# From cells to islands: An unified model of cellular parallel Genetic Algorithms

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**Abstract.** This paper presents the Anisotropic selection scheme for cellular Genetic Algorithms (cGA). This new scheme allows to enhance diversity and to control the selective pressure which are two important issues in Genetic Algorithms, especially when trying to solve difficult optimization problems. Varying the anisotropic degree of selection allows swapping from a cellular to an island model of parallel genetic algorithm. Measures of performances and diversity have been performed on one well-known problem: the Quadratic Assignment Problem which is known to be difficult to optimize. Experiences show that, tuning the anisotropic degree, we can find the accurate trade-off between cGA and island models to optimize performances of parallel evolutionary algorithms. This trade-off can be interpreted as the suitable degree of migration among subpopulations in a parallel Genetic Algorithm.

### Introduction

In the context of cellular genetic algorithm (cGA), this paper proposes the Anisotropic selection as a new selection scheme which accurately allows to adjust the selective pressure and to control the exploration/exploitation ratio. This new class of evolutionary algorithms is supervised in a continuous way by an unique real parameter  $\alpha$  in the range [-1..1]. The work described in this paper is an attempt to provide a unified model of parallel genetic algorithms (pGA) from fine grain massively parallel GA (cGA) to coarse grain parallel model (island GA). As extreme cases, there are the cGA that assumes one individual resides at each cell, and at the opposite, a pGA where distinct subpopulations execute a standard GA; between them we find models of pGA where migration allows to exchange to some extend genetic information between subpopulations. Thus the search dynamics of our family of pGA can vary from a diffusion to a migration process. To illustrate our approach we used one well-known problem: the Quadratic Assignment Problem (QAP). We study the performances of our class of parallel evolutionary algorithms on this problem and we show that there is a threshold for parameter  $\alpha$  according to the average performances. Section 1 gives a description of the cGA and the island models. Section 2 introduces the anisotropic parallel Genetic Algorithms (apGA) and the anisotropic selection scheme. Section 3 is a presentation of the test problem: the QAP, and gives the performances of the apGA on the QAP. Finally, a study on population genotypic diversity is made in section 4.

## 1 Background

This section introduces the concepts of Cellular and Island Models of parallel genetic algorithms.

### 1.1 Cellular Genetic Algorithms

The Cellular Genetic Algorithms are a subclass of Evolutionnary Algorithms in which the population is generally embedded on a two dimensional toroidal grid. In this kind of algorithms, exploration and population diversity are enhanced thanks to the existence of small overlapped neighborhoods [9]. An individual of the population is placed on each cell of the grid and represents a solution of the problem to solve. An evolutionnary process runs simultaneously on each cell of the grid, selecting parents from the neighborhood of the cells and applying operators for recombination, mutations and replacement for further generations. Such a kind of algorithms is especially well suited for complex problems [5]. One of the interests of cGA is to slow down the convergence of the population among a single individual. Complex problems often have many local optima, so if the best individual spreads too fast in the population it will improve the chances to reach a local optimum of the search space. Slowing down the convergence speed can be done by slowing down the selective pressure on the population.

### 1.2 Island Model of pGA

Cellular genetic algorithms and Island Model genetic algorithms are two kinds of Parallel genetic algorithms. The first one is a *fine grain* massively parallel implementation that assumes one individual resides at each cell. The second one, using distinct subpopulations, is a *coarse grain* parallel model; Each subpopulation executes as a standard genetic algorithm, and occasionally the subpopulations would exchange a few strings: *migration* allows subpopulations to share genetical material [4]. Many topologies can be defined to connect the islands. In the basic island model, migration can occur between any subpopulations, whereas in the *stepping stone* model islands are disposed on a ring and migration is restricted to neighboring islands.

# 2 Anisotropic Parallel Genetic Algorithms

This section presents the *anisotropic parallel Genetic Algorithms*, which is a family of parallel genetic algorithms based on cellular GA in which anisotropic selection is used.

#### 2.1 Definition

The Anisotropic selection is a selection method in which the neighbors of a cell may have different probabilities to be selected. The Von Neumann neighborhood of a cell Cis defined as the sphere of radius 1 centered at C in manhattan distance. The Anisotropic selection assigns different probabilities to be selected to the cells of the Von Neumann neighborhood according to their position. The probability to choose the center cell C remains fixed at  $\frac{1}{5}$ . Let us call  $p_{ns}$  the probability of choosing the cells North (N) or South (S) and  $p_{ew}$  the probability of choosing the cells East (E) or West (W). Let  $\alpha \in [-1, 1]$  be the control parameter that will determine the probabilities  $p_{ns}$  and  $p_{ew}$ . This parameter will be called the *anisotropic degree*. The probabilities  $p_{ns}$  and  $p_{ew}$  can be described as:

$$p_{ns} = \frac{(1 - p_c)}{2} (1 + \alpha)$$
$$p_{ew} = \frac{(1 - p_c)}{2} (1 - \alpha)$$

Thus, when  $\alpha = -1$  we have  $p_{ew} = 1 - p_c$  and  $p_{ns} = 0$ . When  $\alpha = 0$ , we have  $p_{ns} = p_{ew}$  and when  $\alpha = 1$ , we have  $p_{ns} = 1 - p_c$  and  $p_{ew} = 0$ .

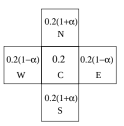


Fig. 1. Von Neumann neighborhood with probabilities to choose each neighbor

Figure 1 shows a Von Neumann Neighborhood with the probabilities to select each cell as a function of  $\alpha$ .

The Anisotropic Selection operator works as follows. For each cell it selects k individuals in its neighborhood ( $k \in [1; 5]$ ). The k individuals participate to a tournament and the winner replaces the old individual if it has a better fitness or with probability 0.5if the fitnesses are equal. When  $\alpha = 0$ , the anisotropic selection is equivalent to a standard tournament selection and when  $\alpha = 1$  or  $\alpha = -1$  the anisotropy is maximal and we have an uni-dimensional neighborhood with three neighbors only. In the following, considering the grid symmetry we will consider  $\alpha \in [0, 1]$  only: when  $\alpha$  is in the range [-1;0] making a rotation of 90° of the grid is equivalent to considering  $\alpha$  in the range [0;1]. When the anisotropic degree is null, there is no anisotropy in selection method, the apGA corresponds to the standard cellular GA. When the anisotropic degree is maximal, selection is computed between individuals in the same column only, the apGA is then an island model where each subpopulation is a column of the grid structured as a ring of cells with no interactions between subpopulations. When the anisotropic degree is set between low and maximum value, according to selection, a number of individuals can be copied from one subpopulation (i.e. column) to the adjacent columns. Thus the anisotropic degree allows to define a family of parallel GA from a cellular model to an island model.

In standard island model, the migration rate is defined as the number of individuals which are swap between subpopulations and migration intervals is the frequency of migration. In apGA, the migration process is structured by the grid. Only one parameter (the anisotropic degree) is needed to tune the migration policy. There is a difference between migration in a standard island model and migration in an apGA. In an apGA it can only happen (when the anisotropic degree allows it) between nearest neighbors in adjacent columns. Migration in that latter case is diffusion as it happens in the standard cGA model, except that the direction is controllable. In the following sections, we study the influence of this parameter on selection pressure, performances and population diversity.

#### 2.2 Takeover times and apGAs

The selective pressure is related to the population diversity in cellular genetic algorithms. One would like to slow down the selective pressure when trying to solve multimodal problems in order to prevent the algorithm from converging too fast upon a local optimum. On the opposite side, when there is no danger of converging upon a local optimum, one would like to increase the selective pressure in order to obtain a good solution as fast as possible. A common analytical approach to measure the selective pressure is the computation of the takeover time [8] [10]. It is the number of generations needed for the best individual to conquer the whole grid when the only active operator is the selection [3]. Figure 2 shows the influence of the anisotropic degree on the takeover time. This figure represents the average takeover times observed on 1000 runs on a  $32 \times 32$ grid for different anisotropic degrees. It shows that the selective pressure is decreasing while increasing anisotropy. These results confirm that the anisotropic selection gives to the algorithm the ability to control accurately the selective pressure. They are fairly consistent with our expectation that selection intensity decreases when the anisotropic degree increases. However, the correlation between takeover and anisotropy is not linear; it fast increases after the value  $\alpha = 0.9$ .

## 3 Test problem

This section presents tests on one well-known instance of the Quadratic Assignment Problem which is known to be difficult to optimize. Our aim is to study the dynamics of the apGA for different tunings, and not to obtain better performances than other optimization techniques. Still, the apGA is implicitly compared to a cellular genetic algorithm when the anisotropic degree is null ( $\alpha = 0$ ).

#### 3.1 The Quadratic Assignment Problem

We experimented the family of apGAs on a Quadratic Assignment Problem (QAP): Nug30. Our purpose here is not to obtain better results with respect to other optimization methods, but rather to observe the behavior of apGAs. Especially we go in the search of a threshold for the anisotropic degree.

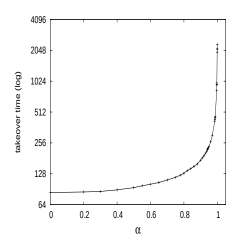


Fig. 2. Average of the takeover time as a function of the anisotropic degree  $\alpha$ .

The QAP is an important problem in theory and practice as well. It was introduced by Koopmans and Beckmann in 1957 and is a model for many practical problems [6]. The QAP can be described as the problem of assigning a set of facilities to a set of locations with given distances between the locations and given flows between the facilities. The goal is to place the facilities on locations in such a way that the sum of the products between flows and distances is minimal. Given n facilities and n locations, two  $n \times n$  matrices  $D = [d_{ij}]$  and  $F = [f_{kl}]$  where  $d_{ij}$  is the distance between locations i and j and  $f_{kl}$  the flow between facilities k and l, the objective function is:

$$\Phi = \sum_{i} \sum_{j} d_{p(i)p(j)} f_{ij}$$

where p(i) gives the location of facility *i* in the current permutation *p*. Nugent, Vollman and Ruml proposed a set of problem instances of different sizes noted for their difficulty [2]. The instances they proposed are known to have multiple local optima, so they are difficult for a genetic algorithm. We experiment our algorithm on the 30 variables instance called Nug30.

#### 3.2 Setup

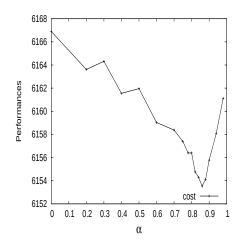
We use a population of 400 individuals placed on a square grid  $(20 \times 20)$ . Each individual represents a permutation of  $\{1, 2, ..., 30\}$ . We need a special crossover that preserves the permutations:

- Select two individuals  $p_1$  and  $p_2$  as genitors.
- Choose a random position *i*.
- Find j and k so that  $p_1(i) = p_2(j)$  and  $p_2(i) = p_1(k)$ .

- exchange positions i and j from  $p_1$  and positions i and k from  $p_2$ .
- repeat n/3 times this procedure where n is the length of an individual.

This crossover is an extended version of the UPMX crossover proposed in [7]. The mutation operator consist in randomly selecting two positions from the individual and exchanging those positions. The crossover rate is 1 and we do a mutation per individual. We perform 500 runs for each anisotropic degree. Each run stops after 1500 generations.

#### 3.3 Experimental results



**Fig. 3.** Average costs as a function of  $\alpha$  for the QAP.

Figure 3 shows the average performance of the algorithm towards  $\alpha$  on the QAP: for each value of  $\alpha$  we average the best solution of each run. The purpose here is to minimize the fitness function values. The performances are growing with  $\alpha$  and then fall down as  $\alpha$  is getting closer to its limit value. The best average performance is achieved for  $\alpha = 0.86$ . This threshold probably corresponds to a good exploration/exploitation trade-off: the algorithm favors propagation of good solutions in the vertical direction with few interactions on the left or the right sides. This kind of dynamics is well adapted to this multi-modal problem as we can reach local optima on each columns of the grid and then migrate them horizontally to find new solutions. The worst average performance is observed for  $\alpha = 0$  when the apGA is a cellular GA.  $\alpha = 0.86$  corresponds to the optimal trade-off between cellular and island models for this problem, with the best migration rate between subpopulations. In our model, the migration rate is not the number of individuals which are swap between subpopulations, but the probability for the selection operator to choose two individuals from separate columns: two individuals from separate subpopulations would then share information. We can tell that there is an optimal migration rate that is induced by the value of the anisotropic degree  $\alpha$ . Performances would probably improve if the migration rate did not stay static during the search process. As in [1], we can define some criteria to self-adjust the anisotropic degree along generations.

## 4 Diversity in apGAs

To understand better why we observe influence of the anisotropic parameter on performances, we felt it is important to measure genetic diversity during runs. We studied changes in diversity during runs according to the whole grid, the rows and the columns.

This section presents measures on population diversity in an apGA for the QAP. We conducted experiences on the average population diversity observed along generations on 100 independent runs for each anisotropic degree. We made three measures on the population diversity. First, we computed the global population diversity gD:

$$gD = \left(\frac{1}{\sharp r \sharp c}\right)^2 \sum_{r_1, r_2} \sum_{c_1, c_2} d(x_{r_1c_1}, x_{r_2c_2})$$

where  $d(x_1, x_2)$  is the distance between individuals  $x_1$  and  $x_2$ . The distance used is inspired from the Hamming distance: It is the number of locations that differs between two individuals divided by their length n.

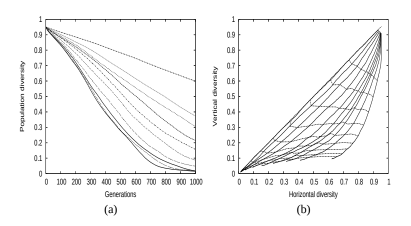
Then, we made measures on diversity inside subpopulations (vertical diversity) and diversity between subpopulations (horizontal diversity). The vertical (resp. horizontal) diversity is the sum of the average distance between all individuals in the same column (resp. row) divided by the number of columns (resp. rows):

$$vD = \frac{1}{\sharp r} \frac{1}{\sharp c^2} \sum_r \sum_{c_1, c_2} d(x_{rc_1}, x_{rc_2})$$
$$hD = \frac{1}{\sharp c} \frac{1}{\sharp r^2} \sum_c \sum_{r_1, r_2} d(x_{r_1c}, x_{r_2c})$$

where  $\sharp r$  and  $\sharp c$  are the number of rows and columns in the grid.

Figure 4(a) shows the average global diversity observed on the 1000 first generations during 100 runs on the QAP. The curves from bottom to top correspond to increasing values of  $\alpha$  from zero to nearly one. Experiments measuring genetic diversity show that small migration rate ( $\alpha$  close to one) causes islands to dominate others and retain global diversity without being able to exchange solutions to produce better results. At the opposite, for the cellular model, as  $\alpha$  is closed to zero, global diversity falls near to zero after 800 generations causing premature convergence and negatively affects performances (see figure 3). Analysis on the QAP show the necessity of maintaining diversity to produce new results and the necessity to have enough information exchanges between columns.

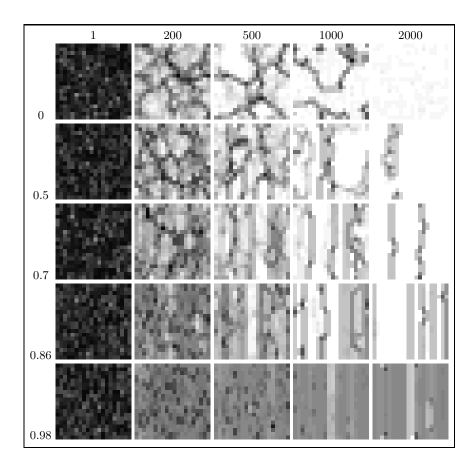
Figure 4(b) represents the vertical diversity against the horizontal diversity. The contour lines plotted every 100 generations give some information on the speed of decrease of diversity. The more the migration rate decreases (i.e.  $\alpha$  increases), the more



**Fig. 4.** Global population diversity against generation, with increasing  $\alpha$  from bottom to top (a) and vertical diversity against horizontal diversity, with increasing  $\alpha$  from left to right (b).

the diversity is maintained on each row and subpopulations converge in each column. The vertical and horizontal diversities are decreasing with the same speed for the cellular model ( $\alpha = 0$ ) and lower number of interactions between subpopulations helps the algorithm to maintain diversity on the rows when  $\alpha$  is high.

Figure 5 shows snapshots of the population diversity during one single run at different generations. The snapshots are taken from left to right at generations 1, 200, 500, 1000 and 2000. The parameter  $\alpha$  takes values in  $\{0, 0.5, 0.7, 0.86, 0.98\}$  from top to bottom. Each snapshot shows the genotypic diversity in the neighborhoods of all cells on the grid. Color black means maximum diversity and color white means that there is no more diversity in the cell's neighborhood. Those snapshots help to understand the influence of the anisotropic selection on the genotypic diversity. First, we can see that the anisotropic degree influences the dynamic of propagation of good individuals on the grid. This propagation is the cause of the loss of diversity in the population. In the standard cellular model ( $\alpha = 0$ ), good individuals propagate roughly circularly. If we slightly privilege the vertical direction ( $\alpha = 0.5$ ) the circles become elliptical. As  $\alpha$  increases, the dynamic changes and good individuals propagate column by column. For extreme values of the anisotropic degree ( $\alpha$  close to 1) the migration rate is so low that good individuals are stuck in the subpopulations and the sharing of genetic information with other subpopulations is seldom observed. In that case, the selective pressure is too low and it negatively affects performances. The crossover operator doesn't have any effect in the white zones, since they represent cells with no more diversity in their neighborhoods. For the standard cellular case, interactions between cells may have some effects on performances only at the frontier between the circles. It represents a litthe proportion of cells on the grid after a thousand generations. For  $\alpha = 0.86$ , we can see vertical lines of diversity, which means that good individuals appear in each subpopulations. For example, when we see two adjacent columns colored in grey it means that



**Fig. 5.** Local diversity in the population along generations (left to right) for increasing  $\alpha$  (top to bottom)

those columns have been colonized by two different individuals. At generation 2000, a good individual has colonized the left of the grid but he still can share information with individuals in the grey zones. This means that the migration rate between subpopulations is strong enough to guarantee the propagation of the genetic information through the whole grid. This study showed that the dynamic of the propagation of individuals on the grid is strongly related to the anisotropic degree. Once again, it would be interesting to see what kind of dynamic appears if we define a local criteria to auto-adapt  $\alpha$  during a run. This parallel model of GA allows to tune separately the anisotropic degree for each cell on the grid and measures during the search process can help to adjust locally the selective pressure.

## **Conclusion and Perspectives**

This paper presents a unified model of parallel Genetic Algorithms where granularity can be continuously tuned from fine grain to coarse grain parallel model. This family is based on the new concept of anisotropic selection. We analysed the dynamics of this class of pGAs on the well-known QAP problem. We have shown that the anisotropic degree plays a major role with regard to the average fitness found. Performances of the apGA increases with  $\alpha$  until a threshold value ( $\alpha = 0.86$ ). After this threshold, the migration rate between subpopulations in columns may be too small to generate good solutions. A study on local diversity shows the interactions between cells for different tunings of the apGA. The dynamic of propagation of individuals, which is strongly related to the genotypic diversity in the population, is dependent from the anisotropic degree of the apGA. Propagation of good individuals is done in circles for low values of  $\alpha$  and turns to vertical lines for high values of  $\alpha$ . Diversity is maintained in the population when the anisotropic degree is high, but when it reaches values close to the extreme case the few interactions between columns penalize the performances of the algorithm. These experimental results lead us to suggest to adjust dynamically the migration ratio during a run: by tuning the control parameter  $\alpha$ , it would be possible to make the algorithm to self-adjust the migration level, depending on global or local measures. While theorical and experimental studies on island models are difficult due to their complexity, the apGA model could be used as a simple framework for calculations on parallel GA. Naturally it would be worth seeing how properties described in this paper extend for even more complex problems.

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