The Evolutionary Unfolding of Complexity

James P. Crutchfield and Erik van Nimwegen

Santa Fe Institute, 1399 Hyde Park Road, Santa Fe, New Mexico 87501. {chaos,erik}@santafe.edu.

Abstract. We analyze the population dynamics of a broad class of fitness functions that exhibit epochal evolution—a dynamical behavior, commonly observed in both natural and artificial evolutionary processes, in which long periods of stasis in an evolving population are punctuated by sudden bursts of change. Our approachstatistical dynamics—combines methods from both statistical mechanics and dynamical systems theory in a way that offers an alternative to current "landscape" models of evolutionary optimization. We describe the population dynamics on the macroscopic level of fitness classes or phenotype subbasins, while averaging out the genotypic variation that is consistent with a macroscopic state. Metastability in epochal evolution occurs solely at the macroscopic level of the fitness distribution. While a balance between selection and mutation maintains a quasistationary distribution of fitness, individuals diffuse randomly through selectively neutral subbasins in genotype space. Sudden innovations occur when, through this diffusion. a genotypic portal is discovered that connects to a new subbasin of higher fitness genotypes. In this way, we identify innovations with the unfolding and stabilization of a new dimension in the macroscopic state space. The architectural view of subbasins and portals in genotype space clarifies how frozen accidents and the resulting phenotypic constraints guide the evolution to higher complexity.

Keywords: punctuated equilibrium, neutrality, epochal evolution, statistical mechanics, dynamical systems

Santa Fe Institute Working Paper 99-02-015

To appear in **Evolution as Computation**, L. F. Landweber, E. Winfree, R. Lipton, and S. Freeland, editors, Lecture Notes in Computer Science, Springer-Verlag (1999). Proceedings of a DIMACS Workshop, 11-12 January 1999, Princeton University.

1 Evolutionary Computation Theory

The recent mixing of evolutionary biology and theoretical computer science has resulted in the phrase "evolutionary computation" taking on a variety of related but clearly distinct meanings.

In one view of evolutionary computation we ask whether Neo-Darwinian evolution can be productively analyzed in terms of how biological information is stored, transmitted, and manipulated. That is, Is it helpful to see the evolutionary process as a computation?

Instead of regarding evolution itself as a computation, one might ask if evolution has produced organisms whose internal architecture and dynamics are capable *in principle* of supporting arbitrarily complex computations. Landweber and Kari argue that, yes, the information processing embedded in the reassembly of fragmented gene components by unicellular organisms is quite sophisticated; perhaps these organisms are even capable of universal computation [31]. It would appear, then, that evolved systems themselves must be analyzed from a computational point of view.

Alternatively, from an engineering view we can ask, Does Neo-Darwinian evolution suggest new approaches to solving computationally difficult problems? This question drives much recent work in evolutionary search—a class of stochastic optimization algorithms, loosely based on processes believed to operate in biological evolution, that have been applied successfully to a variety of different problems; see, for example, Refs. [4,6,8,11,16,20,22,30,33] and references therein.

Naturally enough, there is a middle ground between the scientific desire to understand how evolution works and the engineering desire to use nature for human gain. If evolutionary processes do embed various kinds of computation, then one can ask, Is this biological information processing of use to us? That is, can we use biological nature herself to perform computations that are of interest to us? A partial, but affirmative answer was provided by Adelman, who mapped the combinatorial problem of Directed Hamiltonian Paths onto a macromolecular system that could be manipulated to solve this well known hard problem [2].

Whether we are interested in this middle ground or adopt a scientific or an engineering view, one still needs a mathematical framework with which to analyze how a population of individuals (or of candidate solutions) compete through replication and so, possibly, improve through natural (or artificial) selection. This type of evolutionary process is easy to describe. In the Neo-Darwinian view each individual is specified by a genotype and replicates (i) according to its fitness and (ii) subject to genetic variation. During the passage from the population at one generation to the next, an individual is translated from its genotypic specification into a form, the phenotype, that can be directly evaluated for fitness and so selected for inclusion in the next generation. Despite the ease of describing the process qualitatively, the mechanisms constraining and driving the population dynamics of evolutionary adaptation are not well understood.

In mathematical terms, evolution is described as a nonlinear populationbased stochastic dynamical system. The complicated dynamics exhibited by such systems has been appreciated for decades in the field of mathematical population genetics [24]. For example, the effects on evolutionary behavior of the rate of genetic variation, the population size, and the genotype-to-fitness mapping typically cannot be analyzed separately; there are strong, nonlinear interactions between them. These complications make an empirical approach to the question of whether and how to use evolutionary optimization in engineering problematic. They also make it difficult to identify the mechanisms that drive behavior observed in evolutionary experiments. In any case, one would like to start with the basic equations of motion describing the evolutionary process, as outlined in the previous paragraph, and then predict observable features—such as, the time to find an optimal individual—or, at a minimum, identify mechanisms that constrain and guide an evolving population.

Here we review our recent results that address these and similar questions about evolutionary dynamics. Our approach derives from an attempt to unify and extend theoretical work that has been done in the areas of evolutionary search theory, molecular evolution theory, and mathematical population genetics. The eventual goal is to obtain a more general and quantitative understanding of the emergent mechanisms that control the population dynamics of evolutionary adaptation and that govern other population-based dynamical systems.

2 Epochal Evolution

To date we have focused on a class of population-dynamical behavior that we refer to as *epochal evolution*. In epochal evolution, long periods of stasis (*epochs*) in the average fitness of the population are punctuated by rapid *innovations* to higher fitness. These innovations typically reflect an increase of complexity—that is, the appearance of new structures or novel functions at the level of the phenotype. One central question then is, How does epochal evolutionary population dynamics facilitate or impede the emergence of such complexity?

Engineering issues aside, there is a compelling biological motivation for a focus on epochal dynamics. There is the common occurrence in natural evolutionary systems of "punctuated equilibria"—a process first introduced to describe sudden morphological changes in the paleontological record [23]. Similar behavior has been recently observed experimentally in bacterial colonies [15] and in simulations of the evolution of t-RNA secondary structures [18]. This class of behavior appears sufficiently general that it occurs in artificial evolutionary systems, such as evolving cellular automata [10,34] and populations of competing self-replicating computer programs [1]. In addition to the increasing attention paid to this type of epochal evolution in the theoretical biology community [18,21,26,35,41,49], recently there has also been an increased interest by evolutionary search theorists [5,25]. More directly, Chen et al. recently proposed to test our original theoretical predictions in an experimental realization of a genetic algorithm that exhibits epochal evolution [9].

4

2.1 Local Optima versus Neutral Subbasins

How are we to think of the mechanisms that cause epochal evolutionary behavior? The evolutionary biologist Wright introduced the notion of "adaptive landscapes" to describe the (local) stochastic adaptation of populations to themselves and to environmental fluctuations and constraints [50]. This geographical metaphor has had a powerful influence on theorizing about natural and artificial evolutionary processes. The basic picture is that of a gradient-following dynamics moving over a "landscape" determined by a fitness "potential". In this view an evolving population stochastically crawls along a surface determined, perhaps dynamically, by the fitness of individuals, moving to peaks and very occasionally hopping across fitness "valleys" to nearby, and hopefully higher fitness, peaks.

More recently, it has been proposed that the typical fitness functions of combinatorial optimization and biological evolution can be modeled as "rugged landscapes" [28,32]. These are fitness functions with wildly fluctuating fitnesses even at the smallest scales of single-point mutations. Consequently, it is generally assumed that these "landscapes" possess a large number of local optima. With this picture in mind, the common interpretation of punctuated equilibria in evolving populations is that of a population being "stuck" at a local peak in the landscape, until a rare mutant crosses a valley of relatively low fitness to a higher peak; a picture more or less consistent with Wright's.

At the same time, an increasing appreciation has developed, in contrast to this rugged landscape view, that there are substantial degeneracies in the genotype-to-phenotype and the phenotype-to-fitness mappings. The history of this idea goes back to Kimura [29], who argued that on the genotypic level, most genetic variation occurring in evolution is adaptively neutral with respect to the phenotype. Even today, the crucial role played by neutrality continues to find important applications in molecular evolution, for example; see Ref. [19]. During neutral evolution, when degeneracies in the genotype-phenotype map are operating, different genotypes in a population fall into a relatively small number of distinct fitness classes of genotypes with approximately equal fitness. Due to the high dimensionality of genotype spaces, sets of genotypes with approximately equal fitness tend to form components in genotype space that are connected by paths made of single-mutation steps.

Additionally, due to intrinsic or even exogenous effects (e.g., environmental), there simply may not exist a deterministic "fitness" value for each genotype. In this case, fluctuations can induce variation in fitness such that genotypes with similar average fitness values are not distinct at the level of selection. Thus, genotype-to-fitness degeneracies can, to a certain extent, be induced by noise in the fitness evaluation of individuals.

When these biological facts are taken into account we end up with an alternative view to both Wright's "adaptive landscapes" and the more recent "rugged landscapes". That is, the genotype space decomposes into a set

of neutral networks, or *subbasins* of approximately isofitness genotypes, that are entangled with each other in a complicated fashion; see Fig. 1. As illus-

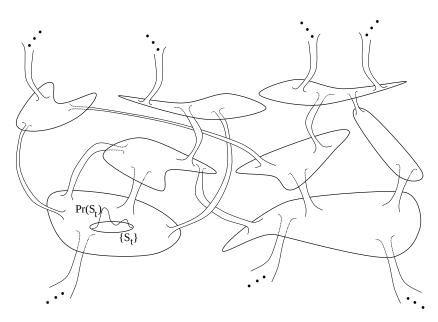


Fig. 1. Subbasin and portal architecture underlying epochal evolutionary dynamics. A population—a collection of individuals $\{S_t\}$ with distribution $Pr(S_t)$ —diffuses in the subbasins (large sets) until a portal (tube) to a higher-fitness subbasin is found.

trated in Fig. 1, the space of genotypes is broken into strongly and weakly connected sets with respect to the genetic operators. Equal-fitness genotypes form strongly connected neutral subbasins. Moreover, since subbasins of high fitness are generally much smaller than subbasins of low fitness, a subbasin tends to be only weakly connected to subbasins of higher fitness.

Since the different genotypes within a neutral subbasin are not distinguished by selection, neutral evolution—consisting of the random sampling and genetic variation of individuals—dominates. This leads to a rather different interpretation of the processes underlying punctuated equilibria. Instead of the population being pinned at a local optimum in genotype space as suggested by the "landscape" models, the population drifts randomly through neutral subbasins of isofitness genotypes. A balance between selection and deleterious mutations leads to a (meta-) stable distribution of fitness (or of phenotypes), while the population is searching through these spaces of neutral genotypic variants. Thus, there is no genotypic stasis during epochs. As was first pointed out in the context of molecular evolution in Ref. [27], through neutral mutations, the best individuals in the population diffuse over the

neutral network of isofitness genotypes until one of them discovers a connection to a neutral network of even higher fitness. The fraction of individuals on this network then grows rapidly, reaching a new equilibrium between selection and deleterious mutations, after which the new subset of most-fit individuals diffuses again over the newly discovered neutral network.

Note that in epochal dynamics there is a natural separation of time scales. During an epoch selection acts to establish an equilibrium in the proportions of individuals in the different neutral subspaces, but it does not induce *adaptations* in the population. Adaptation occurs only in a short burst during an innovation, after which equilibrium on the level of fitness is re-established in the population. On a time scale much faster than that between innovations, members of the population diffuse through subbasins of isofitness genotypes until a (typically rare) higher-fitness genotype is discovered. Long periods of stasis occur because the population has to search most of the neutral subspace before a portal to a higher fitness subspace is discovered.

In this way, we shift our view away from the geographic metaphor of evolutionary adaptation "crawling" along a "landscape" to the view of a diffusion process constrained by the subbasin-portal architecture induced by degeneracies in the genotype-to-phenotype and phenotype-to-fitness mappings. Moreover, our approach is not simply a shift towards an architectural view, but it also focuses on the *dynamics* of populations as they move through the subbasins to find portals to higher fitness.

2.2 Epochal Evolution—An Example

In our analysis [45,46], we view the subbasin-portal mechanism sketched above as the main source of epochal behavior in evolutionary dynamics. We will now discuss a simple example of epochal evolution that illustrates more specifically the mechanisms involved and allows us to introduce several concepts used in our analysis.

Figure 2 shows the fitness dynamics of an evolving population on a sample fitness function that exhibits large degeneracies in the genotype-fitness mapping. This fitness function is an example of the class of Royal Road fitness functions explained in Sec. 3 below. The genotype space consists of all bit-strings of length 30 and contains neutral subbasins of fitnesses 0, 1, 2, and 3. There is only one genotype with fitness 3, 3069 genotypes have fitness 2, 3.14×10^6 have fitness 1, and all others have fitness 0. The evolving population consists of 250 individuals that at each generation are selected in proportion to their fitness and then mutated with probability 0.005 per bit. Figure 2(a) shows the average fitness $\langle f \rangle$ in the population (lower curve) and the best fitness in the population (upper curve) as a function of generation t.

At time t = 0 the population starts out with 250 random genotypes. As can be seen from Fig. 2(a), during the first few generations all individuals are located in the largest subbasin with fitness 0, since both average and

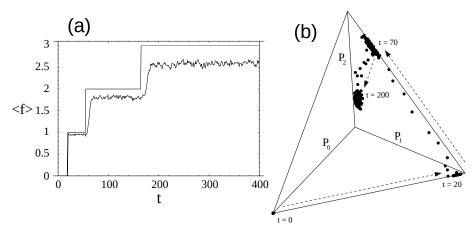


Fig. 2. Dynamics of (a) the average fitness (lower curve) and best fitness (upper curve) and (b) the fitness distribution for a population evolving under a Royal Road fitness function. The fitness function has N=3 constellations of K=10 bits each. The population size is M=250 and the mutation rate $\mu=0.005$. In (b) the location of the fitness distribution at each generation is shown by a dot. The dashed lines there indicate the direction in which the fitness distribution moves from metastable to metastable state through the population's fitness-distribution state space (a simplex). The times at which the different metastable states are first reached are indicated as well.

best fitness are 0. The population randomly diffuses through this subbasin until, around generation 20, a "portal" is discovered into the subbasin with fitness 1. The population is quickly taken over by genotypes of fitness 1, until a balance is established between selection and mutation: selection expanding and deleterious mutations (from fitness 1 to 0) decreasing the number of individuals with fitness 1. The individuals with fitness 1 continue to diffuse through the subbasin with fitness 1, until a portal is discovered connecting to the subbasin with fitness 2. This happens around generation t=60 and by t=70 a new selection-mutation equilibrium is established. Individuals with fitness 2 continue diffusing through their subbasin until the globally optimal genotype with fitness 3 is discovered some time around generation t=170. Descendants of this genotype then spread through the population until around t=200, when a final equilibrium is reached.

The same dynamics is plotted in Fig. 2(b), but from the point of view of the population's fitness distribution $\vec{P} = (P_0, P_1, P_2, P_3)$. In the figure the P_0 axis indicates the proportion of fitness 0 genotypes in the population, P_1 the proportion of fitness 1 genotypes, and P_2 the proportion of fitness 2 genotypes. Of course, since \vec{P} is a distribution, $P_3 = 1 - P_0 - P_1 - P_2$. Due to this, the space of possible fitness distributions forms a three-dimensional simplex. We see that initially $P_0 = 1$ and the population is located in the lower-left corner of the simplex. Later, between t = 20 and t = 60, the

population is located at a metastable fixed point on the line $P_0 + P_1 = 1$ and is dominated by fitness-1 genotypes $(P_1 \gg P_0)$. Some time around generation t = 60 a genotype with fitness 2 is discovered and the population moves into the plane $P_0 + P_1 + P_2 = 1$ —the front plane of the simplex. From generation t = 70 until generation t = 170 the population fluctuates around a metastable fixed point in this plane. Finally, a genotype of fitness 3 is discovered and the population moves to the asymptotically stable fixed point in the interior of the simplex. It reaches this fixed point around t = 200 and remains there fluctuating around it for the rest of the evolution.

This example illustrates the general qualitative dynamics of epochal evolution. It is important to note that the architecture of neutral subbasins and portals is such that a higher-fitness subbasin is always reachable from the current best-fitness subbasin. Metastability is a result of the fact that the connections (portals) to higher-fitness subbasins are very rare. These portals are generally only discovered after the population has diffused through most of the subbasin. Additionally, at each innovation, the fitness distribution expands into a new dimension of the simplex. Initially, when all members have fitness 0, the population is restricted to a point. After the first innovation it moves on a one-dimensional line, after the second it moves within a two-dimensional plane, and finally it moves into the interior of the full threedimensional simplex. One sees that, when summarizing the population with fitness distributions, the number of components needed to describe the population grows dynamically each time a higher-fitness subbasin is discovered. We will return to this observation when we describe the connection of our analytical approach to the theory of statistical mechanics.

3 The Terraced Labyrinth Fitness Functions

As just outlined, the intuitive view of phenotypically constrained, genotype-space architectures—as a relatively small number of weakly interconnected neutral subbasins—is the one we have adopted in our analyses. We will now define a broad class of fitness functions that captures these characteristics. The principal motivation for this is to illustrate the generality of our existing results via a wider range of fitness functions than previously analyzed.

We represent genotypes in the population as bit-strings of a fixed length L. For any genotype there is a certain subset of its bits that are fitness constrained. Mutations in any of the constrained bits lowers an individual's fitness. All the other bits are considered free bits, in the sense that they may be changed without affecting fitness. Of all possible configurations of free bits, there is a small subset of portal configurations that lead to an increased fitness. A portal consists of a subset of free bits, called a constellation, that is set to a particular "correct" configuration. A constellation may have more than one "correct" configuration. When a constellation is set to a portal configuration, the fitness is increased, and the constellation's bits become constrained bits.

That is, via a portal free bits of an incorrectly set constellation become the constrained bits of a correctly set constellation.

The general structure of the fitness functions we have in mind is that fitness is conferred on individuals by having a number of constellations set to their portal configurations. Mutations in the constrained bits of the correct constellations lower fitness; while setting an additional constellation to its portal configuration increases fitness. A fitness function is specified by choosing sets of constellations, portal configurations, and assigning the fitness that each constellation confers on a genotype when set to one of its portal configurations.

3.1 A Simple Example

Let's illustrate our class of fitness functions by a simple example that uses bit-strings of length L=15. The example is illustrated in Fig. 3. Initially, when no constellation is set correctly the strings have fitness f. The first constellation, denoted c, consists of the bits 1 through 5. This constellation can be set to two different portal configurations: either $\pi_1=11111$ or $\pi_2=00000$. When $c=\pi_1$ or $c=\pi_2$ the genotypes obtain fitnesses f_1 and f_2 , respectively. Once constellation $c=\pi_1$, say, there is a constellation c_1 , consisting of bits 9 through 15, that can be set correctly to portal configuration $\pi_{1,1}=1100010$; in which case the genotype obtains fitness $f_{1,1}$. The constellation c_1 might also be set to configuration $\pi_{1,2}=0101101$, leading to a fitness of $f_{1,2}$. Finally, once constellation $c_1=\pi_{1,1}$, there is a final configuration $c_{1,1}$, consisting of bits 6 through 8, that can be set correctly. With $c=\pi_1$ and $c_1=\pi_{1,1}$ configuration $c_{1,1}$ needs to be set to configuration $\pi_{1,1,1}=001$ in order to reach fitness $f_{1,1,1}$. If instead $c_1=\pi_{1,2}$, the final constellation $c_{1,2}$ needs to be set to portal $\pi_{1,2,1}=100$, giving fitness $f_{1,2,1}$.

Alternatively, if constellation $c=\pi_2$, the next constellation c_2 consists of bits 8 through 10, which have portal configuration $\pi_{2,1}=111$. Setting c_2 to $\pi_{2,1}$ leads to fitness $f_{2,1}$. Once c_2 is set correctly, there is a constellation $c_{2,1}$ consisting of bits 13 through 15, which has portal configuration $\pi_{2,1,1}=110$ and fitness $f_{2,1,1}$. Finally, there is the constellation $c_{2,1,1}$ consisting of bits 6, 7, 11, and 12. The portal configuration for this constellation is $\pi_{2,1,1,1}=1000$, leading to fitness $f_{2,1,1,1}$.

Generally, the hierarchical ordering of constellations and their connections via portals can be most easily represented as a tree; as in Fig. 3. Each tree node represents a subbasin of equal-fitness genotypes. The tree branches represent the portals that connect a lower-fitness subbasin to a higher-fitness subbasin. The fitness and structure of genotypes within a subbasin are also shown at each node. Stars (*) indicate the free bits within a subbasin. The constellations at each node indicate which subset of bits needs to be set to a portal configuration in order to proceed further up the tree. Thus, setting a constellation to a portal configuration leads one level up the tree, while

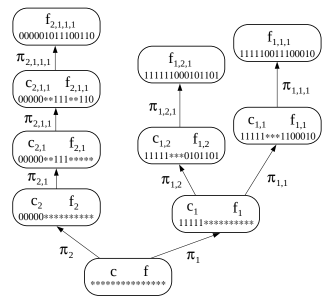


Fig. 3. Tree representation of a Terraced Labyrinth fitness function. The nodes of the tree represent subbasins of genotypes with equal fitness. They are represented by strings that have *'s for the free bits. The fitness f of the genotypes in the subbasins is indicated as well. The constellation c inside each node indicates the subset of bits that needs to be set correctly in order to move up a level in the tree to a higher-fitness subbasin. The portal configurations π that connect subbasins to higher-fitness subbasins are shown as branches.

mutating one or more of the constrained bits leads down the tree. In fact, a single point-mutation might lead all the way back to the root node.

We assume that setting a new constellation correctly leads to an increase in fitness. That is, f_1 and f_2 are larger than f, $f_{1,1}$ is larger than f_1 , and so on. For simplicity in this example, we chose the constellation bits contiguously, except for $c_{2,1,1}$. Since our genetic algorithm, introduced shortly, does not employ crossover, the population dynamics remains the same under arbitrary permutations of the bits in the genome. Note further that we chose the portal configurations rather arbitrarily. In cases where a constellation has only a single portal, this configuration can be chosen arbitrarily without effecting the dynamics. When a constellation has more than one portal, the evolutionary dynamics can be affected by the Hamming distances between the different portal configurations. A key assumption is that portal configurations such as π_1 and π_2 are mutually exclusive. Once evolution follows a certain branch up the tree, it is very unlikely to revert later on. We discuss in Sec. 8 how different evolutionary paths through the tree formalize such notions as historical accident and structural phenotypic constraints.

Finally, in this setting the genotype-to-phenotype map is nonexistent, since fitness is evaluated directly on the genotypes, without an intervening developmental process.

3.2 Definitions

We will now generalize this example by way of defining the class of *Terraced Labyrinth* fitness functions. As we saw in the example, constellations and portals form a hierarchy that can be most easily represented as a tree. Thus, we define Terraced Labyrinth fitness functions using trees, similar to the one illustrated in Fig. 3, as follows.

- 1. The genotypes are bit strings $\mathbf{s} = s_0 s_1 s_2 \cdots s_{L-1}$ of length L with bits $s_i \in \mathcal{A} \equiv \{0,1\}.$
- 2. The hierarchy of subbasins, constellations, and portals form a *tree*, consisting of nodes $\{\vec{\imath}\}$ and branches $\{\pi_{\vec{\imath}}\}$.
 - (a) Tree nodes \vec{i} are specified by a set of indices: $\vec{i} = \{i_1, i_2, i_3, \dots, i_n\}$. The number n of indices denotes \vec{i} 's tree level. A particular setting of the indices labels the path from the root to \vec{i} . That is, one reaches \vec{i} by taking branch i_1 at the root, branch i_2 at node i_1 , and so on. The tree nodes represent both subbasins of genotypes with equal fitness and constellations of bits that, when set correctly, lead out of one subbasin to the next higher-fitness subbasin.
 - (b) Tree branches represent portal configurations that connect the subbasins of equal-fitness genotypes to each other. Branch $\pi_{\vec{i}}$ points to node \vec{i} .
- 3. A constellation is a subset of s's bits. Constellation $c_{\vec{i}}$ is located at node \vec{i} and corresponds to the subset of bits that must be set to a portal configuration in order to move from subbasin $B_{\vec{i}}$ to a higher fitness subbasin. The number of bits in a constellation $c_{\vec{i}}$ is denoted $K_{\vec{i}}$.
- 4. A portal $\pi_{\vec{i},j}$ is one particular configuration of the $K_{\vec{i}}$ bits in constellation $c_{\vec{i}}$ out of the $2^{K_{\vec{i}}}$ possible configurations. The indices \vec{i} of a portal $\pi_{\vec{i},j}$ indicate the node to which it points.
- 5. The subbasin $B_{i_1,i_2,...,i_n}$ is the set of genotypes that have constellations c through $c_{i_1,...,i_{n-1}}$ set to portals π_{i_1} through $\pi_{i_1,...,i_n}$, respectively, but do not have constellation $c_{i_1,...,i_n}$ set to any of its portal configurations.
- 6. All genotypes in the subbasin $B_{\vec{i}}$ have a fitness $f_{\vec{i}}$.
- 7. A leaf-node \vec{i} in the tree represents a set of equal-fitness genotypes that form a local optimum of the fitness function. The fitness of these genotypes is $f_{\vec{i}}$.

The trees that define the hierarchy of constellations, subbasins, and portals are not entirely arbitrary. They have the following constraints.

1. The number of branches leaving node \vec{i} is at most $2^{K_{\vec{i}}}$.

2. A constellation is *disjoint* from the root constellation c and all other constellations that connect it to the root. That is, the set $c_{i_1,i_2,...,i_n}$ is disjoint from the sets c, c_{i_1} , c_{i_1,i_2} , and so on.

This class of Terraced Labyrinth fitness functions incorporates and extends the previously studied $Royal\ Road$ fitness functions of Refs. [45] and [46] and the $Royal\ Staircase$ fitness functions of Ref. [43]. In those fitness functions, all constellations had the same number of defining bits K, and there was only a single portal configuration $\pi=1^K$ for each constellation. A Royal Staircase fitness function corresponds to a Terraced Labyrinth fitness function whose tree is a simple linear chain. Additionally, in the Royal Road fitness functions, constellations were allowed to be set in any arbitrary order.

The architectural approach we have taken here should be contrasted with the use of randomized fitness functions that have been modified to have neutral networks. These include the NKp landscapes of Ref. [5] and the discretized NK fitness functions of Ref. [35]. The popularity of random fitness functions seems motivated by the idea that something as complicated as a biological genotype-phenotype mapping can only be statistically described using a randomized structure. Although this seems sensible in general, the results tend to be strongly dependent on the specific randomization procedure that is chosen; the results might be biologically misleading. For instance, NK models create random epistatic interactions between bits, mimicking spin-glass models in physics. In the context of spin glasses this procedure is conceptually justified by the idea that the interactions between the spins were randomly frozen in when the magnetic material formed. However, in the context of genotype-phenotype mappings, the interactions between different genes are themselves the result of evolution. This can lead to very different kinds of "random" interactions, as shown in Ref. [3].

At a minimum, though, the most striking difference between our choice of fitness function class and randomized fitness functions, is that the population dynamics of the randomized classes is very difficult, if not impossible, to analyze at present. In contrast, the population dynamics of the class of fitness functions just introduced can be analyzed in some detail. Moreover, for biological systems it could very well be that structured fitness functions, like the Terraced Labyrinth class, may contain all of the generality required to cover the phenomena claimed to be addressed by the randomized classes. Several limitations and generalizations of the Terraced Labyrinth fitness functions are discussed in Sec. 9.2.

4 A Simple Genetic Algorithm

For our analysis of epochal evolutionary dynamics we chose a simplified form of a genetic algorithm (GA) that does not include crossover and that uses fitness-proportionate selection. A population of M individuals, each specified by a genotype of length L bits reproduces in discrete non-overlapping

generations. Each generation, M individuals are selected (with replacement) from the population in proportion to their genotype's fitness. Each selected individual is placed into the population at the next generation after mutating each genotype bit with probability μ .

This GA effectively has two parameters: the mutation rate μ and the population size M. A given evolutionary optimization problem is specified, of course, by the fitness function parameters as given by the constellations, portals, and their fitness values. Stated most prosaically, then, our central goal is to analyze the population dynamics, as a function of μ and M, for any given fitness function in the Terraced Labyrinth class. Here we review the essential aspects of the population dynamics analysis.

5 Statistical Dynamics of Evolutionary Search

Refs. [45] and [46] developed an approach, which we called *statistical dynamics*, to analyze the behavioral regimes of a GA searching fitness functions that lead to epochal dynamics. Here we can only briefly review the mathematical details of this approach to evolutionary dynamics, emphasizing the motivations and the main ideas and tools from statistical mechanics and dynamical systems theory. The reader is referred to Ref. [46] for an extensive and mathematically detailed exposition. There, the reader will also find a review of the connections and similarities of our work with the alternative methodologies for GA theory developed by Vose and collaborators [36,47,48], by Prügel-Bennett, Rattray, and Shapiro [37,38,39], in the theory of molecular evolution [13,14], and in mathematical population genetics [24].

5.1 Statistical Mechanics

Our approach builds on ideas from statistical mechanics [7,40,51] and adapts its equilibrium formulation to apply to the piecewise steady-state dynamics of epochal evolution. The microscopic state of systems that are typically studied in statistical mechanics—such as, a box of gas molecules—is described in terms of the positions and momenta of all particles. What is of physical interest, however, are observable (and reproducible) quantities, such as, the gas's pressure P, temperature T, and volume V. The goal is to predict the relationships among these macroscopic variables, starting from knowledge of the equations of motion governing the particles and the space of the entire system's possible microscopic states. A given setting of macroscopic variables—e.g. a fixed P, V, and T—is often referred to as a macrostate; whereas a snapshot of the positions and momenta of all particles is called a microstate.

There are two kinds of assumptions that allow one to connect the microscopic description (collection of microstates and equations of motion) to observed macroscopic behavior. The first is the assumption of *maximum entropy* which states that all microscopic variables, unconstrained by a given macrostate, are as random as possible.

The second is the assumption of self-averaging. In the thermodynamic limit of an infinite number of particles, self-averaging says that the macroscopic variables are expressible only in terms of themselves. In other words, the macroscopic description does not require knowledge of detailed statistics of the microscopic variables. For example, at equilibrium the macroscopic variables of an ideal gas of noninteracting particles are related by the equation of state, PV = kNT, where k is a physical constant, and N is the total number of particles in the box. Knowing, for instance, the frequency with which molecules come within 100 nanometers of each other does not improve this macroscopic description.

Varying an experimental control parameter of a thermodynamic system can lead to a sudden change in its structure and in its macroscopic properties. This occurs, for example, as one lowers the temperature of liquid water below the freezing point. The liquid macrostate undergoes a phase transition and the water turns to solid ice. The macrostates (phases) on either side of the transition are distinguished by different sets of macroscopic variables. That is, the set of macrovariables that is needed to describe ice is not the same as the set of macrovariables that is needed to describe water. The difference between liquid water and solid ice is captured by a sudden reduction in the freedom of water molecules to move. While the water molecules move equally in all directions, the frozen molecules in the ice-crystal possess a relatively definite spatial location. Passing through a phase transition can be thought of as creating, or destroying, macroscopic variables and making or breaking the symmetries associated with them. In the liquid to solid transition, the rotational symmetry of the liquid phase is broken by the onset of the rigid lattice symmetry of the solid phase. As another example, in the Curie transition of a ferromagnet, the magnetization is the new macroscopic variable that is created with the onset of magnetic-spin alignment as the temperature is lowered.

5.2 Evolutionary Statistical Mechanics

The statistical mechanical description can also be applied to evolutionary processes. From a microscopic point of view, the exact state of an evolving population is only fully described when a list \mathcal{S} of all genotypes with their frequencies of occurrence in the population is given. On the microscopic level, the evolutionary dynamics is implemented as a Markov chain with the conditional transition probabilities $\Pr(\mathcal{S}'|\mathcal{S})$ that the population at the next generation will be the "microscopic" collection \mathcal{S}' ; see Refs. [17] and [36] for the microscopic formulation in the context of mathematical population genetics and genetic algorithms, respectively. For any reasonable genetic representation, however, there is an enormous number of these microscopic states \mathcal{S}

and so too of their transition probabilities. The large number of parameters, $\mathcal{O}(2^L!)$, makes it almost impossible to quantitatively study the dynamics at this microscopic level.

More practically, a full description of the dynamics on the level of microscopic states \mathcal{S} is neither useful nor typically of interest. One is much more likely to be concerned with relatively coarse statistics of the dynamics, such as the evolution of the best and average fitness in the population or the waiting times for evolution to produce a genotype of a certain quality. The result is that quantitative mathematical analysis faces the task of finding a macroscopic description of the microscopic evolutionary dynamics that is simple enough to be tractable numerically or analytically and that, moreover, facilitates predicting the quantities of interest to an experimentalist.

With these issues in mind, we specify the macrostate of the population at each time t by some relatively small set of macroscopic variables $\{\mathcal{X}(t)\}$. Since this set of variables intentionally ignores vast amounts of detail in the microscopic variables $\{\mathbf{x}(t)\}$, it is generally impossible to exactly describe the evolutionary dynamics in terms of these macroscopic variables. To achieve the benefits of a coarser description, we assume that the population has equal probabilities to be in any of the microscopic states consistent with a given macroscopic state. That is, we assume maximum entropy over all microstates $\{\mathbf{x}(t)\}$ that are consistent with the specific macrostate $\{\mathcal{X}(t)\}$.

Additionally, in the limit of infinite-population size, we assume that the resulting equations of motion for the macroscopic variables become closed. That is, for infinite populations, we assume that we can predict the state of the macroscopic variables at the next generation, given the present state of only the macroscopic variables. This infinite population limit is analogous to the thermodynamic limit in statistical mechanics. The corresponding assumption is analogous to self-averaging of the macroscopic evolutionary dynamics in this limit.

We use the knowledge of the microscopic dynamics together with the maximum entropy assumption to predict the next macrostate $\{\mathcal{X}(t+1)\}$ from the current one $\{\mathcal{X}(t)\}$. Then we re-assume maximum entropy over the microstates $\{x(t+1)\}$ given the new macrostate $\{\mathcal{X}(t+1)\}$. Since this method allows one to relax the usual equilibrium constraints and so account for the dynamical change in macroscopic variables, we refer to this extension of statistical mechanics as *statistical dynamics*. A similar approach has been developed in some generality for non-equilibrium statistical mechanics by Streater and, not surprisingly, it goes under the same name [42].

5.3 Evolutionary Macrostates

The key, and as yet unspecified step, in developing such a statistical dynamics framework of evolutionary processes is to find an appropriate set of macroscopic variables that satisfy the above assumptions of maximum entropy and self-averaging. In practice, this is difficult. Ultimately, the suitability of a set

of macroscopic variables has to be verified by comparing theoretical predictions with experimental measurements. In choosing such a set of macroscopic variables one is guided by knowledge of the fitness function and the genetic operators. Although not reduced to a procedure, this choice is not made in the dark.

First, there might be symmetries in the microscopic dynamics. Imagine, for instance, that genotypes can have only two possible values for fitness, f_A and f_B . Assume also that under mutation all genotypes of type A are equally likely to turn into type-B genotypes and that all genotypes of type B have equal probability to turn into genotypes of type A. In this situation, it is easy to see that we can take the macroscopic variables to be the relative proportions of A genotypes and B genotypes in the population. The reason one can do this is that all microstates with a certain proportion of A and B types give rise to exactly the same dynamics on the level of proportions of A and B types. That is, the dynamics is symmetric under any transformation of the microstates that leaves the proportions of A and B types unaltered. Neither selection nor mutation distinguish different genotypes within the sets A and B on the level of the proportions of A's and B's that they produce in the next generation. Obviously, one wants to take advantage of such symmetries in a macroscopic description. However, for realistic cases, such symmetries are not often abundant. Simply taking them into account, while important, does not typically reduce the complexity of the description sufficiently.

One tends to make more elaborate assumptions in developing a macroscopic description. Assume that the A and B genotypes are not all equally likely to turn from type A to B and vice versa, but do so only on average. For example, it might be the case that not all A types behave exactly the same under mutation, but that the dominant subset of A's that occurs in a population typically behaves like the average over the set of all A types. This is a much weaker symmetry than the exact one mentioned above. Importantly, it still leads to an accurate description of the dynamics on the level of A and B types under the maximum entropy assumption.

The Neo-Darwinian formalism of biological evolution suggests a natural decomposition of the microscopic population dynamics into a part that is guided by selection and a part that is driven by genetic diversification. Simply stated, selection is an ordering force induced by the environment that operates on the level of the phenotypic fitness in a population. In contrast, genetic diversification is a disordering and randomizing force that drives a population to an increased diversity of genotypes. Thus, it seems natural to choose as macrostates the proportion of genotypes in the different fitness classes (subbasins) and to assume that, due to random genetic diversification within each subbasin, genetic variation can be approximated by the maximum entropy distribution within each subbasin. This intuition is exactly the one we use in our statistical dynamics analysis of the Terraced Labyrinth fitness functions. Specifically, we describe the population in terms of the proportions

 $P_{\bar{i}}$ that are located in each of the subbasins $B_{\bar{i}}$. The maximum entropy assumption entails that within subbasin $B_{\bar{i}}$, individuals are equally likely to be any of the genotypes in $B_{\bar{i}}$. In other words, we assume that all free bits in a constellation are equally likely to be in any of their nonportal configurations.

The essence of our statistical dynamics approach is to describe the population state at any time during a GA run by a relatively small number of macroscopic variables—variables that (i) in the limit of infinite populations self-consistently describe the dynamics at their own level and (ii) can change over time. After obtaining the dynamics in the limit of infinite populations explicitly, one then uses this knowledge to solve for the GA's dynamical behaviors with finite populations.

6 Evolutionary Dynamical Systems

Up to this point we have described our approach in terms of its similarities with statistical mechanics. We appealed intuitively to macroscopic "dynamics", which can be derived in terms of the microscopic equations of motion (of selection and mutation on genotypes) and the maximum entropy assumption. Now we fill in the other half of the story, the half that clarifies what "dynamics" is and that draws out the similarities of our approach with dynamical systems theory.

As we just explained, we approximate the complete finite-population dynamics in two steps. First, we use the maximum entropy assumption together with the microscopic equations of motion to construct an infinite-population "flow" that describes the deterministic (macroscopic) dynamics of the subbasin distribution of an infinite population. Then, we construct the finite-population dynamics by accounting for the finite-population sampling at each generation. The net result is a stochastic nonlinear dynamical system. We now explain these two steps in more detail.

6.1 Infinite Populations

Consider an infinite population with subbasin distribution \vec{P} , where component $P_{\vec{i}} \in [0, 1]$ is the proportional of individuals in the subbasin $B_{\vec{i}}$. Note that the number of components in \vec{P} is equal to the number of nodes in the constellation tree that describes the Terraced Labyrinth fitness function. Given this, the question is how selection and mutation, acting on the distribution $\vec{P}(t)$, create the distribution $\vec{P}(t+1)$ at the next generation.

The effects of selection are simple, since all genotypes in subbasin $B_{\bar{i}}$ have the same fitness. If $\langle f \rangle$ is the average fitness in the population, we simply have that after selection the components are $P_{\bar{i}}^{select} = f_{\bar{i}}P_{\bar{i}}(t)/\langle f \rangle$. To calculate the effects of mutation we have to use our maximum entropy assumption. The probability that a genotype in subbasin $B_{\bar{i}}$ turns into a genotype in subbasin $B_{\bar{i}}$ is simply given by the average probability of a mutation from a genotype

in $B_{\vec{j}}$ to any genotype in $B_{\vec{i}}$. The average is taken with equal weights over all genotypes in $B_{\vec{j}}$. Putting the effects of selection and mutation together, we obtain a generation operator G that specifies the macroscopic evolutionary dynamical system:

$$\vec{P}(t+1) = \mathbf{G}[\vec{P}(t)] . \tag{1}$$

The infinite population dynamics on the level of subbasin distributions is simply given by iterating the operator \mathbf{G} . Following the terminology introduced in molecular evolution theory we call $\vec{P}(t)$ the *phenotypic quasispecies*.

The expected change $\langle d\vec{P} \rangle$ in the fitness distribution over one generation is given by:

$$\langle d\vec{P}\rangle = \mathbf{G}[\vec{P}] - \vec{P}.\tag{2}$$

We visualize the flow induced by the macroscopic equations of motion by plotting $\langle d\vec{P} \rangle$ at a number of states in the simplex of populations. This is shown in Fig. 4; after Ref. [46]. The fitness function and evolution parameters of Fig. 4 are those of Fig. 2. The temporal behavior of the system, starting in an initial condition $\vec{P}(t=0)$, is simply given by following the flow arrows.

For large $(M > 2^L)$ populations the dynamics of the subbasin distribution is simple: $\langle f \rangle$ increases smoothly and monotonically to an asymptote over a small number of generations. (See Fig. 3 of Ref. [45].) That is, there are no epochs. The reason for this is simple: for an infinite population, all genotypes, and therefore all subbasins, are represented in the initial population. Instead of the evolutionary dynamics discovering fitter genotypes over time, it essentially only expands the proportion of globally optimal genotypes already present in the initial population at t = 0.

6.2 Finite Populations

In spite of the qualitatively different dynamics for infinite and finite populations, we showed in Ref. [46] that the (infinite population) operator \mathbf{G} is the essential ingredient for describing the finite-population dynamics with its epochal dynamics as well. Beyond the differences in observed behavior, there are two important mathematical differences between the infinite-population dynamics and that with finite populations. The first is that with finite populations the components $P_{\vec{i}}$ cannot take on continuous values between 0 and 1. Since the number of individuals in subbasin $B_{\vec{i}}$ is necessarily an integer, the values of $P_{\vec{i}}$ are quantized in multiples of 1/M. Thus, the continuous simplex of allowed infinite-population fitness distributions turns into a regular, discrete lattice with spacing of 1/M. Second, due to finite-population sampling fluctuations, the dynamics of the subbasin distribution is no longer deterministic, as described by Eq. (1). In general, we can only determine the conditional probabilities $\Pr[\vec{Q}|\vec{P}]$ that a given fitness distribution \vec{P} leads to another \vec{Q} in the next generation.

¹ It will become clear shortly why we call this change an *expected* change.

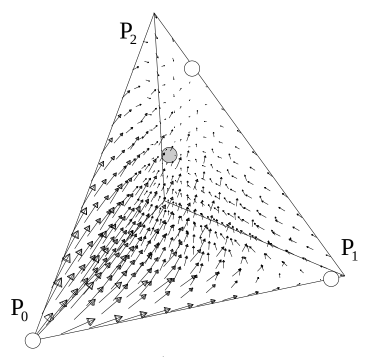


Fig. 4. Fitness distribution flow $\langle d\vec{P} \rangle$ in the simplex for the Royal Road fitness function with N=3 constellations with K=10 bits each and for the simple GA with mutation rate $\mu=0.005$; cf. Fig. 2. Fixed points of the flow are shown as large balls. The grey ball is the stable, asymptotic fixed point inside the simplex. The white balls indicate the locations of the unstable fixed points that are outside the simplex. The latter do not represent valid populations, but nonetheless they can affect the dynamics of allowed populations within the simplex by slowing down (short arrows) the flow near them.

The net result is that the probabilities $\Pr[\vec{Q}|\vec{P}]$ are determined by a multinomial distribution with mean $\mathbf{G}[\vec{P}]$:

$$\Pr[\vec{Q}|\vec{P}] = M! \prod_{\vec{r}} \frac{\left(\mathbf{G}_{\vec{i}}[\vec{P}]\right)^{m_{\vec{i}}}}{m_{\vec{i}}!} . \tag{3}$$

where $Q_{\vec{i}} = m_{\vec{i}}/M$, with $0 \le m_{\vec{i}} \le M$ integers and the product runs over all subbasins \vec{i} . (The stochastic effects of finite-population sampling are illustrated in Fig. 5.) For any finite-population subbasin distribution \vec{P} the operator \mathbf{G} gives the evolution's average dynamics over one time step, since by Eq. (3) the expected subbasin distribution at the next time step is $\mathbf{G}[\vec{P}]$. Note that the components $\mathbf{G}_{\vec{i}}[\vec{P}]$ need not be multiples of 1/M. Therefore, the actual subbasin distribution \vec{Q} at the next time step is not $\mathbf{G}[\vec{P}]$, but is instead one of the allowed lattice points in the finite-population state space

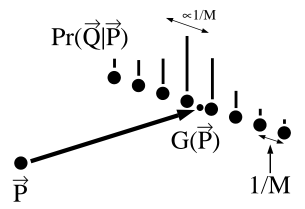


Fig. 5. Illustration of the stochastic dynamics that maps from one generation to the next. Starting with finite population \vec{P} , the arrow indicates the *expected* next population $\mathbf{G}[\vec{P}]$. Due to sampling, the probability that the actual next population is \vec{Q} is given by a multinomial distribution $\Pr[\vec{Q}|\vec{P}]$, Eq. (3). Note that the underlying state space is a discrete lattice with spacing 1/M.

consistent with the distribution $\Pr[\vec{Q}|\vec{P}]$. Since the variance around the expected distribution $\mathbf{G}[\vec{P}]$ is proportional to 1/M, \vec{Q} tends to be one of the lattice points close to $\mathbf{G}[\vec{P}]$.

Putting both the infinite-population dynamical system and the stochastic sampling effects induced by finite populations together, we arrive at the our basic model of evolutionary population dynamics. We can now begin to draw out some consequences.

7 Metastability and the Unfolding of Macrostates

Assume that there are no individuals in a certain subbasin $B_{\vec{i}}$ and that the component $\langle dP_{\vec{i}} \rangle$ is much smaller than 1/M. In that case, the actual change in component $P_{\vec{i}}$ is likely to be $dP_{\vec{i}} = 0$ for a long succession of generations. That is, if there are no individuals in subbasin $B_{\vec{i}}$ and the rate of creation of such individuals is much smaller than 1/M, then subbasin $B_{\vec{i}}$ is likely to stay empty for a considerable number of generations. Consequently, there is no movement to increase fitness to level $f_{\vec{i}}$ during this time. More generally, if the size of the flow $\langle dP_{\vec{i}} \rangle$ (and its variance) in some direction \vec{i} is much smaller than the lattice spacing (1/M) of allowed finite populations, we expect the subbasin distribution to not change in direction \vec{i} . In Refs. [45] and [46] we showed this is the mechanism that causes epochal dynamics for finite populations.

More formally, an epoch corresponds to the population being restricted to a region of an *n*-dimensional subsimplex of the macroscopic state space. Stasis occurs because the flow out of this subspace is much smaller than the finite-population induced lattice spacing. In particular, for the Terraced

Labyrinth fitness functions, an epoch corresponds to the time during which the highest fitness individuals are located in subbasin $B_{i_1,i_2,...,i_n}$. During this time, an equilibrium subbasin distribution is established in the population. Its components are nonzero only for subbasins B, B_{i_1} , B_{i_1,i_2} , through $B_{i_1,...,i_n}$. That is, they are nonzero for all of the lower fitness subbasins that connect $B_{\vec{i}}$ to the root. Since the discovery of a portal configuration of constellation $c_{i_1,...,i_n}$ is rare, the population remains in this n-dimensional subsimplex for a considerable number of generations. The number of generations it remains in this epoch is, of course, directly dependent on the number of portals out of the subbasin $B_{\vec{i}}$ and the number of bits $K_{\vec{i}}$ in constellation $c_{\vec{i}}$.

Recall the example of epochal behavior of Sec. 2.2 and Fig. 2. Initially, the population was located in the zero-dimensional macrostate corresponding to all genotypes located in the root subbasin. Then the first portal configuration was discovered and the population moved onto the line of population states that have some individuals in the root subbasin and some in the basin B_1 . After this epoch, a genotype in subbasin $B_{1,1}$ was discovered and the population moved to a steady-state in the plane of proportions P, P_1 , and $P_{1,1}$. (These were labeled according to their fitnesses— P_0 , P_1 , and P_2 —in Fig. 2.) Finally, the global optimum string in subbasin $B_{1,1,1}$ was discovered, and the population moved to its final fixed point in the three-dimensional simplex.

The global evolutionary dynamics can be viewed as an incremental discovery (an unfolding) of successively more (macroscopic) dimensions of the subbasin distribution space. In most realistic settings, it is typically the case that population sizes M are much smaller than 2^L . Initially, then, the population consists only of genotypes in subbasins of low fitness. Assume, for instance, that genotypes in subbasin $B_{1,2}$ are the highest fitness ones in the initial population. Mutation and selection establish an equilibrium phenotypic quasispecies $\vec{P}^{1,2}$, consisting of nonzero proportions of genotypes in the subbasin B, B_1 , and $B_{1,2}$, and zero proportions of genotypes in all other subbasins. Individuals and their descendants drift through subbasin $B_{1,2}$. The subbasin distribution fluctuates around $\vec{P}^{1,2}$ until a portal configuration $\pi_{1,2,i}$ of the constellation $c_{1,2}$ is discovered and genotypes of (higher) fitness $f_{1,2,i}$ spread through the population. The population then settles into subbasin distribution $\vec{P}^{1,2,i}$ with average fitness $\langle f \rangle_{1,2,i}$ until a portal $\pi_{1,2,i,j}$ of constellation $c_{1,2,i}$ is discovered, and so on, until a local optimum corresponding to a leaf of the fitness function tree is found. In this way, the macroscopic dynamics can be seen as stochastically hopping between the different epoch distributions $\vec{P}^{\vec{i}}$ of subbasins $B_{\vec{i}}$ that are connected to each other in the fitness function tree.

Note that at each stage \vec{P}^{i_1,\dots,i_n} has only n+1 (nonzero) components, each corresponding to a subbasin connecting $B_{\vec{i}}$ to the tree root. All other subbasin components are zero. The selection-mutation balance maintains a constant proportion of genotypes with correct configurations in all constellations that define the epoch. By the maximum entropy assumption, the action of the

generation operator \mathbf{G} is symmetric with respect to all remaining nonportal constellation configurations. That is, \mathbf{G} 's action is indifferent to the various proportions of particular incorrect constellations configurations. The symmetry among constellation $c_{\bar{\imath}}$'s incorrect configurations is broken dynamically when a (typically, rare) portal configuration is discovered. This symmetry breaking adds a new macroscopic variable—a new "active" dimension of the phenotype. This symmetry breaking and stabilization of a new phenotypic dimension is the dynamical analogue of a phase transition.

As alluded to earlier, much of the attractiveness of the Terraced Labyrinth class of fitness functions lies in the fact that, to a good approximation, analytical predictions can be obtained for observable quantities; such as, average epoch fitness $\langle f \rangle_{\vec{i}}$ and the epoch subbasin distribution $\vec{P}^{\vec{i}}$ in terms of the evolutionary and fitness function parameters. For instance, assume that the highest fitness genotypes are in subbasin $B_{i_1,i_2,...,i_n}$ and that the population resides in the steady-state distribution $\vec{P}^{i_1,i_2,...,i_n}$. Denote by

$$L_{i_1,i_2,\dots,i_m} = K + K_{i_1} + K_{i_1,i_2} + \dots + K_{i_1,i_2,\dots,i_{m-1}}, \tag{4}$$

the number of constrained bits in each of the subbasins that have nonzero proportions during this epoch. (Note that L=0 for the root subbasin). Then, up to some approximation,² the average epoch fitness is simply given by

$$\langle f \rangle_{\vec{i}} = f_{\vec{i}} (1 - \mu)^{L_{\vec{i}}}. \tag{5}$$

One can also derive the subbasin distribution $\vec{P}^{\vec{i}}$. In order to express the results most transparently, we introduce the fitness-level ratio using Eq. (5):

$$\alpha_{\vec{i}\vec{j}} = \frac{f_{\vec{j}}}{f_{\vec{i}}} (1 - \mu)^{L_{\vec{j}} - L_{\vec{i}}} \tag{6}$$

Then we have for the highest-fitness component of the subbasin distribution $P_{\overline{i}}$ that

$$P_{\vec{i}}^{\vec{i}} = \prod_{\vec{m} < \vec{i}} \frac{1 - \alpha_{\vec{i}\vec{m}}}{1 - \alpha_{\vec{i}\vec{m}}(1 - \mu)^{K_{\vec{m}}}},\tag{7}$$

where $\vec{m} < \vec{i}$ indicates the set of all nodes lying along the path between \vec{i} and the tree's root, including the root. For the other components of $P_{\vec{i}}$ we have that

$$P_{\vec{j}}^{\vec{i}} = \frac{(1-\mu)^{L_{\vec{j}}} \left(1 - (1-\mu)^{K_{\vec{j}}}\right)}{1 - \alpha_{\vec{i}\vec{j}} (1-\mu)^{K_{\vec{j}}}} \prod_{\vec{m} < \vec{j}} \frac{1 - \alpha_{\vec{i}\vec{m}}}{1 - \alpha_{\vec{i}\vec{m}} (1-\mu)^{K_{\vec{m}}}}.$$
 (8)

Describing the dynamics in and between epoch distributions \vec{P}^i using diffusion approximations and then invoking (dynamical systems) concepts—such

² The approximation here is that, during an epoch, the *back mutations* from lower fitness subbasins to higher subbasins can be neglected. This assumption is generally valid for constellation lengths $K_{\vec{i}}$ that are not too small.

as, stable and unstable manifolds, Jacobian eigenvalues, and their eigenvectors—a number of additional properties of epochal evolution can be derived analytically and predicted quantitatively. The reader is referred to Refs. [46] and [43] for the detailed analysis of the distribution of epoch fluctuations, the stability of epochs, and the average waiting times for portal discovery.

8 Frozen Accidents, Phenotypic Structural Constraints, and the Subbasin-Portal Architecture

The subbasin-portal architecture, whose population dynamics we are analyzing, suggests a natural explanation for the occurrence and longevity of frozen accidents. Generally speaking, frozen accidents refer to persistent phenotypic characters that are selected out of a range of possible, structurally distinct alternatives by specific random events in the evolutionary past. One imagines an arbitrary event, such as a sampling fluctuation, promoting one or another phenotype, which then comes to dominate the population and thereby excludes alternatives that could be equally or even more fit in the long term.

Within the class of Terraced Labyrinth fitness functions frozen accidents occur via a simple mechanism. In particular, a given evolutionary path through the fitness-function tree can be regarded as a sequence of frozen accidents. Since different portals of the same constellation are mutually exclusive, their subbasins are separated by a fitness barrier. Across a wide range of parameter settings, the crossing of such fitness barriers takes much longer than the discovery of new portals, via neutral evolution, in the current subbasin. Once evolution has taken a certain branch up the tree, it is therefore unlikely, that it will ever return. That is, once a subbasin $B_{\vec{i}}$ is discovered, the further course of evolution is restricted to the subtree with its root at \vec{i} . In this way, the genotypic constellations up to \vec{i} become installed in the population.

The alternative evolutionary paths are not merely a case of genetic book-keeping. Different portals of a constellation $c_{\tilde{i}}$ may be associated with very different phenotypic innovations. Once a particular phenotypic innovation has occurred, the phenotype determines which range of future phenotypic innovations can occur. This contingency—how evolutionary futures depend on current phenotypic constraints—goes under the name of structural phenotypic constraints. In the Terraced Labyrinth this phenomenon is reflected in the possibility that fitness-function trees have very dissimilar subtrees. For instance the subtrees rooted at nodes 1 and 2 in Fig. 3 are very dissimilar. This dissimilarity reflects the fact that evolutionary futures starting from the phenotype corresponding to node 1 are very different from those starting from the phenotype associated with node 2.

Naturally, the Terraced Labyrinth class of fitness functions does not indicate which kind of tree structures, reflecting structural constraints, are appropriate or biologically realistic. This will ultimately be decided by ex-

periment. The generality of this class of fitness functions, however, illustrates that qualitative concepts—such as, frozen accidents and structural phenotypic constraints—are very easily represented and analyzed within the statistical dynamics framework.

9 Concluding Remarks

9.1 Summary

We introduced a generalized subbasin-portal architecture by way of defining a new class of fitness functions—the Terraced Labyrinth. The detailed mathematical analysis of the population dynamics that we introduced previously can be adapted straightforwardly to this generalized setting. In this way, statistical dynamics was shown to have a wider applicability and its results on epochal evolution are seen to have wider ranging consequences than the first analyses in Refs. [45] and [46] might have suggested.

We described this more general view of epochal evolution, attempting to clarify the connections to both statistical mechanics and dynamical systems theory. The result is a dynamical picture of a succession of "phase transitions" in which microscopic symmetries are broken and new macroscopic dimensions are discovered and then stabilized. These new macroscopic dimensions then become the substrate and historical context for further evolution.

9.2 Extensions and Generalizations

There are a number of extensions to more complex evolutionary processes that should now be possible. Here we mention a few limitations of the class of fitness functions analyzed and several generalizations.

First, constellations do not overlap constellations higher in the tree. Second, all the subbasins have a similar regular architecture: there is a set of constrained bits (in the portals) that define the subbasin and all other bits are free. This is undoubtedly not the case generally. Different subbasins can have distinct irregular architectures and different kinds of portals. Moreover, the diffusion dynamics through distinct subbasins might be different. For instance, subbasins might also be defined with respect to more complicated genetic operations—such as, gene duplication, unequal crossovers, and gene conversion.

Third, all of a subbasin's portals correspond to configurations of a single constellation. This insures that the topology of the subbasin hierarchy forms a tree, as opposed to the more general topologies suggested by Fig. 1. Extending the analysis to more complicated subbasin architectures is formally straightforward, but becomes considerably more complicated to carry out. For very complicated architectures, the approximations in our analysis may have to be reworked.

Fourth, one would like to extend statistical dynamics to open-ended models in which (say) the genotype length can grow over time, allowing the tree to dynamically grow new branches as well; perhaps along the lines investigated in Ref. [3]. One would hope to see how the evolutionary dynamics adapts as the mutation-genome length error threshold is approached [13]. As long as such open-ended models adhere to the tree topology of the subbasin-portal hierarchy, it would appear that our analyses could easily be extended to them.

Finally, the maximum entropy assumption only holds to some degree of approximation. For instance, whenever a new macrodimension unfolds, the population is initially concentrated around the portal genotype in the neutral network; this is a type of founder effect. The population then spreads out randomly from there, but the genotypes never completely decorrelate due to finite-population sampling fluctuations [12]. Moreover, as we have shown in Ref. [44], the population members in lower-fitness subbasins are closely genetically related to members in the subbasin of currently highest fitness. These facts flatly contradict the maximum entropy assumption that individuals are randomly and independently spread through the subbasins. Since these complications do not generally alter the rate of deleterious mutations from subbasins to lower-fitness subbasins, theoretical predictions—such as, the epoch distributions $\vec{P}^{\vec{i}}$ —are not much affected. However, as shown in Ref. [46], statistics—such as, the average waiting time for the discovery of a portal—may be significantly affected. This leaves open the question of how to extend the set of macroscopic variables to account for these complications.

Acknowledgments. This work was supported at the Santa Fe Institute by grants from the NSF, ONR, and Sandia National Laboratory.

References

- C. Adami. Self-organized criticality in living systems. Phys. Lett. A, 203:29–32, 1995.
- 2. L. M. Adelman. Molecular computation of solutions to combinatorial problems. *Science*, 266:1021–1024, 1994.
- L. Altenberg. Genome growth and the evolution of the genotype-phenotype map. In W. Banzhaf and F. H. Eeckman, editors, Evolution and Biocomputation. Computational Models of Evolution, Monterey, California, July 1992, pages 205–259, Berlin, 1995. Springer Verlag.
- T. Back. Evolutionary algorithms in theory and practice: Evolution strategies, evolutionary programming, genetic algorithms. Oxford University Press, New York, 1996.
- L. Barnett. Tangled webs: Evolutionary dynamics on fitness landscapes with neutrality. Master's thesis, School of Cognitive Sciences, University of East Sussex, Brighton, 1997. http://www.cogs.susx.ac.uk/lab/adapt/nnbib.html.
- R. K. Belew and L. B. Booker, editors. Proceedings of the Fourth International Conference on Genetic Algorithms. Morgan Kaufmann, San Mateo, CA, 1991.

- J. J. Binney, N. J. Dowrick, A. J. Fisher, and M. E. J. Newman. The Theory of Critical Phenomena: An Introduction to the Renormalization Group. Oxford Science Publications, 1992.
- 8. L. Chambers, editor. *Practical Handbook of Genetic Algorithms*. CRC Press, Boca Raton, 1995.
- J. Chen, E. Antipov, B. Lemieux, W. Cedeno, and D. H. Wood. DNA computing implementing genetic algorithms. In L. F. Landweber, E. Winfree, R. Lipton, and S. Freeland, editors, *Evolution as Computation*, pages 39–49, New York, 1999. Springer Verlag.
- J. P. Crutchfield and M. Mitchell. The evolution of emergent computation. Proc. Natl. Acad. Sci. U.S.A., 92:10742–10746, 1995.
- L. D. Davis, editor. The Handbook of Genetic Algorithms. Van Nostrand Reinhold, 1991.
- B. Derrida and L. Peliti. Evolution in a flat fitness landscape. Bull. Math. Bio., 53(3):355–382, 1991.
- M. Eigen. Self-organization of matter and the evolution of biological macromolecules. *Naturwissen.*, 58:465–523, 1971.
- M. Eigen, J. McCaskill, and P. Schuster. The molecular quasispecies. Adv. Chem. Phys., 75:149–263, 1989.
- 15. S. F. Elena, V. S. Cooper, and R. E. Lenski. Punctuated evolution caused by selection of rare beneficial mutations. *Science*, 272:1802–1804, 1996.
- L. Eshelman, editor. Proceedings of the Sixth International Conference on Genetic Algorithms. Morgan Kaufmann, San Mateo, CA, 1995.
- 17. W. J. Ewens. Mathematical Population Genetics, volume 9 of Biomathematics. Springer-Verlag, 1979.
- W. Fontana and P. Schuster. Continuity in evolution: On the nature of transitions. Science, 280:1451–5, 1998.
- W. Fontana, P. F. Stadler, E. G. Bornberg-Bauer, T. Griesmacher, I. L. Hofacker, M. Tacker, P. Tarazona, E. D. Weinberger, and P. Schuster. RNA folding and combinatory landscapes. *Phys. Rev. E*, 47:2083–2099, 1992.
- S. Forrest, editor. Proceedings of the Fifth International Conference on Genetic Algorithms. Morgan Kaufmann, San Mateo, CA, 1993.
- 21. C. V. Forst, C. Reidys, and J. Weber. Evolutionary dynamics and optimizations: Neutral networks as model landscapes for RNA secondary-structure folding landscape. In F. Moran, A. Moreno, J. Merelo, and P. Chacon, editors, Advances in Artificial Life, volume 929 of Lecture Notes in Artificial Intelligence. Springer, 1995. SFI preprint 95-20-094.
- D. E. Goldberg. Genetic Algorithms in Search, Optimization, and Machine Learning. Addison-Wesley, Reading, MA, 1989.
- 23. S. J. Gould and N. Eldredge. Punctuated equilibria: The tempo and mode of evolution reconsidered. *Paleobiology*, 3:115–251, 1977.
- 24. D. L. Hartl and A. G. Clark. *Principles of population genetics*. Sinauer Associates, second edition, 1989.
- 25. R. Haygood. The structure of Royal Road fitness epochs. *Evolutionary Computation*, submitted, 1997. ftp://ftp.itd.ucdavis.edu/pub/people/rch/ StrucRoyRdFitEp.ps.gz.
- M. Huynen. Exploring phenotype space through neutral evolution. J. of Mol. Evol., 43:165–169, 1996.

- M. Huynen, P. F. Stadler, and W. Fontana. Smoothness within ruggedness: The role of neutrality in adaptation. *Proc. Natl. Acad. Sci. USA*, 93:397–401, 1996.
- S. A. Kauffman and S. Levin. Towards a general theory of adaptive walks in rugged fitness landscapes. J. Theo. Bio., 128:11-45, 1987.
- M. Kimura. The Neutral Theory of Molecular Evolution. Cambridge University Press, 1983.
- J. R. Koza. Genetic Programming: On the Programming of Computers by Means of Natural Selection. MIT Press, Cambridge, MA, 1992.
- L. F. Landweber and L. Kari. Universal molcular computation in ciliates. In L. F. Landweber, E. Winfree, R. Lipton, and S. Freeland, editors, *Evolution as Computation*, pages 51–66, New York, 1999. Springer Verlag.
- C. A. Macken and A. S. Perelson. Protein evolution in rugged fitness landscapes. Proc. Nat. Acad. Sci. USA, 86:6191–6195, 1989.
- M. Mitchell. An Introduction to Genetic Algorithms. MIT Press, Cambridge, MA, 1996.
- M. Mitchell, J. P. Crutchfield, and P. T. Hraber. Evolving cellular automata to perform computations: Mechanisms and impediments. *Physica D*, 75:361–391, 1994.
- M. Newman and R. Engelhardt. Effect of neutral selection on the evolution of molecular species. Proc. R. Soc. London B., 256:1333–1338, 1998.
- A. E. Nix and M. D. Vose. Modeling genetic algorithms with Markov chains. Ann. Math. Art. Intel., 5, 1991.
- A. Prügel-Bennett. Modelling evolving populations. J. Theo. Bio., 185:81–95, 1997.
- 38. A. Prügel-Bennett and J. L. Shapiro. Analysis of genetic algorithms using statistical mechanics. *Phys. Rev. Lett.*, 72(9):1305–1309, 1994.
- M. Rattray and J. L. Shapiro. The dynamics of a genetic algorithm for a simple learning problem. J. of Phys. A, 29(23):7451–7473, 1996.
- L. E. Reichl. A Modern Course in Statistical Physics. University of Texas, Austin, 1980.
- C. M. Reidys, C. V. Forst, and P. K. Schuster. Replication and mutation on neutral networks of RNA secondary structures. *Bull. Math. Biol.*, submitted, 1998. SFI Working Paper 98-04-036.
- 42. R. F Streater. Statistical Dynamics: A Stochastic Approach to Nonequilibrium Thermodynamics. Imperial College Press, London, 1995.
- 43. E. van Nimwegen and J. P. Crutchfield. Optimizing epochal evolutionary search: Population-size dependent theory. *Machine Learning*, submitted, 1998. Santa Fe Institute Working Paper 98-10-090. adap-org/9810004.
- 44. E. van Nimwegen and J. P. Crutchfield. Optimizing epochal evolutionary search: Population-size independent theory. Computer Methods in Applied Mechanics and Engineering, to appear, 1998. Special issue on Evolutionary and Genetic Algorithms in Computational Mechanics and Engineering, D. Goldberg and K. Deb, editors. Santa Fe Institute Working Paper 98-06-046; adaporg/9810003.
- E. van Nimwegen, J. P. Crutchfield, and M. Mitchell. Finite populations induce metastability in evolutionary search. *Phys. Lett. A*, 229:144–150, 1997.
- 46. E. van Nimwegen, J. P. Crutchfield, and M. Mitchell. Statistical dynamics of the Royal Road genetic algorithm. *Theoretical Computer Science*, in press,

- 1998. Special issue on Evolutionary Computation, A. Eiben and G. Rudolph, editors. SFI working paper 97-04-35.
- 47. M. D. Vose. Modeling simple genetic algorithms. In L. D. Whitley, editor, Foundations of Genetic Algorithms 2, San Mateo, CA, 1993. Morgan Kauffman.
- 48. M. D. Vose and G. E. Liepins. Punctuated equilibria in genetic search. *Complex Systems*, 5:31–44, 1991.
- 49. J. Weber. Dynamics of Neutral Evolution. A case study on RNA secondary structures. PhD thesis, Biologisch-Pharmazeutischen Fakultät der Friedrich Schiller-Universität Jena, 1996. http://www.tbi.univie.ac.at/papers/PhD_theses.html.
- S. Wright. Character change, speciation, and the higher taxa. Evolution, 36:427–43, 1982.
- J. M. Yeomans. Statistical Mechanics of Phase Transitions. Clarendon Press, Oxford, 1992.