Energy-speed-accuracy relation in complex networks for biological discrimination
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An energy-speed-accuracy relation in complex networks for biological discrimination

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Discriminating between correct and incorrect substrates is a core process in biology but how is energy apportioned between the conflicting demands of accuracy ($\mu$), speed ($\sigma$) and total entropy production rate ($P$)? Previous studies have focussed on biochemical networks with simple structure or relied on simplifying kinetic assumptions. Here, we use the linear framework for timescale separation to analytically examine steady-state probabilities away from thermodynamic equilibrium for networks of arbitrary complexity. We also introduce a method of scaling parameters that is inspired by Hopfield’s treatment of kinetic proofreading. Scaling allows asymptotic exploration of high-dimensional parameter spaces. We identify in this way a broad class of complex networks and scalings for which the quantity $\sigma \ln(\mu)/P$ remains asymptotically finite whenever accuracy improves from equilibrium, so that $\mu_{eq}/\mu \rightarrow 0$. Scalings exist, however, even for Hopfield’s original network, for which $\sigma \ln(\mu)/P$ is asymptotically infinite, illustrating the parametric complexity. Outside the asymptotic regime, numerical calculations suggest that, under more restrictive parametric assumptions, networks satisfy the bound, $\sigma \ln(\mu/\mu_{eq})/P < 1$, and we discuss the biological implications for discrimination by ribosomes and DNA polymerase. The methods introduced here may be more broadly useful for analysing complex networks that implement other forms of cellular information processing.

I. INTRODUCTION

In cellular information processing, a biochemical mechanism is coupled to an environment of signals and substrates and carries out tasks such as detection [1–5], amplification [6–8], discrimination [9–24], adaptation [25], searching [26] and learning [27–30]. As Hopfield pointed out in his seminal work on discrimination [9], systems operating at thermodynamic equilibrium have limited information processing capability and energy must be expended to do better [8, 32].

We focus here on the widely-studied task of discrimination between correct and incorrect substrates, an essential feature of many core biological processes. The accuracy of discrimination may have to be traded off against speed while energy remains a limiting resource [25, 31]. How can energy be apportioned between such desirable properties as accuracy and speed and the inevitable dissipation of heat to the environment? Quantitative insights into this question can help us distill the principles underlying cellular information processing despite the pervasive complexity of the underlying molecular mechanisms.

Previous studies of discrimination have largely focussed on particular systems, such as Hopfield’s original proofreading mechanism [9, 19, 20], McKeithan’s T-cell receptor mechanism [14, 18], minimal feedback mechanisms [25], irreversible multi-step mechanisms [23], or ladder mechanisms [12, 15, 16, 21]. Murugan et al. analysed general systems using simplifying assumptions about where energy is expended and showed how discriminatory regimes also depend on the topology of the mechanism [15, 16]. Several studies have analysed the relationship between energy expenditure and other properties away from thermodynamic equilibrium. These have often been limited to networks with simplifying assumptions [11–13, 15, 19–24] or have considered different questions in the context of kinetic proofreading [13, 15, 16, 22–24, 33, 34]. The results in [17, 25, 35] show some formal similarities to those presented in this paper and these studies are reviewed further in the Discussion (§IX).

One of the challenges in dealing with general systems away from thermodynamic equilibrium is that the steady-state probabilities can be complex algebraic functions of the parameters (see the Discussion) [8, 32], which makes it difficult to identify any universal behaviour. We address this issue here in two ways. First, we use a graph-based treatment of Markov processes called the “linear framework” [36], which allows steady-state probabilities to be analytically calculated for processes of arbitrary structure away from thermodynamic equilibrium (§II, §III). Second, we introduce a way of exploring parameter space by scaling the parameters. This idea is inspired by Hopfield’s original analysis of kinetic proofreading, which we revisit here to point out certain subtleties that are not always appreciated (§IV). The scaling method allows us to calculate the asymptotic behaviour of steady-state properties of general systems, despite the difficulties arising from high-dimensional parameter spaces and algebraic complexity. In this way, we are able to exhibit a universal asymptotic relationship between energy, speed and accuracy for a broad class of discriminatory systems, without simplifying assumptions as to where energy is expended (§VI). We further explore whether this asymptotic relationship also has significance for finite parameter values and for actual biological discrimination mechanisms (§VIII).

II. THE LINEAR FRAMEWORK

The linear framework [36–38] is a graph-based interpretation of biochemical processes which has been used to analyse protein post-translational modification [39], covalent modification switches [40] and eukaryotic gene regulation [8, 32]. In the stochastic setting considered here, the framework follows the treatment previously developed by Hill [41] and Schnakenberg [42]. A finite-state Markov process $M$ is repre-

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sented by a directed graph, \( G \), with labelled edges and no self-loops (Fig. 1(a)), hereafter a “graph”. The vertices \( 1, \ldots, n \), are interpreted as microstates and a labelled edge, \( i \xrightarrow{a} j \), as a transition between microstates whose label, \( a \), is the infinitesimal transition rate of the Markov process. The equations of motion are defined as if the edges are reactions under mass action kinetics, with the labels being the corresponding rates.

This yields the master equation of the Markov process in the form \( \frac{dp_i(t)}{dt} = \mathcal{L}(G)p_i(t) \), in which \( p \in (\mathbb{R}_{\geq 0})^n \) is the vector of microstate probabilities and \( \mathcal{L}(G) \) is the Laplacian matrix of \( G \). For instance, for the subgraph \( G_C \) in Fig. 1(a),

\[
\mathcal{L}(G_C) = \begin{pmatrix}
-k'_C + l'_C & k_C & l_C + W \\
-k'_C & k_C + m' & m \\
l'_C & m' & -(l_C + W + m)
\end{pmatrix}.
\]

Since probability is conserved, there is a conservation law, \( \sum_i p_i(t) = 1 \), where \( p_i(t) \) is the probability that \( M \) is in microstate \( i \) at time \( t \).

If the graph is strongly connected, so that any two vertices can be joined by a path of edges in the same direction, then there is a unique steady state up to a scalar multiple. A representative steady state, \( \rho(G) \), can be calculated in terms of the labels by the Matrix-Tree theorem (MTT): if \( \Theta_i(G) \) denotes the set of spanning trees rooted at \( i \) (Fig. 1(c)), then \( \rho_i(G) \) is the sum of the product of the labels on the edges of each tree,

\[
\rho_i(G) = \sum_{T \in \Theta_i(G)} \prod_{j \xrightarrow{a} k \in T} a.
\]

The steady-state probabilities are then

\[
p_i^* = \left( \frac{\rho_i(G)}{\rho_1(G) + \cdots + \rho_n(G)} \right).
\]

If the steady state is one of thermodynamic equilibrium, so that detailed balance is satisfied, then the framework gives the same result as equilibrium statistical mechanics, with the denominator in equation (2) being the partition function (up to a constant factor). However, equation (2) is also valid away from equilibrium.

In contrast to eigenvalues or determinants, the MTT gives the steady state analytically in terms of the labels (equation (1)). This makes it feasible to undertake a mathematical analysis, without relying on numerical simulation, whose scope is necessarily more restricted. Substantial algebraic complexity can arise in equation (1) through the combinatorial explosion of enumerating spanning trees (Discussion) but, as we show here, with the appropriate mathematical language, it is possible to draw rigorous conclusions about structurally complex systems away from thermodynamic equilibrium. In particular, we exploit the fact that steady-state probabilities are rational functions in the labels, which allows us to determine asymptotic behaviors.

**III. STEADY STATES OF A BUTTERFLY GRAPH**

Discrimination typically requires a mechanism for choosing a correct substrate from among a pool of available substrates, as in DNA replication, in which DNA polymerase must choose at each step one correct deoxyribonucleoside triphosphate from among the four available (dATP, dGTP, dCTP, dTTP). We follow Hopfield in assuming a single correct substrate, \( C \), and a single incorrect substrate, \( D \), and describe this mechanism by a graph \( G \) (e.g. Fig. 1(a)) whose vertices represent the microstates of the discriminatory mechanism, such as DNA polymerase in the case of replication. This graph is naturally composed of two subgraphs, \( G_X (X = C, D) \), corresponding to the states in which substrate \( X \) is bound. \( G_C \) and \( G_D \) share a common vertex, but no edges, so that \( G \) has a butterfly shape.

We will denote such a butterfly graph \( G = G_C \oplus_v G_D \), where \( v \) is the shared vertex. For the task of discrimination, the subgraphs \( G_X \) are structurally symmetric, with symmetric vertices, \( 1_X, \ldots, n_X \), of which \( 1_C = 1_D = 1 \) is shared, and symmetric edges, \( i_C \rightarrow j_C \) if, and only if, \( i_D \rightarrow j_D \). The labels on these corresponding edges may, however, be distinct. The vertices \( i_X \) with \( i > 1 \) are the microstates in which \( X \) is bound, while vertex 1 is the empty microstate in which no \( X \) is bound. All directed edges are assumed to be structurally reversible, so that, if \( i_X \rightarrow j_X \), then \( j_X \rightarrow i_X \). The graphs \( G_C, G_D, \) and \( G \) are therefore all strongly connected.

Let \( G = G_C \oplus_v G_D \) be any butterfly graph. Even if \( G_C \) and \( G_D \) are not structurally symmetric, as above, it follows readily from equation (1) that

\[
\rho_i(G_C \oplus_v G_D) = \begin{cases} 
\rho_i(G_C)\rho_v(G_D) & \text{if } i \in G_C \\
\rho_i(G_D)\rho_v(G_C) & \text{if } i \in G_D.
\end{cases}
\]
IV. THE ERROR FRACTION FOR THE HOPFIELD MECHANISM

The original Hopfield kinetic proofreading mechanism is described by the discriminatory butterfly graph $G = G_C \oplus_1 G_D$ in Fig. 1(a). The substrates $C$ and $D$ are treated as “slow” components and assumed to have constant concentration over the timescale of interest. These concentrations are absorbed into the “on-rates” $k'_C, k'_D, W, l'_C, l'_D$. The discrimination mechanism itself is assumed to have the “fast” components and to be at steady state. The rate $W$ for exit from $3_X$ ($X = C, D$) corresponds to product generation and release of $X$, so that the overall system is open whenever $W > 0$, with $C$ and $D$ being transformed into correct and incorrect product, respectively.

In this mechanism, discrimination occurs twice, through binding and unbinding of $X$ to form $2_X$ and to form $3_X$. It is assumed that unbinding, rather than binding, causes discrimination, as is often the case in biology [14], so that $l'_C = l'_D$ and $k'_C = k'_D$. The correct substrate has a longer residence time, so that $k_C < k_D$, which reflects the free energy difference of $\Delta > 0$ between $2_C$ and $2_D$ (Fig. 1(b)): if energy is measured in units of $k_B T$, where $k_B$ is Boltzmann’s constant and $T$ is the absolute temperature, then $l_D = k_D e^{\Delta}$. There is assumed to be no difference in discrimination between $2_X$ and $3_X$, so that $k_C / k_D = l_C / l_D = e^{-\Delta} < 1$.

Hopfield defines the steady-state error fraction, $\varepsilon$ as the probability ratio of the incorrect to the correct exit microstate, which, using equation (2), is given by $\varepsilon = \rho_{3D} / \rho_{3C} (\varepsilon$ is the inverse of the accuracy $\mu$ in the Abstract; we will work with the former). Using equations (1) and (3),

$$\varepsilon = \frac{l'_D (k_D + m') + m' k'_D)}{(l'_C (k_C + m') + m' k'_C)} \frac{(W + l_C) + m k_C}{(W + l_D) + m k_D}. \tag{4}$$

If the overall system remains closed, so that $W = 0$, while the mechanism operates at thermodynamic equilibrium, then it has the error fraction, $\varepsilon_0 = k_C / k_D = l_C / l_D = e^{-\Delta}$ (Supplementary Material). If the overall system becomes open, so that $W > 0$, while the mechanism remains at equilibrium, then $\varepsilon$ increases monotonically with increasing $W$ (Supplementary Material). If the mechanism itself operates away from equilibrium, then

$$\varepsilon > \varepsilon_0 \frac{l_C + m + W}{l_D + m + W} > \varepsilon_0^2 \tag{5}$$

for all positive values of the parameters (Supplementary Material). Hopfield shows that $\varepsilon$ approaches the minimal error, $\varepsilon_0^2$, as certain parametric quantities become small (Supplementary Material) and suggests how this could be achieved in practice by expending energy to drive the transition from $2_X$ to $3_X$ through the label $m'$. This is kinetic proofreading.

There are two aspects of Hopfield’s analysis which have not always been fully appreciated. First, increasing $m'$ is not sufficient for $\varepsilon$ to approach $\varepsilon_0^2$. Indeed, it follows from equation (4) that, when $W = 0, \varepsilon \to \varepsilon_0$ as $m' \to \infty$. Too much energy expenditure can increase the error fraction, which behaves non-monotonically with respect to $m'$. (Similar non-monotonicity has been observed for kinetic proofreading with the T-cell receptor mechanism in Supplementary Fig. 1 [18].) The parameter $m'$ must neither be too high nor too low for the error fraction to approach $\varepsilon_0^2$. Second, parameters other than $m'$, $m$ and $W$ must also take adequate values for the accuracy to approach this bound: the “on-rate” for $1 \to 2_X$ must be much larger than that for $1 \to 3_X$, so that $l'_D / k'_D = l'_C / k'_C \to 0$ (Supplementary Material). The lower bound of $\varepsilon_0^2$ is only reached asymptotically as several parameters take limiting values.

For more complex systems, the appropriate parameter regime for the minimal error is not readily found using Hopfield’s approach. We therefore sought an alternative strategy. If we let $x = e^\Delta = \varepsilon_0^{-2}$ and substitute $k_D = x k_C$ and $l_D = x l_C$ into equation (4), we see that, if no other parameters change, the error fraction $\varepsilon$ behaves like $x^{-1}$ as $x$ increases. We reasoned that to approach the minimal error of $x^{-2}$, the fold change in other parameter values should be some function of $x$. By retaining only the highest-order term in $x$ as $x \to \infty$, the behaviour of $\varepsilon$ could be determined while bypassing the parametric complexity. Let $R(x) \sim Q(x)$ mean that $R(x) / Q(x) \to c$ as $x \to \infty$, where $0 < c < \infty$. It can be seen from equation (4) that if either $k'_D = k'_C \sim x$ or $l'_C = l'_D \sim x^{-1}$, while none of the remaining parameters depend on $x$, then $\varepsilon \sim x^{-2} = \varepsilon_0^2$. This scaling of the “on-rates” corresponds to what was required in the previous paragraph for Hopfield’s limiting procedure. This suggests a strategy for exploring parameter space that can be extended to more complex systems. We exploit this below to examine the relation between energy, speed and accuracy.

V. DISSOCIATION-BASED MECHANISMS

We introduce here a class of discrimination mechanisms for which such a relation can be determined. We consider a discriminatory butterfly graph of the form $G = G_C \oplus_1 G_D$ consisting of structurally symmetric subgraphs $G_C$ and $G_D$ of arbitrary complexity. The vertex $n_X$ is taken to be the only exit microstate in which product is generated, so that there is a return edge $n_X \to 1$. No further structural assumptions are made but the product generation rate, $W$, makes an additive contribution to the label of the return edge $n_X \to 1$, as in Fig. 1(a).

Multiple internal microstates and transitions are allowed in $G_X$ as well as multiple returns to the empty microstate, 1, although only a single one of these, through $n_X$, also generates product. As in Hopfield’s original scheme, we think of the mechanism as coupled to sources and sinks of energy, which may alter the edge labels. In Hopfield’s scheme, the labels on edges which do not go to the reference microstate 1 were assumed to be the same between $C$ and $D$ (Fig. 1(a)). In other words, there was no “internal discrimination” between correct and incorrect substrates. Here, we allow internal discrimination between $C$ and $D$: when $j \neq 1$, the label on $i_C \to j_C$ may be different from that on $i_D \to j_D$.

Graphs of this form have been widely employed in the literature. In addition to the original Hopfield mechanism (Fig. 1(a)), they include the “delayed” mechanism [10], the multi-step mechanism [12, 13, 43], the T-cell receptor mechanism
FIG. 2. Structure of dissociation-based mechanisms. Shown is a schematic of a labelled, directed “butterfly” graph illustrating the structure of any dissociation-based mechanism, which consists of the subgraph $G_D$ for the incorrect substrate $D$ and the subgraph $G_C$ for the correct substrate $C$ sharing the common vertex 1. $G_C$ and $G_D$ are assumed to be symmetric, but the internal transitions within them can be arbitrary, as suggested by the “clouds.” $n_X$ is taken to be the only exit microstate in which product is generated, so there is a return edge $n_X → 1$. As in Fig. 1(a), cyan and magenta denote correct and incorrect substrate binding, respectively.

[14, 18] and generalised proofreading mechanisms [15–17].

We follow Hopfield in using the steady-state error fraction and work from now on with probabilities. Let $p^*$ be the vector of steady state probabilities. The discrimination error fraction, $ε$, is the steady-state probability ratio of the incorrect exit microstate, $n_D$, to the correct exit microstate, $n_C$,

$$\varepsilon = \frac{p^*_{n_D}}{p^*_{n_C}}. \quad (6)$$

We will analyse the behaviour of $G$ under the assumption that some of the labels are functionally dependent on the non-dimensional variable $x \in \mathbb{R}$. A function $R(x)$ is said to be allowable if it is positive, $R(x) > 0$ for $x > 0$, and has a degree, $\deg(R)$, given by $R(x) \sim x^{\deg(R)}$ as $x \to \infty$. This is well defined because $x^a \sim x^b$ if, and only if, $a = b$. The degree determines the asymptotics of allowable functions: $R \sim Q$ if, and only if, $\deg(R) = \deg(Q)$. Note that $\deg(R) = 0$ if, and only, $R(x) \to c$ as $x \to \infty$, where $c > 0$, is the case if $R$ does not depend on $x$.

The labels in the graph $G$ are assumed to be allowable functions of $x$. (The product generation rate $W$ couples the mechanism to the environment and is assumed not to depend on $x$.) If $R$ and $Q$ are allowable functions, then so are $R^{-1}$, $RQ$, and $R + Q$ and (Supplementary Material)

$$\begin{align*}
\deg(R^{-1}) &= -\deg(R) \\
\deg(RQ) &= \deg(R) + \deg(Q) \\
\deg(R + Q) &= \max(\deg(R), \deg(Q)).
\end{align*} \quad (7)$$

Accordingly, any rational function of the labels with only positive terms, such as $p^*$, which acquires this structure through equation (2) and equation (1), or $\varepsilon$, which acquires it through equation (6), becomes in turn an allowable function of $x$.

We define a dissociation-based mechanism to be a general discrimination mechanism for which, for the edges between the exit microstates and 1,

$$\begin{align*}
\deg(\ell_{i \to n_D}) &= \deg(\ell_{i \to n_C}) \\
\deg(\ell_{n_D \to 1}) &= \deg(\ell_{n_C \to 1}) + 1. \quad (8)
\end{align*}$$

Here, we use $\ell_{i \to j}$ to denote the label on the edge $i \to j$. Eq. 8 is analogous to the assumption $\ell'_j = \ell'_D$ and $xI_C = \ell_D$ for the Hopfield mechanism. Unlike the Hopfield mechanism, we do not restrict what happens at non-exit microstates.

With such general assumptions on the labels, a dissociation-based mechanism may not reach thermodynamic equilibrium. However, if it can, with $W > 0$, so that the overall system remains open, then equation (8) ensures that the equilibrium error fraction, $\varepsilon_{eq}$, has a simple form. Since detailed balance requires that each pair of edges is independently at steady state [36], the exit states, $n_X$, satisfy $\ell_{n_X \to 1} p^*_{n_X} = \ell_{1 \to n_X} p^*_1$, so that

$$\varepsilon_{eq} = \frac{p^*_{n_D}}{p^*_{n_C}} = \frac{\ell_{1 \to n_D}}{\ell_{1 \to n_C}} \frac{\ell_{n_C \to 1}}{\ell_{n_D \to 1}}. \quad (9)$$

Applying equation (8) and using equation (6), we see that, if equilibrium is reached, the resulting error fraction, $\varepsilon_{eq}$, satisfies

$$\varepsilon_{eq} \sim x^{-1}. \quad (10)$$

VI. THE ASYMPTOTIC RELATION

We now define the measures of speed and energy expenditure in terms of which our main result will be stated. A reasonable interpretation for the speed of the mechanism, $\sigma$, is the steady-state flux of correct product [44],

$$\sigma = W p^*_{n_C}. \quad (11)$$

As for energy expenditure, this is determined at steady state by the rate of entropy production. Schnekenberg put forward a definition of this [42] that has been widely used [2, 7]: for a pair of reversible edges, $i \rightleftharpoons j$, the steady-state entropy production rate, $P(i \rightleftharpoons j)$, is the product of the net flux, $J(i \rightleftharpoons j) = \ell_{j \to i} p^*_j - \ell_{i \to j} p^*_i$, and the thermodynamic force, $A(i \rightleftharpoons j) = \ln(\ell_{j \to i} p^*_j/\ell_{i \to j} p^*_i)$:

$$P(i \rightleftharpoons j) = J(i \rightleftharpoons j) A(i \rightleftharpoons j). \quad (12)$$

Here, we omitted Boltzmann’s constant $k_B$ for convenience, so that $P$ has units of (time)$^{-1}$. If $T$ is the absolute temperature, then $k_B T P(i \rightleftharpoons j)$ is the power irreversibly expended through $i \rightleftharpoons j$. The total entropy production rate of the system is then given by $P = \sum_{i \rightleftharpoons j} P(i \rightleftharpoons j)$. Note that $P(i \rightleftharpoons j) \geq 0$ (and so also $P \geq 0$) with equality at thermodynamic equilibrium when detailed balance implies that $J(i \rightleftharpoons j) = 0$. Positive entropy production, with $P > 0$, signifies energy expenditure away from thermodynamic equilibrium.

Both $\sigma$ and $P$ are functions of $x$ and $\sigma$ is evidently allowable. However, $A(i \rightleftharpoons j)$ is not a rational function with positive terms and $\ln(x) \not\sim x^\alpha$ for any $\alpha$, so $P(i \rightleftharpoons j)$ and $P$ are not allowable functions. Nevertheless, the asymptotic behaviour of $P$ can be estimated. Some further notation is helpful to do this. If $R(x)$ and $Q(x)$ are functions which are not necessarily allowable, then $R \prec Q$ means that $R/Q \to 0$ as
This relation is transitive, so that, if $S \prec R$ and $R \prec Q$, then $S \prec Q$. If both functions are allowable, then $R \prec Q$, if, and only if, $\deg(R) < \deg(Q)$. We will say that $R \preceq Q$ if $R/Q \to c$, where $0 \leq c < \infty$, and corresponding remarks about transitivity and allowable degrees hold for this relation. Note that $\prec$ and $\preceq$ dominate over $\sim$ when forming products, so, for instance,

$$
if \ T \preceq S \ and \ R \sim Q \ then \ RT \preceq SQ,
$$

which we will make use of below.

Each summand $P(i \rightleftharpoons j)$ has the form $(R - Q)\ln(R/Q)$, where $R$ and $Q$ are allowable. Let $\alpha = \deg(R)$, $\beta = \deg(Q)$ and $c_1 = \lim R/x^\alpha$ and $c_2 = \lim Q/x^\beta$ as $x \to \infty$. By definition, $c_1, c_2 > 0$. Note that, if $S$ is allowable, then (Supplementary Material)

$$
\ln(S) \sim \begin{cases} 
\ln(x) & \text{if } \deg(S) > 0 \\
\ln(x^{-1}) & \text{if } \deg(S) < 0.
\end{cases}
$$

The asymptotic behaviour of $P(i \rightleftharpoons j)$ then falls into the following three cases (Supplementary Material), as specified on the right:

- **case 1:** $\alpha \neq \beta \sim x^{\max(\alpha, \beta)} \ln(x)$
- **case 2:** $\alpha = \beta, c_1 \neq c_2 \sim x^\alpha$
- **case 3:** $\alpha = \beta, c_1 = c_2 \prec x^\alpha$.

The third case is awkward because the leading-order asymptotics are lost, which leads to the $\prec$ relation instead of $\sim$. However, $c_1$ and $c_2$ are rational expressions in the parameters which do not involve $x$ and the equation $c_1 = c_2$ defines a hypersurface in the space of those parameters. The reversible edges which fall into case 3 therefore determine a set of measure zero in the space of parameters. Provided this set is avoided, the asymptotic behaviour of the summands in $P$ fall into the first two cases and can be controlled. In particular, suppose that the total entropy production rate $P$ is written as $P = \sum P_u$, where $P_u$ is a term coming from a pair of reversible edges $i \rightleftharpoons j$, as in equation (12). In Appendix A, we show that, if $P_k$ is any summand in case 1 of equation (15), then, outside the measure-zero set defined by case 3, $P_k \preceq P$.

Let us now assume, for any dissociation-based mechanism as defined previously, that

$$
\varepsilon(x) \prec x^{-1}.
$$

This forces the error fraction to be asymptotically better than if the system were able to reach equilibrium (equation (10)) and thereby ensures that energy expenditure is contributing to an improvement in accuracy. Consider any general discrimination mechanism which is dissociation-based, as described in equation (8). If its error fraction obeys equation (16) then, outside the measure-zero set in parameter space defined by case 3 of equation (15), we show in Appendix B that the mechanism satisfies the asymptotic relation,

$$
\sigma \ln(\varepsilon^{-1}) \preceq P.
$$

The exact asymptotics of $\sigma \ln(\varepsilon^{-1})/P$ are difficult to estimate for a general dissociation-based mechanism with allowable labels because each pair of reversible edges must be examined. However, for the Hopfield mechanism (Fig. 1(a)), under the conditions described above for which $\varepsilon \sim x^{-2}$, we find (Supplementary Material)

$$
\lim_{x \to \infty} \frac{\sigma \ln(\varepsilon^{-1})}{P} = \frac{2W}{\varepsilon c + W}
$$

outside the parametric set of measure-zero noted above.

## VII. A NON-DISSOCIATION BASED MECHANISM

The requirements in equation (8) for being dissociation-based are necessary for the validity of equation (17). In the Supplementary Material, we consider a discrimination mechanism with a structure identical to that of the Hopfield mechanism (Fig. 1(a)) but with labels that do not follow equation (8) (Supplementary Fig. 3). If the mechanism reaches thermodynamic equilibrium, then it follows from equation (9) that its equilibrium error fraction satisfies $\varepsilon_{eq} \~ x^{-1}$. However, with a particular choice of allowable functions for the labels, for which the mechanism is no longer at equilibrium, its error fraction improves asymptotically, with $\varepsilon \sim x^{-3/2}$, while its speed remains constant, $\sigma \sim 1$, and its entropy production is either constant or vanishes, $P \preceq 1$, outside a set of measure zero. This evidently does not obey equation (17) and shows the existence of a different asymptotic interplay between energy, speed and accuracy.

## VIII. NUMERICAL CALCULATIONS OUTSIDE THE ASYMPTOTIC REGIME

To examine further the energy-speed-accuracy relation found by the asymptotic analysis above, we used more restrictive assumptions on the allowable labels to facilitate numerical exploration. We considered discrimination-based mechanisms in which the $x$-dependency was similar to Hopfield’s original analysis. For any return edge to 1 from a non-exit microstate, we assumed that

$$
\ell_{i_D \to 1} = \ell_{i_C \to 1} x (i \neq n),
$$

with an additive contribution of $W$ in the exit microstate $(i = n)$, $\ell_{n_D \to 1} = ax + W$, $\ell_{n_C \to 1} = a + W$ with $a \in \mathbb{R}_{>0}$. As for the other edges, we assumed no internal discrimination, so that the labels were the same for $C$ and $D$,

$$
\ell_{i_D \to j_D} = \ell_{i_C \to j_C} (j \neq 1),
$$

with no $x$-dependency. By equation (9), the equilibrium error function when the system is closed ($W = 0$) satisfies $\varepsilon_0 = x^{-1}$. We set $x = e^{20}$, sampled the values $\ln(\ell_{i_C \to j_C})$, $\ln(a)$, and $\ln(W)$ uniformly in $[-100, 100]$, and determined $\hat{\ell}_{i_D \to j_D}$ from equations (19) and (20). We plotted $P/\sigma$ against $\ln(\varepsilon_0/\varepsilon)$, when $\varepsilon < \varepsilon_0$, for the Hopfield mechanism (Fig. 3(a)), the T-cell receptor mechanism (Supplementary Fig. 1) and for a mechanism different from both of these (Supplementary Fig. 2). In each case, the resulting region was confined
FIG. 3. Numerics for the Hopfield mechanism. (a) Plot of $P/\sigma$ against $\ln(\varepsilon_0/\varepsilon)$ for the Hopfield mechanism (Fig. 1(a)) for approximately $10^5$ points. The sampling and the dashed lines are described in the text. (b) Similar plot to (a) for the Hopfield mechanism with internal discrimination between correct and incorrect substrates, as described in the text, with the light blue points having a lower asymmetry range ($A = 1$) and the dark blue points having a higher range ($A = 5$). The coloured overlays represent values from experimental data for ribosomes (orange, red and brown regions) and DNA polymerase (green point), with the former being samples of values estimated for a parameter for which no experimental data exists. Only those samples for which $\varepsilon > \varepsilon_0$ are shown and the asterisks, *, give the averages of the plotted values. The inset gives the plotted averages (for the ribosome variants) and values (for DNAP) of error fractions, $\varepsilon$ and $\varepsilon_0$, speed, $\sigma$ and entropy production rate, $P$ (all in units of $s^{-1}$). The data from which these values were calculated are shown in Supplementary Table 1. See [45] and the caption of Supplementary Table 1 for more details.

to the left of a vertical line (Fig. 3(a), black dashed line) and above the diagonal (Fig. 3(a), red dashed line). For the Hopfield mechanism, the vertical bound comes from equation (5) and similar bounds on $\varepsilon$ exist for the other mechanisms (not shown). The diagonal bound, however, is unexpected and implies the bound

$$\sigma/\ln(\varepsilon_0/\varepsilon) < P \quad (\text{for } \varepsilon < \varepsilon_0)$$  \hspace{1cm} (21)

for finite parameter values. It is possible that equation (21) holds for any discrimination-based mechanism whose edge labels satisfy equations (19) and (20).

The calculations leading to equation (21) assumed no internal discrimination between correct and incorrect substrate, as specified in equation (20). We were interested to find that experimental data for ribosomes and DNA polymerase, based on the original Hopfield mechanism, showed substantial internal discrimination, extending even to the product generation rate $W$ [19]. To examine the impact of this, we proceeded as follows. For any return edge to 1 from a non-exit microstate, we introduced an asymmetry between $C$ and $D$ so that

$$\ell_{1D} = \alpha_{11} \ell_{1C} \ell_{1C} (i \neq n).$$  \hspace{1cm} (22)

For the exit state ($i = n$), the product generation rate makes an additive contribution, $W_X$, which now depends on the substrate $X$, so that $\ell_{nC} = a + W_C$ and $\ell_{nD} = \alpha_{11} ax + W_D$, where $a \in \mathbb{R}_{>0}$ and $W_D = \alpha_W W_C$. For the other edges, we similarly introduced an asymmetry

$$\ell_{iD \rightarrow jD} = \alpha_{ij} \ell_{iC \rightarrow jC} (j \neq 1).$$  \hspace{1cm} (23)

The multiplicative factors $\alpha_{ij}$ and $\alpha_W$ carry the asymmetry between $C$ and $D$ in internal discrimination.

In view of the asymmetry in product generation rates, it is natural to redefine the error fraction as

$$\varepsilon = \frac{W_D p_{1D}}{W_C p_{1C}}.$$

Using equation (9), the equilibrium error fraction when the system is closed ($W = 0$) is given by $\varepsilon_0 = \ell_{1 \rightarrow nD} \alpha_{1n}/(\ell_{1 \rightarrow nC} \alpha_{1n}).$

We chose the asymmetry factors by sampling $\ln(\alpha_{ij})$ and $\ln(\alpha_W)$ uniformly in the range $[-A, A]$, for $A = 1$ and $A = 5$, and chose the other parameters as described previously for Fig. 3(a). Fig. 3(b) shows that both the vertical bound and the diagonal bound in Fig. 3(a) are broken, with the extent of the breach increasing with increase in the asymmetry range from $A = 1$ (Fig. 3(b), light blue points) to $A = 5$ (Fig. 3(b), dark blue points). Similar results were found for the other mechanisms that we numerically calculated (Supplementary Figs. 1(c) and 2(c)). We see that the absence of internal discrimination is essential for the vertical and diagonal bounds shown in Fig. 3(a) and Supplementary Figs. 1(b) and 2(b).

Banerjee et al. have provided parameter values for the Hopfield mechanism based on experimental data for discrimination in mRNA translation by the E. coli ribosome, including also an error-prone and a hyperaccurate mutant, and in DNA replication by the bacteriophage T7 DNA polymerase (DNAP) [19]. We used these parameter values to calculate entropy production, speed and accuracy as defined here and overlaid the resulting $\ln(\varepsilon_0/\varepsilon)$, $P/\sigma$ points on the previous numerical calculation (Fig. 3(b)) [45].

The data show a striking difference between mRNA translation and DNA replication (Fig. 3(b)). All three ribosome variants (orange, wild type; brown, error-prone; red, hyperaccurate) have much higher $P/\sigma$ values than DNAP (green), with the former lying comfortably above the diagonal bound given by (21) and the latter lying well below. Nevertheless, all systems exhibit substantial internal discrimination (Supplementary Table 1). As the inset in Fig. 3(b) shows, the separation between translation and replication arises from a decrease of two orders of magnitude in entropy production rate and an increase of two orders of magnitude in speed. Furthermore, DNAP not only shows the smallest error fraction, $\varepsilon$, by three orders of magnitude, but also the greatest fold change over the equilibrium error fraction, $\varepsilon_0/\varepsilon$. In contrast, the ribosome variants, while showing the expected differences in error fraction, have lower fold changes over their equilibrium error fractions. Evolution seems to have tuned the energy dissipation, speed and accuracy of the replication machinery to a much greater degree than the translation machinery.
IX. DISCUSSION

One of the challenges in dealing with nonequilibrium systems in general has been the algebraic complexity, as epitomized by the enumeration of spanning trees in the MTT, Eq. (1). For a complicated graph, the combinatorial explosion in enumerating spanning trees can be super-exponential in the size of the graph [8]. This difficulty may have been apparent to earlier workers like Hill [41] and Schnakenberg [42] and may have discouraged a more analytical approach. The combinatorial complexity has largely been avoided by focussing on simple or highly-structured examples and by astute use of approximation.

In this paper, we have developed a way to address this complexity that is inspired by Hopfield’s analysis of kinetic proofreading. Here, the minimum error fraction can only be reached asymptotically (equation (5)) and only when multiple labels change their values consistently. This has suggested a method of exploring parameter space by treating the labels as allowable functions of a scaling variable $x$. In this way, a system of arbitrary structure can be analysed away from equilibrium, with relaxed assumptions on how energy is being deployed, while rising above the combinatorial explosion from history dependence.

Perhaps the most interesting conclusion from this analysis is the emergence of the quantity $\sigma \ln(\mu)/P$. Our main result, as expressed in equation (17), says that this quantity is asymptotically finite, for any graph obeying the dissociation-based condition on exit edges (equation (8)) and for any scheme of allowable scaling through which energy increases ($\text{deg} > 0$) or decreases ($\text{deg} < 0$) the rates, provided that the accuracy improves over equilibrium (equation (16)).

The advantage of the asymptotic analysis undertaken here is that it reveals a universal behaviour in $\sigma \ln(\mu)/P$ that transcends network structure and parametric complexity. Interestingly, our numerical calculations suggest that universality may still be found for finite parameter values, in the form of the bound in equation (21), as shown in Fig. 3 and Supplementary Figs. 1 and 2. However, this bound depends crucially on the absence of internal discrimination between correct and incorrect substrates, in contrast to the asymptotic behaviour in equation (17), for which internal discrimination is allowed. Experimental data shows that evolution has discriminated internally to a substantial extent but with very different effects on this bound. All E. coli ribosome variants for which we have data comfortably obey the bound, while the T7 DNA polymerase breaks it. This reflects a striking reduction in $P/\sigma$ for the latter, with far less difference between the ribosomes and the DNA polymerase in the fold change over their equilibrium error fractions (Fig. 3(b)). It would be interesting to know if these same comparative relationships are maintained for other ribosomes and polymerases. While recent work has shown that local trade-offs between speed and accuracy can differ markedly between different parametric regions [19], the quantities introduced here may be helpful for more global comparisons between discriminatory mechanisms.

As noted in the Introduction, the previous work of [17, 25, 35] shows formal similarities to the results presented here and it may be helpful to clarify these connections. In [35], England and colleagues determine the minimal energy dissipation cost for a general finite-state Markov process to maintain an arbitrary nonequilibrium steady-state. While their setting is similar to our graph-theoretic approach, they assume that energy is introduced through additional control transitions and their analysis of maintenance does not involve notions of speed or accuracy as used to analyse discrimination. In [25], Tu and colleagues study the costs of adaptation and derive an approximate relationship between the rate of energy dissipation and the speed and accuracy of adaptation (their equations (5) and (S18)). Their formulas resemble our equation (17). They infer their results from a continuum model of a three-state negative-feedback system, using the Fokker-Planck equation for the time evolution of the probability density and undertake the approximation by steepest descent on the adaptation error. The similarity in results despite the very different methods supports the analogy that has been drawn previously between adaptation and discrimination [3]. However, the work of [25] is limited to a three-state system, while our results apply to systems of arbitrary complexity. In [17], Sartori and Pigolotti determine the thermodynamic cost of copying a biopolymer. The copying process includes a discrimination mechanism that chooses between right and wrong monomers for incorporation in the polymer. They adopt a general finite-state Markov process, as in [35], and rely on the second law of thermodynamics to derive a bound for the total entropy production per wrongly incorporated monomer (their equation (3)). They identify three operating regimes, of which the third, called “error correction”, resembles that studied here and yields a formula which is similar to our equation (17). Their formula expresses a finite bound and involves only quantities related to discrimination of the wrong monomer, while our formula is valid asymptotically and compares right and wrong discriminations. The two studies can be seen as offering complementary approaches to the problem of algebraic complexity away from thermodynamic equilibrium: Sartori and Pigolotti appeal to the second law of thermodynamics while we exploit leading-order asymptotics as $x \to \infty$.

In summary, our work offers methods to rise above the algebraic complexity characteristic of nonequilibrium Markov processes and suggests that the quantity $\sigma \ln(\mu)/P$ may be significant for a broader context of cellular information processing that includes discrimination, adaptation, and other processes required for life. Because of their generality, the methods used here may be particularly useful for developing such a broader perspective.

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APPENDIX A: PROOF OF $P_k \lesssim P$

Suppose that $P_k \sim x^{\alpha_k} \ln(x)$. If $P_i$ is also in case 1 and $P_i \sim x^{\alpha_i} \ln(x)$, then $P_i/P_k \to c$, where $c = 0, 1, \infty$, depending on the relative values of $\alpha_i$ and $\alpha_k$. Similarly, if $P_j$ is in case 2 and $P_j \sim x^{\alpha_j}$, then $P_j/P_k \to c$, where $c \neq 0, \infty$, depending on the relative values of $\alpha_j$ and $\alpha_k$. By assumption, there are no other cases to consider (if $P_k$ were in case 3, we could not estimate $\lim_{x \to \infty} P_k/P_k$). Since $P_k$ is one of the summands in $P$, it follows that $P/P_k \to c$, where $1 \leq c \leq \infty$. Equivalently, $P_k/P \to c^{-1}$, where $0 \leq c^{-1} \leq 1$. In particular, $P_k \lesssim P$, as required.

APPENDIX B: PROOF OF THE ASYMPTOTIC RELATION

Suppose first that $1 \rightleftarrows n_C$ falls into case 1 in equation (15). Let $\alpha = \deg(\ell_{n \to 1} p_{n_C}^*)$ and $\beta = \deg(p_{1 \to n_C} p_1^*)$, so that $\alpha \neq \beta$. Then, $x^{\alpha} \ln(x) \lesssim x^{\max(\alpha, \beta)} \ln(x) \sim P(1 \rightleftarrows n_C)$. Since the product generation rate, $W$, appears additively, $\ell_{n \to 1} = W + U(x)$, for some allowable function $U$. It follows from equation (7) that $\alpha = \deg(\ell_{n \to 1}) + \deg(p_{n_C}^*) = \max(0, \deg(U)) + \deg(p_{n_C}^*) \geq \deg(W p_{n_C}^*) = \deg(\sigma)$. Hence, $\sigma \lesssim x^{\alpha}$. Furthermore, since equation (16) tells us that $\deg(\epsilon) < -1$, it follows from equation (14) that $\ln(\epsilon^{-1}) \sim \ln(x)$. Using equation (13), we deduce that

$$\sigma \ln(\epsilon^{-1}) \lesssim P(1 \rightleftarrows n_C).$$

If $1 \rightleftarrows n_C$ does not fall into case 1 in equation (15), then $\alpha = \beta$. Let us then consider $1 \rightleftarrows n_D$. According to equations (6) and (16), $\deg(p_{n_D}^*) < \deg(p_{n_C}^*) - 1$. Using equation (7) to combine this with equation (8.2), we see that $\deg(\ell_{n \to 1} p_{n_D}^*) < \deg(\ell_{n \to 1} p_{n_C}^*) = \alpha = \beta = \deg(\ell_{1 \to n_C} p_1^*)$. But now, by equation (8.1) and equation (7),

$$\deg(\ell_{1 \to n_C} p_1^*) = \deg(\ell_{1 \to n_D} p_1^*).$$

It follows that

$$\deg(\ell_{n \to 1} p_{n_D}^*) < \deg(\ell_{1 \to n_D} p_1^*),$$

so that $1 \rightleftarrows n_D$ falls into case 1 even though $1 \rightleftarrows n_C$ does not. Therefore, by equation (15), $P(1 \rightleftarrows n_D) \sim x^\gamma \ln(x)$, in which, because of equation (25), $\gamma = \deg(\ell_{1 \to n_D} p_1^*)$. But according to equation (24), $\gamma = \deg(\ell_{1 \to n_C} p_1^*) = \beta = \alpha$. Hence, by the same argument as above for $1 \rightleftarrows n_C$, we deduce that

$$\sigma \ln(\epsilon^{-1}) \lesssim P(1 \rightleftarrows n_D).$$

We can now appeal to the result in Appendix A to complete the proof.


[45] We note that Banerjee et al. imposed one additional constraint on their numerical values, following [3], by assuming that the correct and incorrect substrates used the same external chemical potential to break thermodynamic equilibrium; see equation (2) in [19]. While this is reasonable, we have followed Hopfield here and not made that assumption in our analysis and calculations. We also had to estimate two parameter values for the ribosome variants in Fig. 3(b), for which experimental data were not available, and did so by random sampling, as explained in Supplementary Table 1.