Overcoming Problems in the Measurement of Biological Complexity

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In a genetic algorithm, fluctuations of the entropy of a genome over time are interpreted as fluctuations of the information that the genome’s organism is storing about its environment, being this reflected in more complex organisms. The computation of this entropy presents technical problems due to the small population sizes used in practice. In this work we propose and test an alternative way of measuring the entropy variation in a population by means of algorithmic information theory, where the entropy variation between two generational steps is the Kolmogorov complexity of the first step conditioned to the second one. As an example application of this technique, we report experimental differences in entropy evolution between systems in which sexual reproduction is present or absent.

I. INTRODUCTION

The evolution over time of the entropy of a genome within a population is currently an interesting problem which is conjectured to be connected to the evolution of the complexity of organisms in a genetic algorithm [1, 2]. The complexity of the genome of an organism is considered to be the amount of information about its environment it stores. That is, evolution would cause the appearance of more complex sequences, which correspond to more complex phenotypes. This hypothesis states that natural selection acts as a Maxwell demon, accepting only those changes which adapt better to the environment and give rise to more complex individuals with genomes of lower entropy. This idea was tested by the simulation of a very simple system of asexual individuals in fixed environmental conditions.

However, it is well known that that the computation of the entropy as

$$H(\text{genome}) = - \sum_{x \in \text{genomes in the population}} p(x) \log p(x)$$

(1)

has technical complications, due to the large size of the sample needed to estimate it with accuracy [1, 3, 7]. In practice, it is usually estimated as

$$H(\text{genome}) = \sum_{i=1..\text{size(genome)}} H(i),$$

(2)

i.e., as the sum of the entropy contributions of each locus in the genome. This estimation misses the entropy contributions due to epistatic effects. Some sophisticated statistical methods can be used to remedy this (see Appendix in [2]), although we will not deal with them in this work.

An still unexplored way to overcome this problem is to estimate the entropy of a genome as its average Kolmogorov complexity

$$\langle K(\text{genome}) \rangle_{\text{genomes in the population}} \rightarrow H(\text{genome}),$$

(3)

(see [5, 8, 9]). However, this result only holds for infinitely long sequences, and therefore it cannot be applied to finite (sometimes short) genomes.

If we are only interested in the entropy evolution of the genome, and not in the particular value estimation, we can resort to the following trick: the genetic algorithm can be modelled as a thermodynamic system which evolves over time, where every population is just a measurement by an observer, i.e. the system is modeled as a statistical ensemble of genomes and each measurement is just a sample from that ensemble.

Now we can measure the entropy evolution of the system from two different viewpoints. The first is the system itself, where the entropy is calculated a la Shannon (equation 2) by estimating the probabilities of the loci alleles, using the frequencies of the ensemble sample.

The second way of measuring the entropy is from the viewpoint of the observer, where a measurement is made of the population at each time step, and the information about the system is updated, i.e., the observer measures the system at time \(t\) and stores this information \(S_t\). At time \(t+1\) the observer makes another measurement and substitutes \(S_t\) by \(S_{t+1}\). The entropy variation due to this substitution can be calculated for both equilibrium and non-equilibrium thermodynamic systems [4, 12, 14]. Since evolution cannot be modeled as a system in equilibrium, the second case applies: the mutation and generational replacement operators may increase or decrease the entropy of the system [10, 11].

Thus the entropy variation from the observer viewpoint is \(K(S_t) - K(S_{t+1})\) bits. As \(K(\cdot)\) is an incomputable measure, we estimate it by using the Lempel-Ziv algorithm [12] as \(\lim_{n \to \infty} (1/n) |LZ(x)| = (1/n)K(x[n])\), where \(x\) is an infinite string and \(|LZ(x)|\) is the size of the same string compressed [5]. Now our measurement of the system at time \(t\), \(|S_t|\) is much larger than with equation 3 (just the genome), so the estimation becomes possible.

II. EXPERIMENTS

We want to test whether the evolution of \(K(\text{population sample})\) can help in the study of the evolution of \(H(\text{genome})\). Both measurements have their
limitations, but their agreement would provide evidence that the entropy evolution is being studied correctly.

We have evaluated this experimentally, using the genetic algorithm proposed by [6], which is able to reproduce sexual behaviour in a very detailed way, because it includes several features absent in the [2] experiments, such as sexual reproduction, different inter-locus and intra-locus interactions across the genotypic or phenotypic distance, and the evolutionary mechanisms of mutation and natural selection.

We have implemented and run the same simulation proposed by Hayashi et al. The model’s sexual dynamics can be summarized as follows: there are two different sexes (male and female). The likelihood that a female with trait $a$ will mate with a male with trait $b$ is defined by $\psi(d) = e^{-\alpha d^2}$, where $d$ is the genetic or phenotypic distance measuring compatibility between the sexes, and $\alpha$ is a parameter that represents the compatibility between traits. The value of $\alpha$ used in the simulations is 0.005.

The overall number of offspring produced by a female in each coupling is given by $W_f = B_{max}e^{-sc(P-P_{opt})^2}$, where $P$ is the proportion of males with which the female has been able to mate. The parameter $P_{opt}$ (which can take the values $\{0.2, 0.4, 0.6, 0.8\}$) defines the fertility of the female. The parameter $sc$ (ranging from 1.02 to $4 \times 1.02$) stands for sexual conflict selection in females. $B_{max}$ is the maximum possible number of offspring (5). The sex and the father are randomly chosen for each offspring, and the number of males each female encounters is $n = 20$.

Twenty two different experiments have been performed. $K$(population sample) and $H$(genome) have been estimated for each generation step with the methods described above. The system evolved for 10,000 generations with a population size of 1,000 individual genomes.

In all our results, both measures were highly correlated (see fig. 1), giving evidence that the dual measurement of the evolution of entropy by means of Kolmogorov complexity confirms the use of the Shannon entropy.

III. DISCUSSION

When sexual dynamics are introduced in the system, the increase of complexity observed by [2] is not present anymore.

The typical result observed in our experiments is a chaotic behavior of the entropy (fig. 2). Only large
FIG. 4: Example of genetic differentiation without co-evolutionary chase or sympatric speciation, obtained with parameters $B_{\text{max}} = 5$, $P_{\text{opt}} = 0.2$, $\mu = 5.0 \times 10^{-5}$, $\alpha = 0.05$, $sc = 1.02$, 32 loci, phenotypic model: dominance. Experimental correlation coefficient: 0.8571.

Our hypothesis for this behavior of the entropy is the absence of natural selection in the Hayashi et al. [6] model, which could explain the similarities between male and female evolution (fig. 5). Without natural selection, the environment for females is reduced to random boundary conditions (mutations). On the other hand, males are selected by females as mating partners. In this way, females can be considered to become the environment for males, since they determine the way in which the entropy of the males evolves. On the other hand, females have no environment to adapt to. Perhaps if the pressure of natural selection was applied both to male and female (not necessarily in the same way), more complex patterns in the behavior of their entropies would appear. The fact that natural selection is not taken into account may be the cause of the differences in entropy evolution between the models by [2] and [6], and the reason why global decreases in entropy are not observed in the latter.

IV. CONCLUSIONS AND FUTURE WORK

Studying a genetic algorithm from the observer point of view allows us to have large-scale estimates of the entropy evolution via Kolmogorov Complexity. This overcomes many limitations that arise from epistatic effects between loci and provide an easy way to do study the dynamics without resorting to complex mathematical trickeries. We also use this methodology to study what is the effect of sexual reproduction in terms of the evolution of complexity, as a decrease of entropy. We show that when sexual reproduction is present the population enters in a chaotic regime of complexity driven by the complexity drifts of the female organisms. We suggest that this might be cause for female organism evolving chaotically without natural selection, and male organisms evolving with females as the boundary conditions, which gives an overall chaotic evolution of complexity. The next immediate step is to introduce natural selection in the experiments and study whether this will change the evolution of complexity. We plan to do it by implementing the typical natural selection operators from genetic algorithms such as tournament selection, steady state-selection and so forth. We conjecture that introducing natural selection will remove the chaotic complexity dynamics and might probably get closer to what Hayasi et. al. formerly reported: an increase in complexity by removing genetic mutations that do not improve the fitness function.

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