Resource Sharing and Coevolution in Evolving Cellular Automata

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Abstract

Evolving one-dimensional cellular automata (CAs) with genetic algorithms has provided insight into how improved performance on a task requiring global coordination emerges when only local interactions are possible. Two approaches that can affect the search efficiency of the genetic algorithm are coevolution, in which a population of problems—in our case, initial configurations of the CA lattice—evolves along with the population of CAs; and resource sharing, in which a greater proportion of a limited fitness resource is assigned to those CAs which correctly solve problems that fewer other CAs in the population can solve. Here we present evidence that, in contrast to what has been suggested elsewhere, the improvements observed when both techniques are used together depend largely on resource sharing alone.

1 Introduction

Using evolutionary algorithms to design problem-solving strategies often involves the use of test cases to estimate fitness, since the space of possible strategies is typically too large to evaluate exhaustively. An important issue for improving statistical estimates of fitness in such situations is how to sample test cases and then weight their contribution to fitness estimates. This is particularly significant if one wishes to avoid premature convergence, in which a mediocre solution strategy with no nearby fitter variants takes over the population and prevents the emergence of better solutions.

Techniques that have been proposed to ameliorate this difficulty include shared sampling, in which test cases are chosen so as to be unsolvable by as many of the strategies in the population as possible [11, 10]; competitive fitness functions, in which a tournament-style selection scheme determines that one strategy is fitter than another if the number of test cases solved by the first, but not by the second, is greater than the number solved by the second, but not the first [6]; and resource-sharing fitness functions, in which strategies receive a higher fitness if they are able to solve test cases that are unsolvable by a large fraction of other strategies [6, 11, 10].

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The motivation behind resource sharing is to promote diversity by rewarding strategies that can solve test cases that few other strategies are also able to solve. In this way strategies receive less payoff for pursuing approaches that put them into “niches” already heavily occupied. Instead they are encouraged to explore new approaches, particularly those which allow solving test cases that the population as a whole finds difficult. Presumably, the population ends up more spread out over the space of possible strategies. In other words, resource sharing is intended to preserve diversity, to prevent mediocre solutions from taking over the population, and to make more likely the emergence of exceptional new strategies through recombinations of previously discovered strategies.

Another technique that has been proposed to improve the strategies discovered by evolutionary search methods is that of coevolution, as introduced by Hillis [4]. Any particular static method for generating test cases strongly affects the evolutionary course of strategies. Moreover, there appears to be no single best method. If the test cases are too easy, there is no strong pressure for high-performance strategies to emerge; if the test cases are too hard, then all low-performance strategies appear equally poor, reducing fitness variance, and evolution cannot proceed.

In a coevolutionary scheme, a population of test cases is maintained and evolves along with the evolving population of strategies. The fitness of a strategy is then some function of the number of test cases it is able to solve and the fitness of a test case is some inverse function of the number of strategies that are able to solve it, often with some penalty for being too difficult a test. The desired effect is that the test-case population will evolve so as to present an incrementally increasing but appropriate level of difficulty for the evolving population that forces strategies to become successively more capable of solving hard problems.

Past work [9], in accord with our own earlier investigations, showed that a straightforward version of coevolution, on its own, fails to produce high-performing strategies in a task in which cellular automata were evolved to perform a computation. The two populations—candidate solutions and a set of test cases—enter temporal oscillations in which each in turn performs well against the other population. The individuals in both populations, however, generally perform poorly against opponents chosen from outside the populations. Resource sharing has produced more promising results on other tasks [5, 6, 10, 11].

Combinations of different approaches for improving performance often work better than each approach alone [11]. In particular, Juillé and Pollack [7, 8] recently investigated a combination of coevolution and resource sharing in evolving cellular automata (CAs) to perform a density classification task [1, 2, 3], where the CAs play the part of the strategies, and the initial configurations of the CA lattice act as the test cases. Juillé and Pollack found that using both approaches led to the production of significantly better CA strategies than did the use of neither approach. They attributed this success to the effectiveness of coevolution.

Since a somewhat different version of coevolution, acting alone, has been shown not to produce effective strategies for this problem [9], it seems natural to ask whether Juillé and Pollack’s success is due more to coevolution or to resource sharing, or to their particular combination of the two.

2 Methods and Results

In Refs. [7] and [8], Juillé and Pollack described results of combining resource sharing and coevolution in the evolving cellular automata framework of Crutchfield, Das, and Mitchell [1, 2, 3]. In that framework, a genetic algorithm (GA) was used to evolve cellular automata
rule tables (strategies) to perform a density classification task. The fitness of each strategy was a function of its classification performance on a random sample of test cases: initial configurations (ICs) of the CA lattice. The ultimate success of the GA was measured in terms of (1) the performances of the best evolved strategies—their classification performance on larger, more difficult sets of test cases than were used to calculate fitness during evolution, and (2) the GAs search efficiency: the percentage of runs on which high-performance strategies were evolved; see Ref. [2] for details.

In Refs. [1, 2, 3], we identified three classes of CA computational strategy evolved by the GA, only one of which (particle-based) resulted in high performance and generalized well to large lattice sizes. On 149-cell lattices, these sophisticated strategies had performances \( P \) of 0.7 or greater. We define search efficiency \( \mathcal{E}_P \) as the percentage of runs on which strategies with performance \( P \) are evolved (i.e., at least one CA with performance \( \geq P \) appears in the population). In our original experiments, \( \mathcal{E}_{0.7} \) was approximately 3%. For reference, we note that to date the best known CAs for density classification, evolved or designed by hand, have performances above 0.8 (on a scale of 0.0 to 1.0).

Juillé and Pollack showed that a particular combined form of resource sharing and coevolution resulted in strategies with higher performance \( (P \approx 0.86) \) \([7, 8]\) and a larger percentage of GA runs that produced high-performance \( (P \geq 0.7) \) strategies; namely, search efficiencies above 30% (H. Juillé, personal communication).

As noted above, coevolution alone produced only low-performance \( (P << 0.7) \) CAs \([8]\). The results are substantially worse than those of a GA alone—that is, a GA without coevolution and without resource sharing. In particular, coevolution does not produce high-fitness particle-based CAs and so results in a search efficiency \( \mathcal{E}_{0.7} \) of 0%. Thus, we need not consider this alternative GA further.

The experiments described here used GA and CA parameters, resource sharing fitness functions, and a coevolution scheme identical to those of Juillé and Pollack. The experiments were designed to probe their effects in more detail than reported in Refs. [7] and [8].

The populations of CAs and ICs each had 200 members. The CAs were tested on 149-cell lattices. We performed three experiments, each consisting of 50 GA runs initiated with independent random number seeds, where each run consisted of 1000 generations. The experiments evaluated three search techniques: (1) the GA alone, with neither resource sharing nor coevolution, with ICs drawn from a density-uniform distribution (any IC density was equally probable) (the same algorithm was used in Refs. [1, 2, 3] but with smaller values for population size and number of generations, and subsequent lower search efficiencies); (2) the GA with resource sharing only, with ICs drawn from a density-uniform distribution; and (3) the GA with resource sharing and coevolution combined. Under this alternative, ICs were initially drawn from a density-uniform distribution and allowed to evolve thereafter.

For each experiment, we recorded the number of runs in which some individual reached a performance level \( P \) or greater for four values of \( P \): 0.65, 0.7, 0.75, and 0.8. From this we estimated the mean generation \( t_P \) of first occurrence of threshold \( P \). The results are given in Table [1]. The standard deviation \( \sigma_{t_P} \) of \( t_P \) across the 50 runs of each alternative GA is also reported there. In addition, the table gives each alternative’s observed search efficiencies.

The results for experiments involving resource sharing agree, within statistical uncertainty, with results found by Juillé for the percentage of runs in which a CA exceeding these thresholds occurs and for the mean first generation of such occurrences (H. Juillé, personal communication).

Runs with resource sharing alone produce CAs with higher performance more consistently.
Table 1: Statistics for the evolutionary emergence of CAs with performance $P$ exceeding various thresholds. The percentage of the 50 runs reaching threshold $P$ is given as an estimate of each alternative GA’s search efficiency $E_P$. $t_P$ is the mean number of generations to first occurrence of performance $P$ estimated across 50 GA runs. $\sigma_{t_P}$ is the standard deviation in $t_P$ estimated across the runs. $s$ is the rate of change in population diversity $\langle d \rangle$ (quoted in bits per 1000 generations) and $\sigma_s$ its standard deviation estimated in the least-squares fits of Figs. 1(a), 1(b), and 1(c).
performance. While that latter improvement is helpful, we conclude that resource sharing, rather than coevolution, is the key technique leading to improvements in the performance of CAs evolved by the GA and in the frequency of their discovery across GA runs.

3 The operation of resource sharing

We may ask further whether the effectiveness of resource sharing is actually due, as was intended in its design, to a preservation of diversity in the respective populations, or whether its success results from some other mechanism entirely.

One rough measure of diversity in a population is the average pairwise genetic Hamming distance $\langle d \rangle$. The Hamming distance $d$ between two CAs is simply the number of bits by which the genetic specification of their update rules differ. CAs with different strategies are likely to differ in more bits and thus to be separated by a greater Hamming distance than CAs with similar strategies. When averaged over the population, $\langle d \rangle$ is greater if a population is more strategically diverse overall and its members are more spread out across the genotype space. Concerned about possible long-tailed distributions governing $d$, we estimated median, in addition to average, pairwise Hamming distances. There was no qualitative change to the results. Moreover, the median distance never differed from the average by more than a single bit after the first few generations. For these reasons, we report here only average Hamming distances.

To give a sense of the scale of Hamming distances here, we recall several facts about the type of CA evolved. The state of an individual cell in one of our CAs is determined by its own state and the states of its six nearest neighbors at the previous time step. There are then 7 bits in the input to a CA’s rule table. Since the cell states are binary, the rule table is specified by $2^7 = 128$ output bits. Thus, $0 \leq \langle d \rangle \leq 128$.

Figure 1 shows $\langle d \rangle$ at each generation for runs—typical of those that evolved high-fitness particle-based CAs—with (a) neither resource sharing nor coevolution, (b) resource sharing alone, and (c) both techniques. In all cases, $\langle d \rangle$ starts out very large ($\langle d \rangle \approx 128/2 = 64$ bits), since the initial CA population was randomly initialized. $\langle d \rangle$ quickly decreases over a few generations, as the fittest CAs and their descendants take over the population, which settles down to CAs with similar strategies.

Beyond the transient phase, over each run $\langle d \rangle$ fluctuates about 10% to 20% as evolution progresses. Nonetheless, as the plots show, each run does follow an overall trend in population diversity. We measured these trends using a least-squares fit to estimate the average rate $s$ of change in population diversity. We also estimated the standard deviation $\sigma_s$ of the fit. Both estimates for each run are reported in Table 1. Since such trends are interrupted as the GA discovers progressively more effective CAs, the fits were made only over a stationary "epoch" (a period in which average population fitness remains roughly constant). For Fig. 1(a), the fit is made for an epoch lasting from generation 50 to generation 800; in Fig. 1(b), from generation 200 to 1000; and in Fig. 1(c), from 100 to 750.

In runs with neither resource sharing nor coevolution, $\langle d \rangle$ decreases slowly over time, with temporary increases each time a new, more effective type of strategy is discovered. In Figure 1(a), for example, $\langle d \rangle$ declines slowly over nearly 800 generations, from about 18 bits to a minimum of approximately 12 bits. At that point, a new strategy appears around generation 900 and $\langle d \rangle$ increases again. The estimated trend shows a negative slope and one concludes that with this type of GA population diversity steadily decreases in an epoch.

In contrast, when the GA with resource sharing is used, as illustrated by the run in Fig.
\(\langle d \rangle\) remains roughly constant around 17.5 bits. There also appears to be wider fluctuation in the population diversity about this trend than in the alternative GAs; see \(\sigma_s\) in Table 1.

When coevolution is added to resource sharing, as shown by the run in Fig. 1(c), \(\langle d \rangle\) increases over time, after the population initially settles down. \(\langle d \rangle\) typically goes from about 15 bits to about 21 bits over the course of 700 generations.

It would appear then that resource sharing maintains diversity, as it was intended to do. Its use prevents the slow decrease in total Hamming distance that otherwise occurs as the population converges on a narrower range of strategies. In other words, it maintains a wider variation in the space of CAs. The addition of coevolution appears to enhance the effect of resource sharing, when the latter is also used: the total Hamming distance increases and reaches noticeably higher values over a similar number of generations.

4 Further Work

If coevolution does, in fact, augment the operation of resource sharing, as suggested by the above statistical analysis, it remains unclear exactly how it acts to do so. A more detailed analysis of how the GA with resource sharing is affected by the addition of coevolution, and why coevolution is useful when it accompanies resource sharing but not on its own, will be necessary for a fuller understanding of the trade-offs between these alternative evolutionary search techniques. This is motivated, of course, by the marked and very useful improvements shown by the augmented GAs over the GA alone.

Underlying these overall concerns and determining, in large measure, the statistical analysis just reported is a complicated problem in nonlinear population dynamics. High-performance CAs evolve via a series of epochs of stasis punctuated by sudden innovations, whether resource sharing, resource sharing and coevolution, or neither are employed. The dynamics of epochal evolution has recently been mathematically analyzed in some detail; see Refs. [12, 13, 14]. It would be useful, therefore, to bring the current investigations together with this mathematical analysis to understand why epochal evolution with resource sharing results in higher variance in the time it takes moderate- and high-performance CAs to emerge, why higher-performance CAs appear more frequently with resource sharing, and how it is that coevolution increases these effects. Here we have begun to understand more systematically how resource sharing and coevolution affect the evolutionary process, but not yet their underlying mechanisms.

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Figure 1: Average pairwise Hamming distance $\langle d \rangle$ over time for single GA runs with (a) neither resource sharing nor coevolution, (b) resource sharing alone, (c) resource sharing and coevolution. $\langle d \rangle$ is large ($\approx 64$) during the initial generations, and so these data points do not appear on the scales plotted. The straight lines show the trends in population diversity. They are least-squares fits over stationary fitness epochs in the population dynamics. The estimated slopes $s$ and their standard deviations $\sigma_s$ are quoted in Table 1. The runs shown here are typical of those that evolved high-fitness particle-based CAs under each alternative GA.