. S. scientists. Thus, the focus is on understanding the common structure that runs through the human species, although the actual DNA used in the experiments may come from a specific individual.

Issues related to molecular biology, instrumentation, and computations do play a critical role in this effort. It is impractical even to attempt to summarize the scope of this project here. A description of the technique used at the Human Genome Center of the Lawrence Livermore National Laboratory (LLNL), for example, can be found in *Genomics* 4, pp. 129–36 (Carrano et al., 1989). To view online information about, say, the LLNL genome program, it is only necessary to point the WWW client at <http://www-bio.llnl.gov/bbrp/genome.html>. This Center is mapping human chromosome 19, which is estimated to be approximately 60 million base-pairs long, a relatively small chromosome.

Current technology is forcing us to limit the sequencing task to small fragments of DNA that are composed of approximately 0.5K base-pairs (Istvanick et al., 1993). In order to divide the DNA into fragments up to this resolution level, techniques using restriction enzymes are used. The restriction enzymes act on the DNA at specific locations that are randomly distributed along the length of the chromosome. Depending on the number of different restriction enzymes used in obtaining the fragments, the data are called single-digest (one enzyme), double-digest (two enzymes), or *n*-digest (*n* enzymes) data. Most mappings are done using single- and double-digest data. Scientists use different restriction enzymes to obtain DNA fragments of the appropriate size.

In practice, scientists try to map one chromosome at a time. Many techniques generally require (a) purifying chromosomal DNA, (b) cutting the DNA into pieces called contigs using restriction enzymes, (c) inserting contigs into DNA cloning vectors, (d) inserting cloning vectors into bacterial host cells for multiplication, (e) cutting clones into fragments,

(f) analyzing the fragments, and (g) reconstructing the original order of the base-pairs on the chromosome.

#### 4.1 Restriction-Fragment Data for Chromosome 19

As the data used in this study are obtained from the Human Genome Center of LLNL, it is useful to summarize the techniques used at LLNL to gather these data.

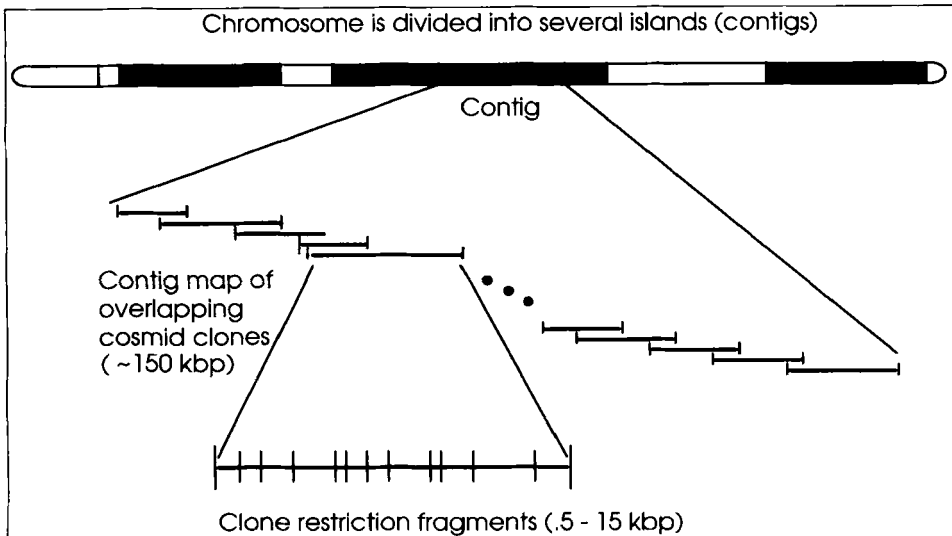
Chromosomes are sorted via a laser flow sorter; restriction enzymes are used to shatter the sorted material into fragments. These fragments are inserted into a host *vector*, which can then be grown in large quantities. Due to the properties of the cloning vector used at LLNL, our so-called *cosmid clones* average 40Kbp (40,000 base-pairs) of DNA each. It is important to note that in each such cloning step, all original order is lost.

The LLNL biologists cloned and selected 15,000 such cosmid clones for human chromosome 19. A “cosmid fingerprint” was obtained for each clone by subjecting it to a double-digest labeled with a fluorescent dye and measuring it on a modified DNA sequencing gel apparatus. These crude fingerprints were compared between all pairs (taking two days on a network of 40 workstations in parallel), and a set of 800 unordered islands or “contigs” was formed by an automated algorithm that also determined a near-minimal spanning path for each island. (Note that 15,000 40Kbp clones provides a nominal 10X depth of coverage for chromosome 19.) However, this set of islands remains unordered by this technique.

A wide range of other techniques, not covered here, is used to merge, order, and provide the distance between the contigs. This paper will focus on one technique used to verify the accuracy of the contigs built via pair-wise overlap data, called EcoRI restriction-fragment mapping after the name of the restriction enzyme used. A subset of the clones in a contig is chosen (i.e., the spanning path members plus other clones to attempt to get at least a 2X coverage across the contig) and digested, with fragment lengths in the range of about 0.5-20Kbp being generated, depending on the distribution of EcoRI sites within each cosmid. These maps are of great utility in locating and sequencing genes; since individual EcoRI fragments can be physically cut from gels and prepared for sequencing, the cost of sequencing a gene can often be reduced by an order of magnitude (assuming a gene can be localized to a ~4Kbp fragment, instead of having to sequence an entire 40Kbp clone).

The order of the fragments within each clone is, of course, unknown. At LLNL, the “EcoRI maps” are constructed rapidly by hand, since the existence of a reliable near-minimal spanning path provides a tremendous hint toward obtaining a proper clone ordering. This in effect reduces a 2-D optimization problem to a 1-D one. Once clone order is known, fragment ordering within clones is a much simpler problem (note that fragment sizes can be measured to about 5% and that fragments of the same size can occur in multiple unrelated positions within a single map.)

The process of initially assembling the contigs and thereby deriving the minimal spanning paths is itself quite slow and labor-intensive in terms of laboratory bench work. LLNL researchers were interested in seeing how well a fully automated system could do in assembling EcoRI maps if *a priori* contig information were not available. This is the problem discussed here: Given a number of clones, in unknown order, each with a set of restriction fragments, also in unknown order, construct a maximum-likelihood map of that region, subject to the constraints that, in general, the cosmids should be contiguous in the resulting maps, and apart from end fragments, all fragments should overlap others in their vertical “stack” within about 5% of their lengths. If these maps could be built accurately over large regions (i.e., at least 200–500Kbp), a potential exists for reducing the overall cost of obtaining a high-resolution physical map.



**Figure 7.** Physical map for island using fragments from a set of overlapping clones.

Toward the goal stated above, in this paper, we propose to establish the relative position of each cosmid clone on a contig by establishing the possible locations of the fragments in each cosmid clone in such a manner that the fragment overlap among the clones is maximized while a suitably defined total error between the overlapped fragments is minimized. There are other constraints, such as the total length of the assembly must be equal to the contig's original size. The problem is one of assembling cosmid clone sequences, as shown in Figure 7. The problem is complicated further by the uncertainty in the data, the possibility of data loss (fragments of the same size are hard to distinguish during fingerprinting), and the known fact that data related to *corner fragments* (i.e., fragments near the fragment boundaries) is almost always unreliable. Problems of this type are known to be hard (Opatrny, 1979). For example, in the case with 10 cosmid clones, there are  $10!/2$  possible clone sequences.

#### 4.2 Related Work and Scope of Present Work

The DNA restriction-fragment assembly resembles, and is somewhat related to, the *restriction-site mapping*, which deals with the equivalent problem of determining the absolute location of a fragment within a cosmid clone. Here also one uses digestion data from restriction enzymes, but the focus is on finding the absolute location of a fragment on a clone. Stefik (1978) used a branch and bound technique with rules to exhaustively eliminate wrong answers from the digest fragment data. This approach is sensitive to error in the data and is computationally intensive. Pearson (1982) exhaustively generated permutations of the single-digest data to compute the error between the generated double-digest and the actual (experimental) double-digest data. This approach is faster, but it is limited to a small number of restriction sites. Krawczak (1988) developed a divide-and-conquer technique that groups the fragments into compatible clusters and then determines the order of the fragments within each cluster. This approach can process a greater number of restriction sites. Platt and Dix (1993) used genetic algorithms for restriction-site mapping using double-digest data. In

their work they did not consider operators suited for multimodal search spaces and mating that preserves adjacency information.

Other techniques are available to sequence larger DNA regions. Branscomb, Slezak, Pae, Galas, Carrano, & Waterman (1990) developed a greedy algorithm to order the most probable clone sequence using overlap probabilities between the clones. The algorithm works well when a large amount of overlap between the clones exists and the fragment data has small errors. This approach is prone to getting stuck in local minima and does not use all the available data gathered at great expense. Techniques using larger clones are also being tried to order, orient, and connect the islands in the original DNA (Olson et al., 1986; Waterman & Griggs 1986; Stallings et al., 1990; Fickett & Cinkosky, 1993).

Cuticchia, Arnold, and Timberlake (1992) constructed maps using simulated annealing techniques. In their work, clones are ordered according to a measure of similarity between them given by the presence or absence of specific sequences. A signature is assigned to each clone and the algorithm uses it to minimize the error between the actual length of the contig and the given length by the hypothetical clone ordering. Matching signatures are used to order the clones. In their work they only considered the relationship between consecutive clones.

Recently, Parsons, Forrest, and Burks (1993, in press) applied the *edge recombination crossover* (Whitley, Starkweather, & Fugway, 1989) in conjunction with several specialized operators and solved a 10 Kbp sequencing problem consisting of 177 fragments with no manual intervention. Previously, we too had experimented with, but did not report, a variety of permutation-based crossover operators, such as those one finds in Traveling Sales Person type problems. In that context we also found and reported that the edge recombination operator outperformed the permutation-based operators by a wide margin (Cedeño & Vemuri, 1993). In this paper, we now describe a genetic algorithm that uses single-digest restriction data on a set of overlapping clones to find multiple solutions for the clone sequences. We use genetic operators suited for multimodal function optimization (Cedeño & Vemuri, 1992) to determine solutions. A mating operator based on genetic edge recombination was used to preserve adjacency information and improve convergence toward multiple solutions at the same time. Results are given for two data sets of overlapping clones from human chromosome 19.

Before delving into the details of the application of the MNC algorithm to the DNA problem, it is useful to pause and justify the need for multimodal optimization to solve this problem. We are aware of the fact that molecular biologists, unlike engineers, cannot use “sub-optimal” solutions. As any geneticist knows, one missing or misplaced base-pair could lead to the synthesis of an entirely different protein. Ideally, what a molecular biologist wants is the exact sequence, which should correspond in our mathematical formulation to the global optimum. But the search for this global optimum is intimately tied up with the selection of the optimization criterion, say the fitness function in GAs or the optimization criterion such as an energy function in classical optimization. As long as this criterion is fabricated by our methods, we will never be able to zero in on the “correct” solution using mathematical methods. The best we can do is to ease the burden on the biologist by showing him or her possible avenues for further experimental investigation. We believe that this is the place the sub-optimization plays a role, and that is why we were searching for the k-best sub-optima.

## 5. Problem Representation

It is worth repeating a cautionary word about the terminology used here. Hereafter, the word DNA is used synonymously with the word chromosome(s) found in a cell, whereas the

ALLELE NUMBER	CLONE ID	FRAGMENT SIZES (in thousands of base-pairs)
C0	5154	16.55, 4.4, 1.68, 1.07, 4.81, 8.5
C1	7442	0.79, 0.79, 2.6, 4.35, 8.24, 2.7, 6.9, 5.16
C2	21230	0.96, 1.68, 1.08, 4.77, 8.47, 1.44, 2.37, 6.29, 0.62
C3	8131	0.92, 3.73, 19.8, 4.43, 1.69, 1.25, 4.68, 5.63
C4	18993	0.96, 6.31, 5.48, 8.61, 7.29, 0.81, 0.81, 2.6, 4.36, 1.92
C5	5435	2.89, 8.24, 2.7, 6.9, 5.14, 5.14, 2.89, 1.54
C6	7255	1.04, 8.21, 2.69, 6.89, 5.12, 5.12, 2.88, 1.94, 2.42, 1.37, 3.33
C7	12282	4.52, 5.13, 5.13, 2.87, 1.94, 2.42, 1.39, 3.35, 5.41
C8	27714	6.69, 5.07, 5.41, 2.88, 1.92, 2.32, 1.4, 3.35, 5.46, 17.65, 1.0, 10.49, 0.58, 1.74
C9	10406	2.03, 1.43, 2.34, 6.28, 5.46, 8.58, 7.27

**Figure 8.** Cosmid clones with fragment data.

word *chromosome* will be used to describe the computational data structure used in genetic algorithms. This interesting situation arose because genetic algorithms are computational processes that imitate natural processes of selection and survival. Coincidentally, we are applying this metaphorical computational paradigm to solve a problem in genetics! So it is important to distinguish meanings of terms used in the problem domain from those used in the paradigm domain.

In this section some problem-dependent genetic operators are defined. First, the encoding of the chromosomes to describe clone sequences is examined. Second, the use of fragment sizes to define the fitness function is described. Third, the modified edge recombination mating operator is described. And last, the function that measures similarity between two clone sequences is described.

Before going into detail about the operators, it is important to show how the data for the problem is presented to the GA. Figure 8 shows fragment sizes obtained from fingerprinting for a set of overlapping cosmid clones. For example, cosmid clone with the ID number 5154, which is also labeled as Allele Number C0, is known to comprise six fragments, containing 16550, 4400, 1680, 1070, 4810, and 8500 base-pairs, in that order. Also, cosmid clone with the ID number 8131, also labeled as Allele Number C3, is known to comprise eight fragments, containing 920, 3730, 19800, 4430, 1690, 1250, 4680, and 5630 base-pairs, in that order. By comparing these fragment sequences one can surmise that the third and fourth fragments of Allele C0 are probably the same as the fifth and sixth fragments of Allele C3 mainly because the fragment lengths are so nearly equal to each other. If this is true, then Allele C3 can be “aligned” below Allele C0 in such a manner that the fifth and sixth fragments of Allele C3 fall right below the third and fourth fragments of Allele C0, as shown in Figure 9. The matching of the fragment lengths is not perfect. Indeed, the mismatch at other positions is large. The goal of this problem is to maximize this type of matching while minimizing the number and degree of mismatches and keeping the total length of the assembly within reasonable limits. The data for this problem consist of the  $n$  cosmid clones with their fragment lengths and the tolerance measure  $\epsilon$ , which is used to determine if two fragments are of the same size. That is, two fragments F1 and F2 are considered to be of the same size if  $|F1 - F2| < \epsilon$ .

Clone	Fragments																			
C3 8131	0.92	3.73	19.8	4.43	1.69	1.25	4.68	5.63												
C0 5154			16.55	4.4	1.68	1.07	4.81	8.5												
C2 21230				0.96	1.68	1.08	4.77	8.47	1.44	2.37	6.29	0.62								

Figure 9. An example of fragment assembly.

Number of matches between clones											Total error in the matches									
CLONE	C0	C1	C2	C3	C4	C5	C6	C7	C8	C9	C0	C1	C2	C3	C4	C5	C6	C7	C8	C9
C0	6	1	4	2	1	0	1	0	2	1	0	5	8	4	4	0	3	0	13	8
C1	1	8	0	1	4	4	4	1	1	0	5	0	0	8	5	2	9	3	9	0
C2	4	0	9	3	2	1	3	2	5	3	8	0	0	14	2	10	16	10	23	5
C3	2	1	3	8	2	0	0	1	2	0	4	8	14	0	11	0	0	9	13	0
C4	1	4	2	2	10	1	3	2	3	4	4	5	2	11	0	10	19	9	6	10
C5	0	4	1	0	1	8	6	3	2	0	0	2	10	0	10	0	10	4	8	0
C6	1	4	3	0	3	6	11	7	7	3	3	9	16	0	19	10	0	7	26	23
C7	0	1	2	1	2	3	7	9	7	4	0	3	10	9	9	4	7	0	20	26
C8	2	1	5	2	3	2	7	7	14	3	13	9	23	13	6	8	26	20	0	5
C9	1	0	3	0	4	0	3	4	3	7	8	0	5	0	10	0	23	26	5	0

Figure 10. The  $M$  matrix on the left whose entries are the number of fragment matches and the  $E$  matrix on the right whose entries are the total errors between any two clones with  $\varepsilon = 10$ .

### 5.1 Chromosome Encoding

The encoding for this problem is simple. Each allele in the chromosome has a label between 0 and  $n-1$  corresponding to one of the cosmid clones. No two alleles have the same label, and mating and mutation will preserve this constraint. In Figure 8, for example, an allele with the label C0 corresponds to the clone with ID 5154, and an allele with the label C9 corresponds to clone ID 10406. The clone sequence (5154, 21230, 10406, 7255, 12282, 27714, 8131, 18993, 7442, 5435), for example, is represented by the chromosome (0 2 9 6 7 8 3 4 1 5). The initial population is generated by picking, at random, values between 0 and  $n-1$  without replacement.

### 5.2 Fitness Function

To calculate the fitness of an individual, the number of fragment matches between all consecutive clones and the error between the fragments are considered. The fragment sizes are represented using integer numbers (by multiplying the number given in Figure 8 by 100), to accelerate computation of the fitness function. Prior to the execution of the GA, two matrices are calculated. One matrix, the match matrix  $M$  shown on the left side of Figure 10, contains the number of fragments that match between two clones  $C_i$  and  $C_j$ , within an error tolerance  $\varepsilon$ . The other matrix, the error matrix  $E$  shown on the right side of Figure 10, contains the total error between the clones being matched. The error between two clones is given by the sum of the errors between all fragments that matched. For example, between clone No. 8131 (C3) and clone No. 5154 (C0) there are two pairs of fragments that match within the specified tolerance of  $\varepsilon = 10$ . The lengths of these fragments are 169 and 168 for one pair and 443 and 440 for the second pair. Thus a two appears in row 2, column 4 of the match matrix  $M$ . The total error between both pairs of fragments is  $(169 - 168) + (443 - 440) = 4$ , which is shown in row 1, col. 4 of the error matrix  $E$ .



Our goal is to arrange the cosmid clones as shown in Figure 7 so that the lengths of the overlapping fragments match with each other as closely as possible. The necessary matching information is already gathered in the matrix  $M$ , and the degree of accumulated mismatch per clone is gathered in the matrix  $E$ . However, we believe that this information alone is not sufficient to establish which two clones are “adjacent” to each other in the arrangement shown in Figure 7. For example, consider how clone C0 (i.e., allele No. 5154) matches with other clones. Inspection of the match matrix  $M$  indicates that the degree of match between clone C0 and clone C2 (or, equivalently, allele No. 21230), is four matches. Also, fragments in clone C0 match with fragments in clone C3 as well as C8, each with two matches. By interpreting this to mean that C2 should be placed nearer to C0 than C3 or C8, we are ignoring information contained in the C0-C3 matches and C0-C8 matches. One more example suffices to make the point. Clone C3 should be placed closer to C2 because they match with each other the maximum number of times, namely three, although C3 matches with three other clones, each with only two matches. This phenomenon makes us think that using only the number of matches between clones is not sufficient to establish the partial order between clones when they possess the same match count. We believe that part of this problem is due to false matches, between fragments of similar sizes, that may occur by chance. We tried to overcome this problem by incorporating the total error in the matches, shown in matrix  $E$ , in order to enable our GA to discriminate further between clones. Using the same example, notice that clone C3 has less total error when matched with C0 than clone C8 and therefore indicates that C0 is adjacent to C3. The following equation for fitness captures the essence of the method described so far.

$$fitness = \sum_{i=0}^{n-1} \sum_{\substack{j=-1 \\ j \neq 0}}^1 \frac{M[a_i, a_{i+j}]}{Count[a_{i+j}]} \left( 1 - \frac{E[a_i, a_{i+j}]}{M[a_i, a_{i+j}] \cdot \varepsilon} \right)$$

Here  $a_i$  refers to the cosmid clone placed in the  $i^{\text{th}}$  position of the chromosome,  $a_{i+j}$  refers to the clone to the right or left (if any) of the  $i^{\text{th}}$  position. The first term,  $M[a_i, a_{i+j}] / Count[a_{i+j}]$ , gives us a normalized count for the degree of match. The term  $E[a_i, a_{i+j}] / M[a_i, a_{i+j}] \cdot \varepsilon$  refers to the normalized error per fragment. When this normalized error reaches unity, it means that the total error is so large that any apparent matches are worthless. With this interpretation, the second term of the equation essentially tells us the degree of confidence we can place on the normalized matches we are counting in the first term. In the above equation,  $n$  refers to the number of alleles in the chromosome, which is equal to the number of clones.

By defining the fitness function as above, we are assigning a higher fitness to those clone pairs that match a higher percentage of their regions. For example, Allele 0 with 6 fragments has two matches each with Alleles 3 and 8, each having 8 and 14 fragments, respectively. Since  $2/8$  represents a higher percentage than  $2/14$ , we designed a fitness function that prefers a configuration that places Allele 3 closer to Allele 0 than Allele 8. This is achieved by dividing the number of matches by the number of fragments in the clone.

Before settling on the fitness function described above, others were considered. For example, fitness functions that just counted the number of matches between clones with no regard to normalization failed to produce the correct answer. A fitness function that just counted the number of matches and then subtracted the total error in those matches also failed to give satisfactory results. It is possible that other fitness functions may give results that are even better than those reported here. In the future, we plan to include the number

Parent 1	Common links (traits)	Offspring 1
(6 7 8 1 2 3 9 4 5 0)	(- 7 8 1 - - - - 5 0)	(3 7 8 1 4 6 2 9 5 0)
Parent 2	Common links (traits)	Offspring 2
(1 8 7 9 5 0 2 6 3 4)	(1 8 7 - 5 0 - - - -)	(1 8 7 3 5 0 6 2 4 9)

**Figure 11.** Modified genetic edge recombination for clone sequencing.

of matches as well as errors among groups of three clones or more as components of the fitness function and study its effect on performance.

### 5.3 Mating and Mutation Operators

The mating operator used in this method is based on a slight modification to the genetic edge recombination operator that was applied successfully to solve the TSP (Traveling Sales Person) problem. As in the TSP problem, the important information here is the adjacency of the alleles, although the order the alleles appear in the chromosome can be derived from the adjacency information. The idea is to recombine the links (pairs of clones) between two parents such that common links are inherited by the offspring. This operator is implemented in two steps, as shown in Figure 11. First, those links (or traits) that are common to both the parents are identified and passed on to the offspring; the links occupy the same absolute positions in the offspring chromosome. In the example shown in Figure 11, the relevant link-pairs are 7-8, 8-1, and 5-0 in the first parent and 1-8, 8-7, and 5-0 in the second parent. Notice that these links are passed on to the two offspring undisturbed. Second, those alleles that are not passed to the offspring (indicated by dashes, in Figure 11) are randomly assigned to the available positions while observing the constraint that no link label is repeated.

The differences between this operator and the original edge recombination operator are in the number of offspring generated and in the assignment of alleles not transferred from the parent. We generated two offspring instead of one because the location of the links in the clone sequence is important to our problem. In TSP the chromosome is circular; thus, the location did not matter. We allow both parents to pass the location of the links to their offspring. To assign the other alleles we select them at random from those clones not passed by their parents. In the original operator the links are assigned from those present in either of the two parents. Alleles with fewer links are assigned first to prevent running out of links for a given allele.

In the mating operator used here, there is excessive exploration of the search space primarily due to the random filling of the unassigned slots, in the second step, while creating the offspring chromosomes. Part of this exploration difficulty is alleviated by the fact that mates are selected using crowding selection and, therefore, they share common features. Exploration is therefore localized to a smaller region within the entire search space. On the other hand, by allowing unassigned clones to be chosen at random, we are allowing links to reappear that might not have done so using mutation alone. Other crossover operators based on clone positions alone did not perform as well in this problem; we give the results only for the modified edge recombinator operator.

Mutation is applied on an individual basis. After an offspring is generated it is mutated if the outcome from the flip of a biased coin is true. When this happens, a link from the offspring is selected at random and all alleles from that link to the last position of the

chromosome are reversed. For example, the offspring (1 8 7 3 5 0 6 2 4 9) after mutation can result in (1 8 7 9 4 2 6 0 5 3) if the link between allele 7 and 3 is selected to mutate. This mutation operator is known as *inversion* (Holland, 1975).

The mating and mutation operators are compatible with each other in the sense that they both operate on links. The building blocks of this problem are based on the links between clones in the sequence. The GA operates on these links so that the most useful ones are passed from generation to generation.

#### 5.4 Similarity Function

The similarity function is very simple also. It counts the number of dissimilar links between two individuals. Using the parents from Figure 11 once again as an example, notice that there are six dissimilar links, corresponding to the five alleles not assigned to the offspring. For concreteness, these six dissimilar links in Parent 1 chromosome are 6-7, 1-2, 2-3, 3-9, 9-4, and 4-5; for Parent 2 they are 7-9, 9-5, 0-2, 2-6, 6-3, and 3-4. This metric measures the proximity between two clone sequences by counting the different links they have, not the position of the alleles. For example, the sequences (0 1 2 3 4 5 6 7 8 9) and (9 8 7 6 5 4 3 2 1 0) have a distance of zero since all the links are the same. This metric captures the essential aspect of the problem since both solutions are equivalent in our problem.

### 6. Results and Discussion

The results presented in this section were obtained on an SGI IRIS 4D computer under IRIX O.S. running the GA application written in C. The parameters for the GA are the following:

Population size	100
Mutation probability	0.06
Crossover probability	1.00
Crowding selection group size ( $C_s$ )	20
Crowding factor group size ( $s$ )	10
Crowding factor ( $C_f$ )	5
Maximum number of generations to execute	100
Tolerance $\varepsilon$	10

These parameters were selected after various trials. In each trial, different values for each of the six parameters were tried, varying one at a time while holding the others constant. A population size of 100 was found satisfactory. Other sizes (50, 150, 200, 250, and 300) were tried. Higher sizes did not provide new information about the problem, although they produced, on average, the solution in fewer generations. Lower sizes in some cases did not converge to the best solutions seen before in the allowed number of generations. The population size of 100 is a compromise between speed of convergence and execution time.

Mutation was set at 0.06; therefore, an average of six individuals were mutated every generation. This low value of mutation works well in this problem. It seems to allow individuals to escape local optima without causing any major disruption in the elements of a niche. In the MNC method, diversity is maintained implicitly when the population converges to different optima, reducing the need for higher mutation values. On the other hand, the mating probability was set to a high value of 1.0. As described before, the MNC method

is basically a steady-state GA, and highly fit individuals have a good chance of surviving for many generations. Also, crowding selection exploits the similarity between individuals in the population by allowing localized solutions to pass common traits between them.

The group size for crowding selection was set at 20. This high value emphasizes the importance of similarity between parent and mate for this problem. Then the mating operator will likely generate an offspring within the same region as its parents. The size for the crowding factor groups was set at 10. This allowed competition between multiple optima to occur while maintaining multiple solutions. Good results were obtained with other  $C_r$  and  $s$  values (5, 10, 15, 20) as well. The crowding factor was set at 5. This value allowed the MNC method to eliminate low fitness individuals from the niches more rapidly. These parameter values allowed a diverse population to coexist during the number of generations allowed and did not restrict competition between individuals from different niches. The tolerance value  $\varepsilon$  was set to 10 to minimize false matches due to chance. Higher values of  $\varepsilon$  increased the false matches more than true matches and, therefore, more possible clone sequences were found.

The MNC method took an average of 50 seconds for each run. Some of the best sequences obtained for two different data sets of overlapping clones are shown in Figure 12. These results were obtained from 10 different runs. The figure shows the actual sequence for the data sets and the clone sequences (with their fitness) obtained by the algorithm.

Data for set 1 is shown in Figure 8, and data for set 2 is shown in the Appendix. There was nothing really special about these data sets except that they were in regions containing various gene(s) of interest to some of our collaborators. They had already mapped them manually, so we could judge the answers our GA technique provided. Finally, and perhaps most importantly, the data sets were large enough to avoid being toy problems. One of the data sets contained two vertical “columns” in the final map that had fragments of the same size. The existence of independent columns of the same size is, of course, one of the things that makes this problem so tough, and so interesting. Many of the runs using data set 2 generated multiple solutions with relatively close fitness values.

For data set 1 the MNC method was able to find the actual clone sequence. The best sequence as described by the fitness function did indeed match the solution. The best sequence was found in all the runs. From the other sequences found, the fitness values of the next best is 15 less than the actual sequence. Similar gaps exist between all the sequences shown in Figure 12 for data set 1. From the solutions we can see that the other sequences are a single mutation from the actual sequence.

Data set 2, shown in the Appendix, presented a more challenging problem for the MNC method. In this case the best sequence found had only clones 6722 (C7) and 26999 (C2) transposed from the actual sequence. The fitness for the actual sequence is 750, which is the sixth best score when compared with all solutions found. The actual sequence was obtained in 8 of the 10 runs. As shown in Figure 12, different solutions with equal fitness values were found and maintained in the runs. Two different solutions with a distance of 2 (two different links) were found with fitness equal to 757. Four others were found with fitness equal to 750 (including the actual sequence). Another observation is that there is a difference of 8 or less in the fitness between all the sequences found. Some of the sequences are mutations of others, but there is more diversity when compared with the solutions for data set 1. The biologists later confirmed via independent techniques (i.e., hybridization probing) that the clones in data set 2 were indeed from two distinct regions in the DNA. That is, the sequence (6722 26999 12595 29626 29064 18301) and (19811 29035 17755 28828 20235) belong to separate maps in the DNA. The best solutions found by the MNC method kept both sequences apart.

Data set 1 actual sequence and its fitness:

(8131 5154 21230 10406 18993 7442 5435 7255 12282 27714) 764

The three best sequences found by the GA and their corresponding fitnesses:

(8131 5154 21230 10406 18993 7442 5435 7255 12282 27714) 764

(8131 5154 21230 27714 12282 7255 5435 7442 18993 10406) 749

(8131 5154 21230 10406 27714 12282 7255 5435 7442 18993) 744

Data set 2 actual sequence and its fitness:

(12595 6722 26999 29626 29064 18301 19811 29035 17755 28828 20235) 750

The 12 best sequences found by the GA and their corresponding fitnesses:

(12595 26999 6722 29626 29064 18301 28828 20235 17755 29035 19811) 757

(20235 28828 17755 29035 19811 18301 29064 29626 6722 26999 12595) 757

(12595 26999 6722 29626 29064 18301 20235 28828 17755 29035 19811) 756

(28828 20235 17755 29035 19811 18301 29064 29626 6722 26999 12595) 755

(12595 26999 6722 29626 29064 18301 28828 20235 17755 19811 29035) 754

(12595 26999 6722 29626 29064 18301 20235 28828 17755 19811 29035) 753

(12595 6722 26999 29626 29064 18301 19811 29035 17755 28828 20235) 750

(20235 28828 17755 29035 19811 18301 29064 29626 12595 6722 26999) 750

(19811 29035 17755 20235 28828 18301 29064 29626 12595 6722 26999) 750

(19811 29035 17755 20235 28828 18301 29626 29064 26999 6722 12595) 750

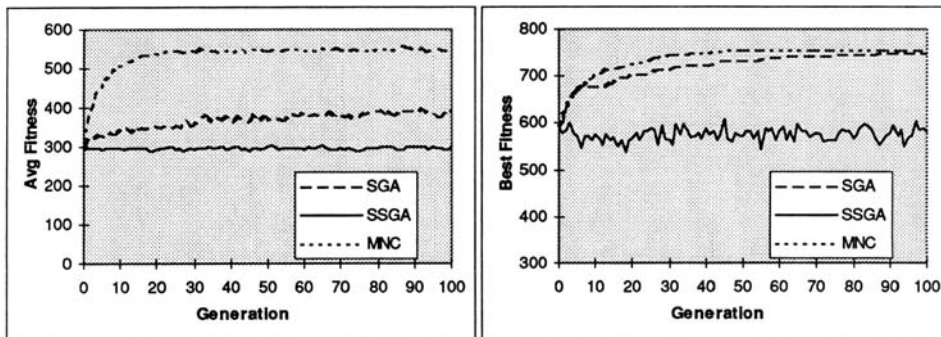
(19811 29035 17755 28828 20235 18301 29064 29626 26999 6722 12595) 749

(19811 29035 17755 28828 20235 18301 29064 29626 12595 6722 26999) 749

**Figure 12.** Clone sequences obtained by GA and actual sequence.

To see how the MNC method compares with more traditional GAs we applied the same problem (i.e., using data set 2) to the SGA and SSGA with crowding. For both the SGA and SSGA the crossover probability was set at 0.6. The crowding factor was set to 10 for the SSGA. All other parameters are the same as in the MNC method. We observed poor results by the SGA using these parameters alone. To improve the performance of the SGA the 10 best individuals in the population—known as generation gap (De Jong, 1975)—were transferred from the current generation to the next. Figure 13 shows the population's average fitness (on the left) and the fitness of the best solution (on the right) for the MNC, SGA, and SSGA methods averaged over 10 runs.

The MNC method outperformed both the SGA and the SSGA. The population converged faster and at the same time kept multiple solutions throughout the run. Use of a generation gap allowed the SGA to retain the good individuals found in previous runs, but it did not improve convergence toward multiple solutions. The SSGA did not find the best sequence in any of the runs. The SGA found one of the two best sequences in two of the runs. As mentioned before, the MNC found the two best sequences and others in eight of



**Figure 13.** Population average fitness (left) and best fitness (right) for different methods.

the runs. The ability of the MNC method to exploit similarity and increase competition among localized solutions proved to be successful in this problem.

## 7. Comments and Conclusions

Two points deserve further discussion. First, data containing clones with fewer than five fragments were normally sequenced erroneously by the GA. This is due to the lack of opportunity for sufficient fragment matches. Also, data pertaining to corner fragments (i.e., fragments lying near the cosmid clone boundaries) is generally more prone to errors. Consequently, corner segments will not match well with a high probability with their counterparts in the preceding and succeeding clones. For clones with fewer than five fragments, this means that, on the average, at least half of the data is not useful and in some cases leads to more false matches. Clones with fewer than five fragments were usually placed first or last in the clone sequence by the GA.

Second, when a large number of overlaps existed between three or four clones, the GA experienced difficulty deciding the correct sequence. An example of this behavior was observed with data set 2. This phenomenon, we believe, is happening because the fitness function is looking only for matches between the clones to the left and right without accounting for the fragments that are common to three or more clones. Permutations of these clones generally had similar fitness values. An improved fitness measure is needed to account for fragment matches between three or more clones.

Overall, the MNC worked well with the data presented to it. Using the correct set of genetic operators was very important to find a MNC model that will find good solutions to the problem. Using a multimodal approach was very useful for this problem also because it prevented premature convergence and at the same time explored the search space in a more efficient manner. Defining the operators for mating, mutation, fitness, and similarity measure to work with adjacency information between the clones rather than clone positions gave the MNC method the correct set of tools to converge toward the most probable solutions. More information must be incorporated into the fitness evaluation to distinguish even further between the best clone sequences and other similar ones.

The results on multimodal optimization and DNA fragment assembly show the ability of the MNC to converge to multiple solutions, maintain stable subpopulations, and succeed

in complex search spaces. Exploiting similarity during selection and replacement allows a diverse population to coexist while competition for slots in the population evolve naturally. Convergence toward the best solution(s) is not affected by the increased diversity. The MNC method uses the diversity present in the population to guide the population toward multiple optima in the search space.

## Appendix

Fragment data for data set 2 of overlapping cosmid clones:

CLONE ID    FRAGMENTS (k base-pairs)

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29064	15.42, 3.46, 1.50, 9.12, 4.30
19811	3.13, 7.89, 7.89, 3.02, 4.35, 14.31, 2.65, 0.64
26999	4.64, 19.69, 1.10, 1.48, 2.82, 0.77, 9.61
29626	3.46, 1.50, 9.14, 6.47, 13.48
17755	1.26, 2.62, 6.32, 2.73, 3.54, 7.88, 7.88, 3.02, 4.35, 1.74
12595	1.48, 2.83, 0.76, 9.68, 12.75, 1.48, 6.06
20235	8.01, 12.56, 2.62, 6.32, 2.74, 3.54, 5.71
6722	2.84, 19.72, 1.16, 1.49, 2.84, 0.77, 9.69, 9.20
28828	12.45, 2.61, 6.27, 2.72, 3.52, 7.89, 3.89
18301	2.53, 13.99, 2.64, 16.88, 3.44
29035	2.45, 2.74, 3.53, 7.82, 7.82, 3.02, 4.35, 7.44

CLONE	C0	C1	C2	C3	C4	C5	C6	C7	C8	C9	C10
C0 :	5	1	1	3	2	1	1	2	1	1	2
C1 :	1	8	0	0	5	0	1	0	2	1	5
C2 :	1	0	7	1	1	4	1	6	1	0	1
C3 :	3	0	1	5	1	1	1	2	1	1	1
C4 :	2	5	1	1	10	1	4	1	5	2	6
C5 :	1	0	4	1	1	7	1	4	0	0	1
C6 :	1	1	1	1	4	1	7	1	4	2	2
C7 :	2	0	6	2	1	4	1	8	0	0	1
C8 :	1	2	1	1	5	0	4	0	7	2	3
C9 :	1	1	0	1	2	0	2	0	2	5	3
C10:	2	5	1	1	6	1	2	1	3	3	8

CLONE	C0	C1	C2	C3	C4	C5	C6	C7	C8	C9	C10
C0 :	0	5	2	2	13	2	8	9	6	2	12
C1 :	5	0	0	0	5	0	3	0	4	1	23
C2 :	2	0	0	2	9	9	8	20	10	0	8
C3 :	2	0	2	0	8	2	8	7	6	2	7
C4 :	13	5	9	8	0	10	1	10	10	12	14
C5 :	2	0	9	2	10	0	9	4	0	0	9
C6 :	8	3	8	8	1	9	0	10	10	12	1
C7 :	9	0	20	7	10	4	10	0	0	0	10
C8 :	6	4	10	6	10	0	10	0	0	11	10

C9 : 2 1 0 2 12 0 12 0 11 0 27  
 C10: 12 23 8 7 14 9 1 10 10 27 0

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