Casting polymer nets to optimize noisy molecular codes

Tsvi Tlusty*

Department of Physics of Complex Systems, Weizmann Institute of Science, Rehovot 76100, Israel

Edited by Albert Libchaber, The Rockefeller University, New York, NY, and approved April 1, 2008 (received for review October 29, 2007)

Life relies on the efficient performance of molecular codes, which relate symbols and meanings via error-prone molecular recognition. We describe how optimizing a code to withstand the impact of molecular recognition noise may be understood from the statistics of a two-dimensional network made of polymers. The noisy code is defined by partitioning the space of symbols into regions according to their meanings. The “polymers” are the boundaries between these regions, and their statistics define the cost and the quality of the noisy code. When the parameters that control the cost–quality balance are varied, the polymer network undergoes a transition, where the number of encoded meanings rises discontinuously. Effects of population dynamics on the evolution of molecular codes are discussed.

In the living cell, information is carried by molecules. The outside environment and the biochemical circuitry of the cell churn out fluxes of molecular information that are read, processed, and then stored in memory by other molecules. The cell’s information-processing networks often need to translate a symbol written in one class of molecules into another symbol written in a different molecular language. This requires a code-table that translates between the two molecular languages. Perhaps the best-known example is the genetic code-table that translates 64 DNA base triplets into 20 amino acids. The notion of mapping between two molecular spaces occurs also in biological codes of a much larger scale; for example, the transcriptional regulatory network that controls gene expression by DNA-binding proteins. This network may be seen as a mapping—a probabilistic one because of the inherent noise—between the space of molecular symbols, e.g., the base triplets, and the space of molecular meanings, e.g., amino acids. The notion of mapping between two molecular spaces occurs also in biological codes of a much larger scale; for example, the transcriptional regulatory network that controls gene expression by DNA-binding proteins. This network may be seen as a mapping—a probabilistic one because of the inherent noise—between the space of molecular symbols, e.g., the base triplets, and the space of molecular meanings, e.g., amino acids. The notion of mapping between two molecular spaces occurs also in biological codes of a much larger scale; for example, the transcriptional regulatory network that controls gene expression by DNA-binding proteins. This network may be seen as a mapping—a probabilistic one because of the inherent noise—between the space of molecular symbols, e.g., the base triplets, and the space of molecular meanings, e.g., amino acids. The notion of mapping between two molecular spaces occurs also in biological codes of a much larger scale; for example, the transcriptional regulatory network that controls gene expression by DNA-binding proteins. This network may be seen as a mapping—a probabilistic one because of the inherent noise—between the space of molecular symbols, e.g., the base triplets, and the space of molecular meanings, e.g., amino acids. The notion of mapping between two molecular spaces occurs also in biological codes of a much larger scale; for example, the transcriptional regulatory network that controls gene expression by DNA-binding proteins. This network may be seen as a mapping—a probabilistic one because of the inherent noise—between the space of molecular symbols, e.g., the base triplets, and the space of molecular meanings, e.g., amino acids.

The symbol space is a graph, hexagonal in this example, in which vertices are symbols and edges connect symbols that are likely to be confused by reading. The code-table induces a coloring pattern on the symbol space, which divides it into meaning islands (dotted lines). In the triangular dual graph (Right), the boundaries between meaning islands form a network of “polymers” (thick solid lines).

random walks (Fig. 1). Polymer networks are natural in this context because they are related to the notion of space partitioning that is central to coding theory (5). But the resemblance to a polymer network is not merely structural. Optimizing the fitness of the mapping is shown to be equivalent to minimizing the free energy of the polymer network. Such an optimal mapping must balance the three conflicting needs for maximal error tolerance, for maximal diversity, and for minimal cost. During evolution, the code-table adapts by altering the network in response to changes in the equipoise of these three evolutionary forces.

In the present work, we first discuss how the partition of the symbol space by the polymer networks determines the fitness of the code-table, where mathematical definitions of fitness and its determinants, error-load, diversity, and cost are given. Next, we discuss one purpose of the present work, which is to show that the fitness function of the code-table corresponds to the free energy of the polymer network. Thus, it suggests that the problem of optimizing the code-table is equivalent to calculating the equilibrium statistics of the polymer network. A second purpose is then to use this equivalence to examine the code optimization problem in parameter regimes that are hard to access otherwise. In particular, it is used to identify a first-order transition, in which the number of meaning islands changes abruptly in response to varying the error-tolerance–diversity–cost balance. Finally, to put the model within a context of population dynamics, we discuss possible aspects of metastabili-
ity, mutations, and genetic drift that may affect the evolution of the code.

**Model and Results**

**Fitness of Molecular Codes.** Information in the cell is recorded in molecules and then retrieved and translated into other molecules by a code-table. To discuss how the organism optimizes such a code-table, we represent the table as an information channel or a mapping that relates a meaning space, in which \( m \) meanings reside, to a symbol space, which contains \( n \) symbols (Fig. 1). The code-table maps (or encodes) meanings to symbols and thus partitions the space of symbols into meaning islands whose boundaries form a polymer network. In this section, we first calculate the fitness of a code-table as a function of the mapping as specified by the polymer network. The code fitness is composed of contributions due to the diversity of the code-table, its average error load, and the cost of constructing the molecular coding apparatus. There are two sources of noise in our simple description of the coding system, noise while reading and noise in the mapping itself. Recognition errors while reading a symbol are represented by the symbol graph whose edges connect symbols that misreading may confuse. The impact of this noise is included in the error-load component of the fitness. Noise in the mapping, which is implemented in error-prone molecular recognizers, is represented as a statistical average over an ensemble of polymer networks. This second recognition problem requires additional resources and is included in the cost (which together with the error load is defined below).

A code-table does not necessarily map all of the \( m \) available meanings to symbols but may possibly map only \( n_i \leq m \) out of them. The larger is the number of encoded meanings \( n_i \), the more diverse is the code. Diversity contributes to the fitness of the code because it increases the chance that when the organism needs to read, write, or process a certain meaning it can accurately encode it as one of the available symbols. However, diversity of meanings also increases the probability for misreading errors: To reconstruct the meaning encoded by a certain symbol, the organism has to read it. Because the molecular reading apparatus is not perfect, it may sometimes confuse this symbol with one of its neighbors in the graph (Fig. 1). If many meanings are encoded then, on average, the meaning islands defined by the network are smaller. In this finer network, the chance of confusing symbols of different meanings is larger, which costs the organism in a higher error load.

To find the error load, one needs to specify a partition into meaning islands and examine the average chance to cross by misreading the boundaries between the islands. We specify a partition (see Fig. 1) by assigning to each edge \( i-j \) a binary variable \( E_{i,j} \) that indicates whether the edge is on the boundary between two meaning islands \( (E_{i,j} = 1) \) or inside of an island \( (E_{i,j} = 0) \). Misreading along the edge, which confuses \( i \) with \( j \), occurs with a probability \( r_{i,j} \) while \( r_{i,j} \) is the probability to correctly read \( i \). There may be two possible outcomes of misreading: If both symbols are in the same island and are therefore synonymous, then misreading bears no load because the translated meaning does not change. If the symbols reside in two different islands and are therefore nonsynonymous, then the fitness of the organism decreases by one fitness unit. The contribution of an \( i-j \) misreading to the error load can be therefore quantified as \( r_{i,j} E_{i,j} \) and the total error load is a sum over all edges, \( \sum_{i,j} r_{i,j} E_{i,j} \).

The need for diversity is an evolutionary force that counteracts the need to minimize error load. Our model assumes for simplicity that the contribution of diversity to the fitness is linear in \( n_i \), the number of encoded meanings. The quality \( H_{E,C} \) of a given network pattern \( E \), is then a linear combination of the error load and the diversity, where the parameter \( w_D \) measures the significance of diversity relative to error load. We use a sign convention in which an optimal code of high quality corresponds to low values of \( H_{E,C} \).

The quality depends on the coloring pattern of the code-table, which determines its error load and diversity. As illustrated in Fig. 1, each coloring pattern is determined by the network of boundaries between the islands, which is equivalent to a network of polymers. The quality is governed by the interplay between error load and diversity: If the reader were ideal \( r_{i,j} = \delta_{i,j} \), then it would have been advantageous to decode as many meanings as there are available symbols, \( n_i = n \). However, because the molecular reader is not perfect, it is preferable to decode fewer meanings to minimize the effect of misreading errors. The quality \( H_{E,C} \) corresponds to a “macrostate” specified by a deterministic network configuration \( E \). The stochastic mapping of the molecular code is a “microstate,” which is represented by an ensemble of such configurations and is calculated below. The code quality \( H_{E,C} \) is defined as the ensemble average of quality over all microstates, \( H_{C} = \langle H_{E,C} \rangle \).

Besides the quality of the code, which combines its error load and diversity, the code fitness must also account for the cost of the molecular machinery that performs the mapping. Molecular codes are physically implemented by recognition interactions between the meanings, the symbols, and sometimes other intermediary molecules, such as the tRNA in the genetic code (1). High specificity of recognition improves the quality of the code because it enables more accurate mapping. However, highly specific binding requires a higher binding energy, which in general necessitates larger binding sites. It is plausible to assume that the cost of the code is proportional to the average size of the binding sites and therefore to the average binding energy (6). This is because the cost of synthesizing the molecules and maintaining their genes is proportional to their size. To estimate the cost, one notes that the binding probability scales like the Boltzmann exponent of the binding energy (in units of \( k_B T \)), \( P_{b} \sim \exp(E_{b}) \). It follows that the cost, which is equal to the average binding energy \( E_{C} \), can be approximated by the average \( \langle \ln P_{b} \rangle \), which is minus the entropy of the mapping of meanings to symbols, \( -S_{C} \) (see Methods). This entropy averages over the ensemble of all possible mappings, which is determined by all possible networks and the number of possible ways to color every such network.

Finally, the overall fitness of the code \( F_{C} \) is estimated by a weighted sum of the quality and the cost, \( F_{C} = H_{C} - w_{C} \times S_{C} \), where the parameter \( w_{C} \) measures the significance of the cost with respect to quality. \( F_{C} \) is like a free energy of all possible colorings of the code-table and may be derived from the partition function \( Z_{C} \).

\[
H_{E,C} = \sum_{i,j} r_{i,j} E_{i,j} - w_{D} \times H_{C} \tag{1}
\]

\[
Z_{C} = \exp(-F_{C}/w_{C}) = \sum_{mappings} \exp(-H_{E,C}/w_{C}) \tag{2}
\]

Within this analogy, the quality \( H_{E,C} \) plays the role of energy, and \( w_{C} \) is an effective temperature. The ensemble average in \( Z_{C} \) is due to the probabilistic nature of the molecular mapping. At high \( w_{C} \), codes are fuzzy and smeared over many network configurations, whereas at low \( w_{C} \), they are sharper because the mapping is almost deterministic. In principle, one can derive the code-table by performing the summation in Eq. 2 (6), but in practice this is a burdensome task that can be performed only numerically and even this only for codes of limited size. Tractable analytic results exist mostly at the limit of high \( w_{C} \). In this regime, the code table undergoes a second-order phase transition from a noncoding state of no correlation between meanings and sym-
symbols to a coding state, in which such correlation has just emerged (2, 6–8).

The cost–quality balance of the code-table is analogous to the balance in an engineered noisy information channel between the average distortion in the channel $H$, which measures its quality, and the channel’s rate $I$, which measures the cost of the channel by estimating how many bits are required to encode one meaning. Rate-distortion theory (9) focuses on the fundamental problem of optimizing a noisy information channel, which can be formalized as the following question: What is the minimal rate $I$ required to assure that the distortion in the channel will be less than a certain desired value $H'$? This optimal rate-distortion curve is calculated by minimizing a functional $F = H + wCI$, where $H, I$, and $F$ are analogues of $H_E, -SC$, and $FC$, respectively. The “temperature” $wC = -\delta H C/\delta C$ is the slope of the optimal curve, measures the increase in quality due to an additional bit of information. In the biological context, $wC$ is expected to decrease with the complexity the organism and its environment: A complex organism transmits more information. It is therefore in the interest of this organism to pay a larger cost to improve the quality of its codes, and $wC = \delta H C/\delta C$ is lower. Similarly, a richer environment is also “colder.” At low $wC$, the quality $H C$ dominates the free energy $FC$ and the code-table tends to one of the many minima of $H_E$. Derivation of the optimal code in this regime is difficult, even numerically, because of the rugged landscape of $H_E$. As we discuss below, the polymer network formalism offers insight into this regime and, specifically, suggests a first-order coding transition.

Statistics of Polymer Networks on the Dual Symbol Graph. To formalize the analogy of molecular codes to polymer networks, we need to examine the dual of the symbol graph on which the network resides. To find the dual graph, one embeds the symbol graph in a surface (10). In the example of Fig. 1, the symbol graph is a hexagonal lattice that is embedded in a torus and the dual is a triangular lattice (see Methods). It is evident that the vertex-coloring pattern of the symbol graph corresponds to a connected “polymer” network whose monomers are a subset of the edges of the dual graph (Fig. 1).

By counting the number of edges and vertices in the polymer network one can derive the number of meaning islands $n_i$ in the quality $H_C$ (Eq. 1). For this purpose, we introduce another binary variable, $V_i$, that indicates whether a vertex $i$ of the dual graph is part of the polymer network ($V_i = 1$) or not ($V_i = 0$). Then, the numbers of occupied edges $n_v$, occupied vertices $n_v$, and islands $n_i$ are related through the definition of Euler’s characteristic $\chi = n_v - n_e + n_i$ (10).

$\chi$ is determined by the topology of the surface in which the symbol graph is embedded; for example, $\chi = 0$ for the torus in Fig. 1. By substitution of Eq. 3 into Eq. 1, we find that the code quality is $H_E = \Sigma E_{ij} = 1 + wD E_{ij} + wD \Sigma V_i$ and the code’s partition function is therefore $Z_C = \Sigma E_{ij} N_{EV} \exp(-H_E)$, where the summation is over all valid network configurations. The factor $N_{EV}$ is the number of possible ways to color a given pattern specified by the fields $E$ and $V$. As is common in biological codes, it is assumed henceforth that there are many more meanings than available symbols, $n_m \gg n_s \gg n_t$, so the combinatorial factor can be estimated as $N_{EV} \approx \exp(n_t n_m)$. By substituting into $Z_C$ the approximated $N_{EV}$ with the island number $n_i$ taken from Eq. 3, the partition function becomes

$$Z_C = \exp(-\alpha \chi/wC) \sum_{E, V} \left( \prod_{i-j} \exp(-\beta_i E_{ij}/wC) \prod_{i} \exp(-\alpha V_i/wC) \right),$$

with the coefficients $\alpha = w_D + w_C n_m$ and $\beta_i = (r_i - w_D) - w_C n_i$.

The code partition function $Z_C$ (Eq. 4) sums over all possible networks. The building blocks of these networks are self-avoiding polymers that fuse to each other at junctions. Within the analogy to physical networks, $E_{ij}$ may appear as a summation over two chemical “species”—one that resides on the edges with “excitation energies” (or chemical potentials) $\beta_i$, and one on the vertices with the excitation energy $\alpha$. However, these two species are not really independent and cannot be summed separately. A vertex is occupied if and only if its coordination number is at least two because “dangling ends” or isolated vertices are forbidden. Similarly, an edge is occupied if and only if it connects two occupied vertices. The relevant chemical species are the monomers, i.e., pairs of a vertex and a neighboring edge that carry energies of $\alpha + \beta_i$ and the possible $k$-fold junctions ($k > 2$). The formation of a $k$-junction replaces $k$ ends that contribute an energy of $\alpha/2$ by one vertex of energy $\alpha$ so that the overall energy change is $(1 - k/2)\alpha$. Performing the summation over all possible networks proves to be tricky because the vertex and edge occupations are not independent. Below, we employ the $n = 0$ formalism to resolve this configuration counting problem.

Correspondence Between Code Optimization and Spin Networks. The $n = 0$ formalism was devised by de Gennes to examine polymer solutions (11, 12). Recently, it was applied also to microemulsions, micellar solutions, and dipolar fluids (13–15). At the basis of this approach is a mathematical equivalence between a system of self-excluding polymers and a system of interacting $n$-component magnets, in the limit of vanishing number of components, $n \to 0$. The $n = 0$ formalism is reviewed elsewhere (12, 13). Here, we only discuss concisely the basic idea and use this approach to show that the statistics of the code-table (Eqs. 2 and 4) can be mapped to that of the zero-component magnets.

To demonstrate the equivalence between the code-table problem and the spin system, let us consider the dual of the symbol graph on which the polymer network resides (Fig. 1) and assign to each of the edges $i \to j$ an $n$-component magnetic spin $S_j$. The interaction is represented by a spin-Hamiltonian $H_S$ and the spin partition function is $Z_S = \langle \exp(-H_S) \rangle$, where $\langle \ldots \rangle$ denotes the average over all possible spin orientations. A peculiar feature of this system in the limit of zero components (“the $n = 0$ property”) is that all averages over products of spins vanish except for the quadratic averages $\langle S_j S_j \rangle = 1$, where $S_j$ is any of the components of $S_j$. This property enables mapping of the spin lattice to the network ensemble by tailoring a spin Hamiltonian $H_S$ and a consequent spin partition-function $Z_S$, in which an $\langle S_j \rangle$ term appears if and only if the corresponding edge $i \to j$ is occupied in the network partition function $Z_C$. It is shown below that this correspondence is accomplished by the Hamiltonian $H_S$.

$$H_S = -\sum_i h_i, \quad h_i = a_i \left( \prod_{j(i)} (1 + b_j S_j) - \sum_{j(i)} b_j S_j - 1 \right),$$

[5]

where $j(i)$ are all of the neighbors of $i$, and the coefficients $a_i$ and $b_j$ are related to the $\alpha$ and $\beta_i$ parameters (Eq. 4). The functions $h_i$ are the contributions of each vertex $i$ to $H_S$ and consist of all possible products of two or more edges around $i$. Each of these products corresponds to a possible edge occupation state in a network (Fig. 2). The one- and zero-edge configurations are forbidden in the network and are therefore subtracted from $h_i$ (the second and third terms).

How this form of the Hamiltonian $H_S$ ensures the equivalence is clear by expanding $Z_S$ in a power series, $Z_S = \langle \exp(-H_S) \rangle = \langle \prod_i \exp(h_i) \rangle = \langle \prod_i (1 + h_i + h_i^2/2 + \ldots) \rangle$. Because of the $n = 0$
averages of spin products. In this sum, the only nonvanishing term in the partition function is a product of the occupation variables.

To solve for the typical rate-distortion expression (e.g., ref. 6). More importantly, it provides insight into the “low-temperature” (low $w_C$) regime where the landscape of the code’s free energy is rugged and therefore hard to calculate.

To make use of the equivalence between the spin system and the coding network, we need to express the average network occupancies, $\bar{e}_i = (\bar{E}_i)$ and $\bar{v}_i = (\bar{V}_i)$ as a function of the average spin $\bar{s}_i$. The average edge-occupancy is given by

$$\bar{e}_ij = (1/2)\ln Z_S/\ln b_{ij} = g_{ij}/2, \text{a consequence of Eqs. 5–7.}$$

Likewise, the average vertex-occupancy is $\bar{v}_i = \partial \ln Z_S/\ln a_i = \bar{h}_i$ (see Methods). Thus, one can calculate the average network configuration (that is the average code) for any value of the fugacities $a_i$ and $b_{ij}$, or for the equivalent control parameters of the coding system, $w_D$, $w_C$, $r_D$, and $n_m$. This is demonstrated below, where a first-order “coding transition” is deduced from the spin formalism.

Mean-field models similar to the one used here are standard in $n = 0$ treatments of self-assembling systems, such as polymer and micellar solutions (14, 15) and networks (13). The basic idea of the mean-field approach is to replace the spin–spin interactions by an interaction of a single spin with an effective field ($a_{ij}$ polynomials). This procedure vastly simplifies the problem and avoids a relatively simple solution. However, this simplicity comes at the cost of disregarding the long-range spatial correlations between the spins and the vertices. This implies, for example, that one can estimate the mean connectivity in the network but cannot tell how many loops it contains. In general, a mean-field treatment merely approximates qualitatively the behavior of thermodynamic functions. However, the accuracy of this approximation improves when each spin interacts with many neighbors. Therefore, when the symbol graph is highly connected—such as the graph of the genetic code, where each codon has nine neighbors—the mean-field approximation is expected to be relatively accurate and provides a basis to more elaborate models.

A First-Order Coding Transition. The equivalence of the spin system and the code-table enables us to follow the evolution of the code-table in response to variation of the control parameters that govern its optimization: the cost weight $w_C$, the misreading matrix $r_D$, the diversity weight $w_D$, and the number of meanings $n_m$. These parameters are not independent but related through the spin fugacities, $a$ and $b_{ij}$, or the corresponding edge and vertex energies, $\alpha$ and $\beta$. It proves convenient to represent these relations in terms of the normalized diversity, $D = w_D/w_C + \ln n_m = -\ln a = \alpha w_C$, and the normalized misreading probabilities, $R_D = r_D/w_C = -\ln(a b_{ij}) = (\alpha + b_{ij})/w_C$.

To examine the response of the code-table to variation of the four control parameters, we consider, for the sake of simplicity, regular symbol graphs, in which all of the vertices and edges are equivalent. Regular symbol graphs are useful in biological models to one used here are standard in $n = 0$ treatments of self-assembling systems, such as polymer and micellar solutions (14, 15) and networks (13). The basic idea of the mean-field approach is to replace the spin–spin interactions by an interaction of a single spin with an effective field ($a_{ij}$ polynomials). This procedure vastly simplifies the problem and avoids a relatively simple solution. However, this simplicity comes at the cost of disregarding the long-range spatial correlations between the spins and the vertices. This implies, for example, that one can estimate the mean connectivity in the network but cannot tell how many loops it contains. In general, a mean-field treatment merely approximates qualitatively the behavior of thermodynamic functions. However, the accuracy of this approximation improves when each spin interacts with many neighbors. Therefore, when the symbol graph is highly connected—such as the graph of the genetic code, where each codon has nine neighbors—the mean-field approximation is expected to be relatively accurate and provides a basis to more elaborate models.

A First-Order Coding Transition. The equivalence of the spin system and the code-table enables us to follow the evolution of the code-table in response to variation of the control parameters that govern its optimization: the cost weight $w_C$, the misreading matrix $r_D$, the diversity weight $w_D$, and the number of meanings $n_m$. These parameters are not independent but related through the spin fugacities, $a$ and $b_{ij}$, or the corresponding edge and vertex energies, $\alpha$ and $\beta$. It proves convenient to represent these relations in terms of the normalized diversity, $D = w_D/w_C + \ln n_m = -\ln a = \alpha w_C$, and the normalized misreading probabilities, $R_D = r_D/w_C = -\ln(a b_{ij}) = (\alpha + b_{ij})/w_C$.

To examine the response of the code-table to variation of the four control parameters, we consider, for the sake of simplicity, regular symbol graphs, in which all of the vertices and edges are equivalent. Regular symbol graphs are useful in biological
context. For example, a regular graph may approximate the symbol graph of the genetic code, where each of the 64 codons has 9 neighbors (2). Regular graphs may also describe large symbol spaces, for example the space of DNA binding sites of the transcription system, whose structure is not exactly known but whose average coordination number q is well determined. Regularity of the symbol graph implies uniform misreading \( e = gs/2 \) and the ellipse \( 2q^2 + (2e - 1)^2 = 1 \) (Lower). (b) Phase diagram of the coding (network ↔ no network) transition (solid line) in the D-R plane. The dashed line bounds the region of metastability, beyond which the no-network state is destabilized. (C) Vertex occupancy v, edge occupancy e, and ratio meanings/symbol n_m/n_s along the transition line (see text).

In the D-R plane (Fig. 3B), the phase transition line approaches a straight line, \( R = (1 - 2q)D \), which corresponds to a \( q \)-fold junction whose “energy” equals the thermal energy \( \alpha + (q/2)\beta = w_c \). In other words, the system undergoes a phase transition when \( q \)-junctions become thermally excitable. Because \( q \)-junctions are the majority species, the emergent network is dense and highly connected. The transition line indicates various pathways that the system can take toward the formation of a network at a coding state: increasing the number of available meanings \( n_m \), increasing the diversity parameter \( w_D \), and decreasing the misreading \( r \).

At the transition, the number of meaning islands \( n_t \) jumps abruptly and becomes proportional to the number of symbols (Fig. 3C). \( n_t \) is given by Euler’s characteristic (Eq. 3) with the average edge occupancy \( e = gs/2 \) and vertex occupancy \( v = h \). This implies that the number of meanings per symbol is \( n_j/n_s = \chi/n_s + (p/2) - (p/q)v \), where \( p \) is the coordination number in the symbol graph and \( q \) in the dual graph. The curves of the vertex and edge occupancies approach a common high D limit, \( e, v \rightarrow e_0 = a(bs)^q \), where the resulting meanings/symbol ratio is \( n_j/n_s = p(1/2 - 1/q)e_0 \). For planar regular graphs, \( 1/p + 1/q = 1/2 \) and \( n_j/n_s = e_0 \). In Discussion, we examine possible effects of population dynamics on the evolution of the code.

Discussion

The mean-field solution allows us to draw an approximate fitness landscape where the evolution of the code takes place. In general, this code fitness landscape \( F_k(s) \) (see Eq. 6) resides in a high-dimensional code space, whose coordinates are the average spins \( s_j \) or their conjugates, the average edge occupancies \( e_j \). Each point in this space is a vector \( s = (s_j \ldots) \) of dimension equal to the number of edges, which represents a possible code. Symmetry may reduce this dimensionality, and for the regular symbol graph it becomes one-dimensional \( f(s) \) (Fig. 3). We imagine a population of “organisms,” simple information-processing systems, which compete according to the fitness of their codes. Each organism is depicted as a point in code space positioned at its code \( s \), and the population is described by a probability density \( \Psi(s) \). The preceding discussion assumed implicitly that, as the control-parameters change, the evolution of the code more or less follows the track of the optimum in code space. In other words, the population density is a delta-distribution located at the optimal \( F_0 \). We conclude this work by considering several more realistic scenarios. First, we discuss the possibility that the coding system is stuck at metastable suboptimal states. Then, we consider mutations and genetic drift that may broaden the population toward codes of less optimal fitness.

Metastable Suboptimal Codes. Even when the global optimum in the code fitness is at a network state, \( s_c \neq 0 \), the no-network state, \( s = 0 \), may remain locally stable for some parameter range (Fig. 3B). To locate the metastable state, one examines the curvature of the free energy at its \( s = 0 \) extremum, which in the case of an isotropic symbol graph is \( F' = \partial^2 f/\partial s^2 = -2(\alpha - 1)e_0 \). A metastable state exists as long as this curvature is positive. We find that the curvature changes its sign at the line \( R = \ln(2p - 2) \), which corresponds to a “monomer” (a vertex plus an edge) of energy \( \alpha + \beta = w_c \ln(2p - 2) \). This may be further clarified by considering the limit of the vertex/edge occupancy ratio near the no-network state, \( v/e \rightarrow q/2 \). Because there are \( q/2 \) edges per vertex, this indicates that the dominant building block of the dilute network is the monomer. Thus, a coding system becomes unstable exactly when the monomers become “thermally excitable,” compared to the \( q \)-junctions that are excitable at the coding transition.

Effects of Mutations and Genetic Drift in the Codes Space. So far, it was assumed that the population of organisms is sharply con-
centrated around a certain optimal or metastable, suboptimal code. This scenario applies to large populations at negligible mutation rates $\mu$. Let us consider two possible effects of population dynamics that may smear the population over the code space, mutations and genetic drift. These effects were analyzed in detail within the framework of rate-distortion theory (6) and are discussed here only schematically, in the context of the present polymer network model.

We consider a population that is localized around a fitness optimum $f_0$ at an optimal code $s_0$, where the landscape is approximately $f(s) = f_0 + \frac{1}{2} f''(s - s_0)^2$ [a regular graph with a one-dimensional fitness $f(s)$ is assumed for simplicity]. Mutations drive the organisms to diffuse into somewhat less optimal regions of the code space. This effect may be described in terms of reaction-diffusion dynamics of the probability distribution (6), $\partial_t \Psi = \nu \Delta \Psi - f(s) \Psi$, where the first term is due to mutations and the second represents reproduction at a rate $-f(s)$. It is straightforward to find that this dynamics tends to a steady state, in which the population is broadened into a Gaussian, $\Psi(s) \rightarrow \exp[-(f^2/2\nu)(s - s_0)^2]$ of width that scales like $\sim (\mu t)^{1/4}$ (6). It implies that the width of a population at a metastable state $s = 0$ will diverge near the metastability limit $f^* = 0$ just before the Gaussian migrates to the coding state ($s_C \neq 0$).

When the effective population size $N$ is relatively small, fluctuations in the reproduction rate, termed genetic drift, become significant. In this regime, the dynamics is characterized by long periods when the population is localized around a fitness optimum, which are separated by fast diffusive migrations to new optima (6). The dynamics of the distribution is known to reach a Boltzmann partition, $\Psi(s) \rightarrow \exp(-\beta f(s))$, where the population size plays the role of an inverse temperature. Relatively small populations (which are “hot” in this sense) are expected to be partitioned by genetic drift between the available fitness optima. For example, the two minima of the free energy $f(s)$ (Fig. 3) will be populated according to their fitness values.

In the previous sections, we have shown that each organism experiences “internal” noise due to stochastic molecular recognition in its coding system. The internal noise affects the fitness of the code through the error load and the cost (16). On top of this, mutations and genetic drift add “external” sources of noise, which may drive parts of the population away from the optimal code. The existence of metastable states may further delay the transition to a coding state. The present model and its conclusions suggest that the $n = 0$ polymer network formalism is a potential tool to study several other aspects of noisy coding systems.

Methods

Cost of a Code-Table. The cost of a code-table is traditionally measured by the mutual information $I$ between the symbols and the meanings that they encode, $I = S_{SW} + S_{S_Y} - S_{SWY}$, where $S_{SW}$ and $S_{S_Y}$ are the entropies of the meaning space and symbol space, respectively, and $S_{SWY}$ is the joint entropy of these two spaces. The entropy of meanings $S_{SWY}$ is determined by their given distribution $p_{SWY} = -\sum_p \ln p_{SWY}$ and similarly, $S_Y = -\sum_p \ln p_{S_Y}$. These are constant terms in the cost $I$ that we can neglect, and consider only the joint entropy $S_{SWY}$, which can be optimized by tuning the average partition pattern $\psi_{SWY}$ is simply the entropy of all of the possible coloring patterns as determined by all possible networks and the number of possible ways to color every such pattern. It follows that the cost is therefore minus the coloring entropy $I = -S_{SWY} - S_C$.

Graph Embedding and the Dual Graph. The embedded graph divides the surface into faces or cells, hexagons in our example (Fig. 1). Then, one finds the dual graph by the following correspondence (10): Every vertex in the symbol graph corresponds to a cell in the dual (a triangle in this example) whereas every cell in the symbol graph (a hexagon) corresponds to a vertex of the dual. The correspondence between the edges in the symbol graph and its dual is one-to-one, every edge corresponds to the edge that crosses it in the dual. The resulting dual graph is a triangular lattice. The hexagonal lattice is a regular graph in which all of the vertices have the same coordination number. However, the embedding procedure described here applies to any connected graph whether it is regular or not.

Mean-Field Approximation. The spin probability distribution decouples into a product of independent single-spin distributions, $P = \prod_i P_i$. We use a variational inequality, which sets an upper limit on the spin free energy, $F_M = \langle H_t \rangle + T I_p \ln \rho$, where $p$ satisfies probability conservation, $\langle p \rangle = 1$. We augment $F_M$ with a Lagrange multiplier to account for probability conservation, $L = F_M + \eta I_p$, and take the derivative $\partial L \partial \rho$. The resulting distributions are $P_i = \exp[g_i S_i]/\sum_i \exp[g_i S_i]$, where the effective fields are $g_i = \Delta h_i + \delta h_i S_i = S_i \Delta h_i L_i + (1 + S_i) - 1$ with $S_i = S_0 + \Delta h_i L_i$. From the $n$-property, it follows that $\exp[g_i S_i] = \sum S_i g_i^k / k!$. In $g_i = 1 + g_i(1/2)$ (Eq. 7). In a similar fashion, we obtain the mean-field approximation for the free energy, $F_M = \sum_i h_i - \sum_i \ln(1 + g_i/2)$, and it is easy to verify that Eq. 7 defines the extremum $\partial F_M / \partial \rho = 0$. Eq. 7 is analogous to the self-consistency relation of an Ising magnet, $S = \ln \theta/\cosh(\theta)$; the different form is due to the truncation of the power-series expansion of the hyperbolic functions thanks to the $n = 0$ property. Solving Eq. 7 for $g_i$ as a function of $\rho_i$ we find that the solution can be described graphically as the points where the function $g_i = \cosh(g_i) \cosh(\rho_i)$ crosses the ellipse $2g_i^2 + (2g_i - 1)^2 = 1$ (Fig. 3A).

Use of Euler’s Characteristic. When we apply Eq. 3, an underlying assumption is that the embedding is cellular (10); i.e., every meaning island is homeomorphic to an open disk. This is not necessarily true when the number of islands is less than the number of holes in the surface, that is the genus, $\gamma = 1 - 2/\chi$. In this case, some islands are expected to wrap between two holes and therefore are not homeomorphic to a disk. However, in the “thermodynamic limit” of many islands per hole, $n_i \ll \chi$ the embedding is mostly cellular and is well approximated by Eq. 3.

References