Balancing error and dissipation in computing

Paul M. Riechers, Alexander B. Boyd, Gregory W. Wimsatt, and James P. Crutchfield

Abstract: Modern digital electronics support remarkably reliable computing, especially given the challenge of controlling nanoscale logical components that interact in fluctuating environments. However, we demonstrate that the high-reliability limit is subject to a fundamental error–energy-efﬁciency tradeoff that arises from time-symmetric control: requiring a low probability of error causes energy consumption to diverge as the logarithm of the inverse error rate for nonreciprocal logical transitions. The reciprocity (self-invertibility) of a computation is a stricter condition for thermodynamic efﬁciency than logical reversibility (invertibility), the latter being the root of Landauer’s work bound on erasing information. Beyond engineered computation, the results identify a generic error–dissipation tradeoff in steady-state transformations of genetic information carried out by biological organisms. The lesson is that computational dissipation under time-symmetric control cannot reach, and is often far above, the Landauer limit. In this way, time-asymmetry becomes a design principle for thermodynamically efﬁcient computing.

I. INTRODUCTION

Tantalizingly, the thermodynamics of computation tells us that information processing can be achieved with zero energy dissipation if one has sufﬁcient control over a system’s microscopic degrees of freedom and can endure the quasistatic limit of inﬁnitely slow processing [1–5]. To be useful, though, computation must be performed in ﬁnite time. Unfortunately, this requires additional work and guarantees the investment is lost via dissipation. This state of affairs poses a grand challenge to thermodynamic computing: identify control protocols that reliably drive a system between memory states according to the computation’s input-output map. At an appropriate later time, the forces are ramped down (mirroring the ramp-up procedure) so that the computing system returns to its resting state, while new memories are again stored robustly in metastable conﬁgurations. Such implementations represent control protocols that are symmetric in time.

Time-symmetric protocols are common. Most notably, the primary signal controlling information processing in contemporary microprocessors is a time-symmetric clock-voltage signal—a several gigahertz square wave that orchestrates all transformations of the computer’s logic and memory components [12,13].

Time-symmetric protocols are common in other settings that seek reliable transformations. For example, recent explorations of synthetic molecular machines externally drive molecular rotors with sinusoidal (time-symmetric) electric ﬁelds [14]. And, autonomous synthetic molecular machines are known to reliably produce steady-state rotations embedded in a time-invariant nonequilibrium environment [15]—a special case of time-symmetric control. They achieve this through chemical catalysis, much as natural molecular machines operate in vivo. In vivo, reliable genetic transformations of DNA (and reliable directional motion of molecular motors) occur in environments with very low Reynolds number, where there are no inertial variables to freely exert a time-asymmetric inﬂuence [16,17]. More broadly, breaking time symmetry in Brownian motion requires signiﬁcant dissipation [18]. And so, in both the biological and engineered worlds, simple time-symmetric protocol design—turning on and then turning off an interaction—seems to offer an energetically efﬁcient and reliable way to implement change.

Published by the American Physical Society under the terms of the Creative Commons Attribution 4.0 International license. Further distribution of this work must maintain attribution to the author(s) and the published article’s title, journal citation, and DOI.

For example, a set of gate voltages may be applied or, by any other method, a potential energy landscape may be altered to temporarily couple and bias a set of initially independent metastable potential minima.
The goal, naturally enough, is to implement information processing in ways that require no more than the minimal work exertion set by Landauer’s logical-irreversibility bound [19]: \( W \gtrsim k_B T \ln 2 \). One might even hope that this work could then be recycled rather than dissipated. Indeed, as appreciated recently, changes in the nonequilibrium addition to free energy can be leveraged to implement logically irreversible computations in a thermodynamically reversible manner [2–5]. The vision is of a future of hyperefficient computing using orders-of-magnitude less energy than currently.

However, there is a wrinkle in this optimism. The following shows that all time-symmetric protocols for transforming metastable memories engender an irreducible trade-off between computational accuracy and energy efficiency. Specifically, in the limit of highly reliable computing with vanishingly small error probability \( \epsilon \), the minimal dissipation under time-symmetric controls diverges as \( -\ln \epsilon \) with a coefficient proportional to the computation’s nonreciprocity—the non-self-invertibility of its memory-state transitions.

That is, for reliable time-symmetric implementations of a deterministic computation \( C \), which maps each memory state \( m \mapsto C(m) \), we show that the minimal work above the change in local-equilibrium free energy is \( k_B T \ln(1/\epsilon) \) whenever \( C(C(m)) \neq m \). That is, work \( k_B T \ln(1/\epsilon) \) is required whenever the computation iterated twice does not return to the original memory state. Moreover, there is a correction when time-odd variables store memory that highlights the unique advantages of both magnetic and conformational memories.

In this way, nonreciprocity identifies a dominant cost of thermodynamic computing for the broad class of time-symmetric control protocols. While future work addresses more general thermodynamic implications of reliable computation, since time-symmetric control protocols are common and often unavoidable in key applications, they are the subject of our error–dissipation tradeoff analysis here.

Our first step revisits the basics of dissipation in thermodynamic systems and the cost of partial macroscopic knowledge, when the entire microstate cannot be observed or controlled. The dissipation costs are then recast to apply to thermodynamic computing generally and, in particular, to information storage and processing in metastable mesoscopic states. We then analyze the role of time symmetries in control protocols and give a thermodynamic accounting. Dissipation scaling with error level is derived and applied in the limit of highly reliable computing. This is then used to analyze dissipation in erasure, logic gates, and biological systems. We conclude, briefly comparing recent work on control restrictions and noting directions for further exploration. Reference [59] provides details underlying the theory.

## II. COMPUTATIONAL DISSIPATION

To start, consider any physical realization of a memory device with a system Hamiltonian \( \mathcal{H}_0 \) parametrized by control \( x \in \mathcal{X} \). For example, \( x \) could correspond to an applied electromagnetic field, a vector of quantities specifying a potential energy landscape, or a set of piston positions. The Hamiltonian \( \mathcal{H}_0 \) determines the system’s instantaneous energies \( \{E_i(s)\}_{i \in \mathcal{S}} \), where \( \mathcal{S} \) is the set of system microstates. We assume that the system is in contact with an effectively memoryless heat bath at temperature \( T \) that enables the system to relax to both local and global equilibria. The instantaneous stochastic microstate dynamics are completely determined by the system’s instantaneous Hamiltonian and the system’s interaction with the heat bath.

Work is performed by driving the system via a control protocol \( x_{0\to T} = x_0 \cdots x_T \), which is the trajectory of the control parameters from time 0 to \( T \). The work \( W \) measures the accumulated change in the energy of occupied microstates—energy that was supplied by the controller. In the following, we are especially interested in computations implemented by the control protocol and in the associated work cost.

### A. Background

Understanding the relationship between work costs and computations requires tracking the energetics of microstate trajectories \( s_{0\to T} = s_0 \cdots s_T \) of the system from time 0 to \( T \). Via the implied dynamics set by the time-dependent Hamiltonian \( \mathcal{H}_x \) and the coupling with the bath at temperature \( T \), the initial state \( s_0 \) and the control protocol \( x_{0\to T} \) determine the probability of microstate trajectories:

\[
\text{Pr}_{s_{0\to T}}(S_{0\to T} = s_{0\to T}|S_0 = s_0) \quad \equiv \text{Pr}(S_{0\to T} = s_{0\to T}|S_0 = s_0, X_{0\to T} = x_{0\to T}),
\]

where \( S_\tau \) is the random variable for the microstate at time \( \tau \) and \( S_{0\to T} \) is the random variable chain for the full state trajectory. The subscripted probability \( \text{Pr}_{s_{0\to T}}(\cdot) \) denotes the probability distribution induced by the driving protocol \( x_{0\to T} \). This is the same as the probability conditioned on the driving protocol, as described by Eq. (1).

Hamiltonian control implies microscopic reversibility of the instantaneous system dynamics. This means state trajectories that release energy into the heat bath \( (Q > 0) \) are exponentially more likely than the time-reversed microstate trajectory of the system (that absorb heat), if the protocol were run in reverse [20,21]. In the Markovian limit,\(^2\)

\[
\frac{\text{Pr}_{s_{0\to T}}(S_{0\to T} = s_{0\to T}|S_0 = s_0)}{\text{Pr}_{s_{0\to T}}(S_{0\to T} = \mathcal{H}(S_{0\to T})|S_0 = s_0)} = e^{\beta Q(s_{0\to T}, x_{0\to T})},
\]

where \( \beta = (k_B T)^{-1} \) and \( k_B \) is Boltzmann’s constant. \( \mathcal{H} \) denotes time-reversal, including not only reversal of the time ordering, but also conjugation of time-odd variables such as momentum and spin: \( s_0 \cdots s_T \mapsto s_T^\dagger \cdots s_0^\dagger \). For example, if the microstate \( s = (\vec{q}, \vec{p}) \) is a collection of spatial \( \vec{q} \) and momentum \( \vec{p} \) degrees of freedom, then the conjugation simply flips all of the momentum degrees of freedom \( s^\dagger = (\vec{q}, -\vec{p}) \).

Thermodynamic irreversibility is quantified by entropy production \( \Sigma \) that, by the second law, is nonnegative on average: \( \langle \Sigma \rangle \geq 0 \). In our setting, \( \Sigma \) is simply the net change in the component entropies of both the system and the bath.

More specifically, entropy production decomposes into the change in heat-bath entropy \( Q(s_{0\to T}, x_{0\to T})/T \) beyond any

---

\(^2\)In the more general non-Markovian case, Eq. (2) becomes an inequality that nevertheless still upholds our main results bounding computational dissipation.
compensating reduction in the system’s microstate entropy. The system’s instantaneous microstate entropy is given by the nonequilibrium surprisal \(-k_B \ln \mu_i(s_i)\), where \(\mu_i(s_i) = Pr_{\text{sys}}(S_i = s_i)\) is the current microstate’s probability given the entire history of preparation and driving [20,22,23]. Accordingly, the entropy production is

\[ \Sigma = \frac{1}{T} Q(s_{0:T}, x_{0:T}) + \Delta(-k_B \ln \mu(s_i)), \]

where \(\Delta\) indicates the change from time 0 to \(T\). Specifically, \(\mu_0\) is the distribution over microstates given the system’s initial preparation and \(\mu\) is the time-evolved version of \(\mu_0\) under the influence of the driving \(x_{0:T}\).

This results in the trial-specific entropy production [20,21]:

\[
\Sigma = \frac{1}{T} Q(s_{0:T}, x_{0:T}) + k_B \ln \frac{\mu_0(s_0)}{\mu(s_T)} = k_B \ln \left( \frac{Pr_{s_0}(S_{0:T} = s_0 \mid s_0 = s_0) \mu_0(s_0)}{Pr_{\text{sys}}(S_{0:T} = s_T \mid s_0 \sim \mu_0) \mu(s_T)} \right)
\]

where \(\mu_0\) is the distribution such that \(\mu_0(s) = \mu(s')\). The condition \(S_0 \sim \mu\) means the random variable \(S_0\) is distributed according to \(\mu\), as in Ref. [24]. We used the fact that \(\mu_i(s) = \mu_i(s')\) in going from the second to the third line.

When all influences driving the system away from equilibrium are controlled changes to the system Hamiltonian, the total entropy production is proportional to the dissipated work \(W_{\text{diss}}\). This is the accumulation of lost opportunities to extract work:

\[ W_{\text{diss}} = T \Sigma. \]

The entropy production and dissipated work both measure a given computation’s efficiency, as they both quantify the degree to which more work was done than necessary by an unrestricted controller.

The expected dissipated work then quantifies the average amount of work done beyond the change in nonequilibrium free energy [25]:

\[ \langle W_{\text{diss}} \rangle = \langle W \rangle - \Delta F. \]

This is the average work that has been irreversibly lost, since the nonequilibrium free energy \(F\) is the expected amount of energy that possibly could be extracted as work [4].

Calculating the dissipated work from these two quantities seems to require explicit knowledge of the Hamiltonian, since the work is the integrated change in the energy due to changes in control:

\[ \langle W \rangle = \sum_x \int_0^T dt \mu_i(s) \frac{\partial E_x(s)}{\partial x_t} dx_t. \]

\[ 3 \text{Hence, } \mu_i(s_i) \text{ is the expected probability of being in the actual microstate } s_i \text{ over many trials where the system is prepared and driven in exactly the same way. The fact that this distribution is not } \delta \text{-distributed is due to the stochasticity induced by the interaction with the heat bath (and any other uncontrollable degrees of freedom).} \]

and the nonequilibrium free energy is the difference between the average energy and the microstate entropy [4]:

\[ \mathcal{F}(t) = \langle H_S \rangle - k_B T H(S_i), \]

where \(H(Z) = -\sum_s \Pr(Z = s) \ln \Pr(Z = s)\) is the Shannon entropy of the driven system in nats.\(^4\)

However, due to Eq. (3), we can calculate the dissipated work for a control protocol using only the probability of forward trajectories under forward driving \(\rho(S_{0:T} = s_{0:T}) = Pr_{\text{sys}}(S_{0:T} = s_{0:T} | S_0 \sim \mu_0)\) and reverse microstate trajectories under reverse driving \(\rho^R(S_{0:T} = s_{0:T}) = Pr_{\text{sys}}(S_{0:T} \sim \mathcal{H}(S_{0:T}) | S_0 \sim \mu_i)\). Averaging over forward trajectories produces a relative entropy:

\[
\beta \langle W_{\text{diss}} \rangle = \int_0^T dt \mu_i(s) \frac{\partial E_x(s)}{\partial x_t} dx_t.
\]

where \(D_{KL} [\cdot | | \cdot]\) is the Kullback–Leibler divergence.

Equation (5) appears superficially similar to many previous related results, including Refs. [26–30]. However, these previous results all required that the system starts in a steady-state distribution (either equilibrium or a nonequilibrium steady state), which precludes the storage of general initial memories. Equation (5) overcomes that previous limitation via its dependence on \(\mu_0\) and \(\mu_i\)—a crucial generalization that will allow us to address the thermodynamics of general memory transformations.

By expressing the dissipated work and entropy production in terms of the relative entropy between forward and reverse trajectories, Eq. (5) provides a method to reconstruct a system’s thermodynamics without explicit knowledge of the underlying Hamiltonian mechanics. This only requires knowledge of trajectory probabilities under forward and reverse driving and these can be determined experimentally. However, practical considerations can stand in the way: Often direct observations of a thermodynamic system’s underlying microstate trajectories are not available.

Fortunately, via the information processing inequality [24], this powerful relation leads to bounds on dissipated work based on observed trajectories, which contain only partial details of the full microstate trajectory. In particular, any observable coarse-graining of the microstate trajectory yields a lower bound on the dissipation [27,28,31]. This will allow us to infer new lower bounds on computational dissipation from only partial knowledge related to the logical dynamics of the computation itself.

---

\(^4\)In point of fact, knowledge of the Hamiltonian is not strictly necessary if one instead calculates \(W_{\text{diss}} = W_{\text{ex}} - \Delta F_{\text{add}}\), where \(W_{\text{ex}} = W - \Delta F_{\text{eq}}\) and \(F_{\text{add}} = k_B T D_{KL} [\mu | | \pi_s]\). With knowledge (or observation) of the equilibrium distribution \(\pi_s\), associated with each control setting \(s\), one can leverage the Boltzmann relationship \(\pi_s(s) = e^{-\beta E_x(s)} \pi^0_s\) to calculate: \(\beta \langle W_{\text{ex}} \rangle = \sum_x \int_0^T dt \mu_i(s) \frac{\partial E_x(s)}{\partial x_t} \). Still, Eq. (5) presents an even more tempting opportunity for calculating \(W_{\text{diss}}\) directly from observed trajectory probabilities.
B. Thermodynamic computing

To be usefully manipulated, memories must be physically addressable. Accordingly, we define the set $\mathcal{M}$ of memory states as a partition of the set $\mathcal{S}$ of microstates, such that each memory state $m \in \mathcal{M}$ is a subset of the microstates: $m \subset \mathcal{S}$ and $\bigcup m = \mathcal{S}$. In this way, the memory-state random variable $\mathcal{M}_t$ at time $t$ is determined by the microstate $\mathcal{S}_t$. Moreover, for memory states to robustly hold memories between computations, we assume that partitioning is such that each memory state corresponds to a state-space region that is effectively autonomous when no computation is being performed, resulting in metastability of the memory. Figure 1 shows a spatial system that has been partitioned into left (L) and right (R) memory states, each of which corresponds to a local minimum in the energy landscape.

Partitioning microstates into metastable memory states $\mathcal{M}$ introduces an observational channel for monitoring computation. The task of thermodynamic computation is to map an initial memory state $m$ to a final memory state $m'$ according to a given conditional probability distribution:

$$p(m \to m') = \Pr(\mathcal{M}_t = m' | \mathcal{M}_0 = m).$$

For instance, as shown in Fig. 2 for Fig. 1’s system, erasure is composed of two desired transitions with high probabilities—$p(L \to L) \approx 1$ and $p(R \to L) \approx 1$—between the two metastable memory states. (The latter are labeled L and R according to the common use of left and right minima in a double-well potential landscape to store a bit [32].) A computation occurs due to a particular stochastic trajectory, such as the red path shown in Fig. 2. Naturally, the statistics of the memory state transition probability $p(L \to L)$, say, come from the ensemble of trajectories.

Since stochasticity is the rule in thermodynamic computations, all of our results throughout Sec. II are fully general for any level of stochasticity. However, since the nearly deterministic limit of reliable computation is common and practically applicable, Sec. III will subsequently address the drastic consequences of low error. For now, we proceed in the general probabilistic regime.

To lower-bound the thermodynamic dissipation associated with computation, we consider the coarse graining $s_0: \tau \mapsto (m, m')$, where $s_0 \in m$ and $s_\tau \in m'$. This maps the detailed microstate trajectory to the logical transformation it induces between initial and final memory states. For a particular realization $\mathcal{M}_0 = m$ and $\mathcal{M}_\tau = m'$, the time reversal of the initial and final memory states is: $\mathcal{M}(m, m') = (m', m)$, where the conjugate $m' = \{ s' : s \in m \}$. This leads to observable

\[\text{FIG. 1. Bistable memory element: a cross-section (at constant momentum } \mathcal{P}_0 \text{) of both global and local equilibrium. Initial system state } s_0 = (q_0, \mathcal{P}_0) \text{ (red star) lies in a continuum. The energy landscape (black curve) has two symmetric minima that effectively partition the space into robust left (blue) and right (yellow) spatial memory states. The stable equilibrium distribution (green dashed curve) is uniformly distributed over the left and right states. The restriction of the equilibrium distribution to the left well (blue curve) is metastable, only locally in equilibrium.}\]

\[\text{FIG. 2. (Top) Time-symmetric control implementing reliable information processing: erasure to L. (Middle) Controlled potential } E_n \text{ at several times during the erasure protocol. (Bottom) Representative behavior via 1000 microstate trajectories in gray which sketches a profile of } \mu. \text{ A particular microstate trajectory } s_0: \tau \text{ is highlighted in red. (Note that } q_0: \tau, \text{ shown explicitly, implies } \mathcal{P}_0: \tau \text{ by differentiation and so also implies } s_0: \tau.\]

\[\text{5During a computation, though, driving couples memory states to instantiate nontrivial computing.}\]
and the reversal probabilities:
\[ \rho^R((\mathcal{M}_0, \mathcal{M}_\tau) = (m', m)) \]
\[ = \sum_{s_\tau \in \mathcal{M}} \delta_{m' \in \mathcal{M}^e} \rho(\mathcal{S}_{0,\tau} = s_{0,\tau}) \]
\[ = \Pr_{s_{0,\tau}}(\mathcal{M}_0 = m, \mathcal{M}_\tau = m'|\mathcal{S}_0 \sim \mu_0) \]  \hspace{1cm} (6)

where \( \mu_\tau \) is the same preparation-and-driving-induced probability measure as previously introduced, such that \( \mu_\tau(m) = \Pr(\mathcal{M}_\tau = m|\mathcal{S}_0 \sim \mu_\tau) = \sum_{x \in \mathcal{X}} \mu_\tau(x) \), and

\[ \Delta H(M_\tau) = \sum_{m \in \mathcal{M}} \mu_\tau(m) \ln \mu_\tau(0)(m) - \mu_\tau(m) \ln \mu_\tau(m) \]  \hspace{1cm} (7)

is the change in Shannon entropy (in nats) of the coarse-grained memory states.

Note that Eq. (10) defined
\[ d(m, m') = \Pr_{x_{0,\tau}} (M_\tau = m'|\mathcal{S}_0 \sim \mu_0) \]
\[ \times \ln \left( \frac{\Pr_{x_{0,\tau}} (M_\tau = m'|\mathcal{S}_0 \sim \mu_0)}{\Pr (M_\tau = m'|\mathcal{S}_0 \sim \mu_0^{(m')})} \right) \]  \hspace{1cm} (11)

where \( \mu_0^{(m')} = \delta_{s \in \mathcal{M}^e} \mu_\tau(s) / \mu_\tau(m) \) is the renormalized microstate distribution \( \mu_\tau \) restricted to memory \( m \)'s microstates. This compares (i) the probability of transitioning from memory state \( m \) to \( m' \) to (ii) the probability of returning to \( m \) upon subsequent momentum-conjugation of the microstate distribution in \( m' \) and reversal of the control protocol. One concludes that dissipation is due to statistical irreversibility. At first sight, Eqs. (10) and (11) do not appear to significantly simplify the problem of inferring dissipation from partial knowledge. However, the practical constraints of reliable computation greatly simplify \( d(m, m') \), as we will see shortly.

To emphasize as either Eqs. (9) or (10), the bound applies when the conjugated \( m' \equiv \{ s' : s \in m \} \) of a memory state \( m \) is an element of the memory partition, \( m' \in \mathcal{M} \), for all \( m \). (However, see Ref. [59] part A for the more general case.) This occurs, for example, when spatial cells store a memory that is indifferent to rapidly fluctuating momentum degrees of freedom, so that \( m \equiv m' \). A similar simplification occurs for magnetic memory where \( s \) and \( s' \) are always in different memory states, such that \( m' \in \mathcal{M} \) although \( m \neq m' \). These two types of memory, illustrated in Fig. 3, lead to notable physical consequences; ones that we explore in a sequel.

Equation (10) is close to yielding a bound on entropy production divorced from knowledge of microstate dynamics. \( \Pr_{x_{0,\tau}}(\mathcal{M}_\tau = m'|\mathcal{S}_0 \sim \mu_0^{(m)}) \), for instance, can be recovered from observations, given a system’s initialization \( \mu_0^{(m)} \) and the driving \( x_{0,\tau} \). However, for \( \Pr_{x_{0,\tau}}(\mathcal{M}_\tau = m'|\mathcal{S}_0 \sim \mu_0^{(m)}) \), it is unclear how to experimentally prepare the distribution \( \mu_0^{(m)} \) in the presence of time-odd variables, like momentum. Fortunately, as we now show, this obstacle can be removed for thermodynamic computations using metastable memories. Ignoring the underlying microstate dynamics and, instead, only tracking the logical dynamics, we bound the work production while computing.

C. Metastable processing

Further simplifying Eq. (10)’s bound on dissipated work turns on recognizing a common property of information processing systems: They hold memory states as metastable nonequilibrium distributions. This has profound thermodynamic implications.

The implied timescale separation requires that relaxing to the local-equilibrium distribution \( \pi_0^{(m)} \) within each memory region is fast, while relaxing to the global equilibrium distribution \( \pi_\ast \) is much slower than the timescale of (and between) a system’s computational steps. On the one hand, metastable distributions are those that correspond (up to some approximation) to a normalized weighted sum of local-equilibrium distributions. On the other, the global equilibrium distribution \( \pi_\ast \) is the canonical one: \( \pi_\ast(s) = e^{-\beta E_\ast(s)} / \sum_{s \in \mathcal{S}} e^{-\beta E_\ast(s)} \). The system approaches it in the limit of infinite time, if the control setting \( x \) is held fixed. In this, the local-equilibrium distributions \( \pi_0^{(m)}(s) = \delta_{s \in m} \pi_\ast(s) / \sum_{s \in \mathcal{S} \setminus m} \pi_\ast(s) \) are the canonical distributions, associated with each nearly autonomous region of state space, that local densities approach much more quickly due to timescale separation [25].

If metastable memories are stored robustly between computations, then the memory-state distribution \( \mu_\tau(m) \) at times \( t = 0 \) and \( t = \tau \) contain almost all the information about the microstate distributions at the start and end times; or,

---

6Metastable states are indeed stable on the timescale between computations if each local-equilibrium distribution \( \pi_0^{(m)} \) is an approximately stationary distribution of the evolution induced under the fixed potential energy landscape \( E_\ast(s) \) and the interaction with the thermal bath. This can be achieved if the energy-barriers between each memory region are much larger than the thermal energy.
at least, at the beginning and shortly after the end time. Indeed, the local distributions associated with each memory state are then nearly local-equilibrium distributions. If, to the contrary, the ending microstate distribution $\mu_x(s)$ is not yet metastable, then it quickly relaxes to the metastable distribution $\mu_{x,\text{fr}}(s) \approx \sum_{m \in M} \mu_x(m) \pi_{x,m}(s)$. This results in yet more dissipation.

To include all dissipation associated with a computation, we extend a given protocol’s start and end times to include most of the post-computation relaxation to metastability. Assuming a metastable distribution at the beginning and end of the computation means $\mu_x^{(0)} \approx \pi_x^{(m)}$ and $\mu_x^{(m)} \approx \pi_x^{(m)}$ and the full microstate distributions are then $\mu_0 \approx \sum_{m \in M} \mu_0(m) \pi_x^{(m)}$ and $\mu_\tau \approx \sum_{m \in M} \mu_\tau(m) \pi_x^{(m)}$, respectively. Thus the beginning and ending microstate distributions are almost entirely determined by the distribution over memory states.

Initial and final memory-system metastability means that the empirically observable memory transitions are always implicitly conditioned on metastability:

$$\Pr(M_\tau = m^0 | M_0 = m)$$

Similarly, if we operate the control protocol in reverse, starting metastably under the influence of $x'_t$, the observed memory-transition probabilities are

$$\Pr(M_\tau = m^0 | M_0 = m^t)$$

Note, for comparison with Eq. (11), that

$$\pi_x^{(s)}(s) = \pi_{x',t}(s).$$

And so, we have

$$\pi_{x,t}^{(m')} = \pi_{x',t}^{(m')} = \mu_x^{(m')}.$$

This allows us to remove all dependence on microstates from the work bound and estimate dissipation purely from observed memory-state trajectories. In particular, Eq. (11) simplifies to

$$d(m, m') = \Pr(M_\tau = m^0 | M_0 = m)$$

with

$$\Pr(M_\tau = m^0 | M_0 = m^t)$$

This demonstrates how restricting to time-symmetric control leads to unavoidable dissipation that depends explicitly on a computation’s logic.

D. Time symmetric driving

Let us now consider the consequences for reliable computing with time-symmetric protocols—those for which $x_{0,\tau} = \mathbf{R}(x_{0,\tau})$—as shown in the upper panel of Fig. 2’s computation. In these cases, Eq. (13) simplifies considerably:

$$\beta(W_{\text{diss}}) \geq \Delta H(M_t) + \sum_{m,m'} \Pr(M_t = m', M_0 = m)$$

$$\times \ln \frac{\Pr_{x_{0,\tau}}(M_t = m^0 | M_0 = m)}{\Pr_{x_{0,\tau}}(M_t = m^0 | M_0 = m^t)}.$$
which is achieved in the limit of zero dissipation. Here, we recalled that for metastable distributions the nonequilibrium free energy decomposes as
\[
F = F^{(i)}_{x} - k_B T H(M_i),
\]
where \( F^{(i)}_{x} = -k_B T \ln(\sum_{m} e^{-\beta E_i(s)}) \) is the local-equilibrium free energy associated with the memory \( m \) [4,5].

\[
(W)^{\text{sym}}_{\min} = \Delta \left[ F^{(i)}_{x} \right] + (W)_{\text{Landauer}}^{\text{sym}} + (W)_{\text{dir}}^{\text{sym}} + \mu_0(m)p(m \rightarrow m') \ln \frac{p(m \rightarrow m')}{p(m' \rightarrow m')}. \tag{18}
\]

The consequences are immediate. Computing with time-symmetric protocols requires additional work above and beyond the Landauer bound. Moreover, beside the reversible Landauer contribution and the change in local free energy, all of this work must be irreversibly dissipated since it is not stored as non-equilibrium free energy. \( (W)_{\text{dir}}^{\text{sym}} > 0 \), quantified precisely by Eq. (15), contributes to wasted heat in any computer. It adds on top of other beyond-Landauer work bounds including (i) the cost of modular computations, where global correlations are dissipated due to localized control [5,33], and (ii) the cost of neglecting the local statistics of the manipulated memory [5,34].

E. (Non)Reciprocity

We argued that it is common—in practice and, occasionally, out of necessity—for a computation to be implemented by transforming metastable memories with time-symmetric driving. We are now in a position to plainly state the thermodynamic consequences of this type of computation.

Generically, a physical computation produces and is characterized by the set of memory transition probabilities: \( p(m \rightarrow m') \in \mathcal{M} \). Drawing out the consequences of Eq. (18), we see that the work required for such a computation, beyond the change in local free energy (which is often constructed to be zero), is determined by the nonreciprocity of memory transitions. We say that a memory transition is reciprocated if
\[
p(m \rightarrow m') = p(m' \rightarrow m'). \tag{19}
\]
Nonreciprocity \( \Psi \) quantifies the deviation from this
\[
\Psi(m \rightarrow m') \equiv \ln \frac{p(m \rightarrow m')}{p(m' \rightarrow m')}. \tag{20}
\]

An important related but often overlooked result emerges. Even for logically irreversible computations, the minimal work required by this generalized Landauer bound is reversible: it corresponds to a stored change in nonequilibrium free energy that can be retrieved. Thus, ultimately no work needs to be dissipated. We could have \( (W)_{\text{dir}}^{\text{sym}} = 0 \) in the case of unrestricted control. And, this alerts us to additional opportunities for optimizing implementations.

Appealing to Eq. (4) again, we see that the minimum total work under time-symmetric driving must be
\[
(W)_{\min}^{\text{sym}}(m \rightarrow m') = F_{m}^{(m')} - F_{m}^{(m')} + k_B T \Psi(m \rightarrow m'). \tag{23}
\]

Notably, it vanishes when Eq. (19) is satisfied. Keep in mind that \( m' \) and \( m'' \) are themselves valid memory states, so non-reciprocity compares the probability of two different memory transitions.

Reference [59], part B, establishes the transition-specific version of Eq. (18). To summarize, we first derive a useful transition-specific fluctuation theorem:
\[
\langle e^{-\beta W} \rangle_{\Psi_{x}}(S_{0}, s_{0}, m, \pi) = e^{-\beta \Delta F_{x}^{(m)}} \frac{\text{Pr}_{\Psi_{x}}(s_{0}, m, \pi)}{\text{Pr}_{\Psi_{x}}(s_{0}, m', \pi)} \tag{21}
\]
which applies to systems that start in local equilibrium. The appearance of the local-equilibrium free energies makes this a useful generalization of related past results [35]. By Jensen’s inequality, and assuming time-symmetric control, this then implies that the minimal average work required of a memory transition is
\[
(W)_{\min}^{\text{sym}}(m \rightarrow m') = F_{m}^{(m')} - F_{m}^{(m')} + k_B T \Psi(m \rightarrow m'). \tag{22}
\]

In the common scenario where all local-equilibrium free energies are constructed to be equal, this reduces to
\[
(W)_{\min}^{\text{sym}}(m \rightarrow m') = k_B T \Psi(m \rightarrow m'). \tag{23}
\]

In other words, nonreciprocity implies work. This can also be seen from Trajectory Class Fluctuation Theorems if one restricts to trajectories that start and end in the specified memory states [36]. Not all work is bad, though. When \( W_{\min}^{\text{sym}}(m \rightarrow m') \) is negative, some work can be harvested during the transition. In fact, such work harvesting is necessary to achieve minimal dissipation.

Equation (23) implies that strictly reciprocal computations, for which Eq. (19) is satisfied for all \( m, m' \in \mathcal{M} \), can be implemented with a time-symmetric protocol with no expended work at all. The identity map is a trivial example. Two-cycles, like a bit swap, can also be implemented for free if using a time-symmetric memory substrate (for which \( m' = m \) for all \( m \in \mathcal{M} \)). However, most computations require nonreciprocated memory transitions and, so, require work. For example, a non-self-inverting permutation—like a three-cycle—is nonreciprocal for time-symmetric memories.

---

\(^{a}\)Common computational substrates have \( \Delta F_{m}^{(m')} = 0 \), since engineered memories typically have equal local-equilibrium energies. The change in local free energy is nevertheless likely important in the manipulation of biological memories.
Despite being logically reversible, this computation requires work.

This points out the important difference between nonreciprocity and logical irreversibility—the latter of which is typically deemed responsible for the Landauer work cost of computation. Reciprocity is stricter than logical reversibility, with nonreciprocated permutation being a prime example. However, nonreciprocity also has more interesting physical nuance, since it involves the time-reversal symmetries of the memory elements used in the computation.

While the Landauer work cost corresponds to average state-space compression, nonreciprocity corresponds to a localized imbalance of memory currents. Even in the case of heterogeneous local free energies [i.e., Eq. (22)], the minimal dissipation required of time-symmetrically driven computations is the difference between the two:

\[ (W_{\text{diss}})_{\text{min}}^{\text{t-sym}} = k_B T \langle \Psi(A_0 \rightarrow A_1) \rangle - (W_{\text{Landauer}})_{\text{min}}. \]  

(24)

The expected nonreciprocity \( \langle \Psi(A_0 \rightarrow A_1) \rangle \) weights how often a nonreciprocated transition is exercised. This uses the weighted nonreciprocity \( d(m \rightarrow m') \):

\[ d(m \rightarrow m') = p(m \rightarrow m') \Psi(m \rightarrow m'), \]

(25)

which appeared previously in Eq. (14). The nonreciprocity weight is critical to the following analysis. Though \( \Psi(m \rightarrow m') = -\Psi(m' \rightarrow m) \), weighted nonreciprocity can strongly break the symmetry.

Consider a typical computation that enforces logical transitions: some transitions should happen with certainty while others should be forbidden. In particular, consider a nonreciprocated \( m \mapsto m' \) logical transition: \( m \mapsto m' \) should be strongly enforced \( (p(m \rightarrow m') \approx 1) \), while \( m' \not\mapsto m \) should be strongly forbidden \( (p(m' \rightarrow m) \approx 1) \), \( 0 < \epsilon < 1 \) otherwise. Then \( d(m \rightarrow m') \) diverges as \( \ln(1/\epsilon) \) while \( d(m' \rightarrow m) \) vanishes. In short, divergent work is required to implement the desired nonreciprocated \( m \mapsto m' \) logical transition.

A deterministic computation \( C \) entails a deterministic logical transformation \( m \mapsto C(m) \) satisfying the reciprocity condition of Eq. (19) only when

\[ C(C(m)) = m. \]

(26)

If Eq. (26) is not satisfied—i.e., if \( C(m) = m' \) but \( C(m') \neq m' \)—then the logical transformation appears to require infinite work and infinite dissipation since, from Eq. (18), it appears to require work of \( k_B T \ln \frac{1}{\epsilon} \). In practice, since dissipation is bounded, such logical transformations can only be approximated. Reliable logic (with a probability of error less than some small \( \epsilon \)) requires significant dissipation of at least \( k_B T \ln(1/\epsilon) \) for each violation of Eq. (26). Section III develops the consequences of this new relationship between reliability and dissipation. That then allows us to explore how the two can be balanced in several examples.

**F. Time symmetric memory**

The following explores the implications of these results for time-reversal-invariant memories—\( m = m' \)—in which information is stored via time-symmetric variables, such as spatial location or physical conformation. In these cases, Eq. (15) simplifies to

\[ \beta \langle W_{\text{diss}} \rangle_{\text{min}}^{\text{t-sym}} = \Delta H(A_1) + \sum_{m,m'} \mu_0(m)p(m \rightarrow m') \times \ln \frac{p(m \rightarrow m')}{p(m' \rightarrow m)}. \]

(27)

As before, minimal dissipation depends directly on the nonreciprocity between memory transitions, but now we have the simplification that

\[ \Psi(m \rightarrow m') = \ln \frac{p(m \rightarrow m')}{p(m' \rightarrow m)}. \]

And here, reciprocity has an especially straightforward interpretation. If \( m \) maps to \( m' \), then \( m' \) should map back to \( m \) with the same probability. Otherwise work is required and, if it is above the Landauer bound, it must be dissipated.

What is reciprocity in the limit of deterministic computation? For time-symmetric memory, it acts somewhat like logical invertibility. However, it is stricter. Rather, it is logical self-invertibility. Logical noninvertibility, in contrast, gives rise to state-space compression—the origin of the Landauer bound. Indeed, circumventing the latter led early researchers to investigate reversible computing—computing with invertible logical transformations [1]. Logically reversible computing requires no work to operate. However, it is now understood that even logically irreversible computing can be accomplished with zero dissipation, despite requiring some recyclable work [2–4].

The next section draws out the consequences of nonreciprocity in the limit of nearly deterministic computation. In contrast to the Landauer cost of logical irreversibility—that saturates to some small value on the order of the thermal energy in the limit of zero error—the thermodynamic cost of nonreciprocity diverges in the limit of zero error, with no chance of recovering the input energy. It demands an accounting.

**III. ALMOST-DETERMINISTIC COMPUTING**

At this point, we showed that the transition probabilities entailed by a desired logical transformation set a lower bound on the work required to instantiate it physically. We noted that this gives a much stronger bound than that due to Landauer, which only depends on the relative state-space volume supporting memory before and after a computation. More specifically, when time-symmetric control implements a transformation of metastable memories, the minimal work investment is proportional to the nonreciprocity (non-self-invertibility) of memory transitions. Furthermore, we discovered that the time-reversal symmetries of the memory elements themselves can substantially alter the minimal thermodynamic costs.

Drawing out the practical consequences, the following shows that computing with minimum work depends directly on a computation’s error rate \( \epsilon \). Most concretely, time-symmetry-induced dissipation diverges as \( \ln(1/\epsilon) \) for almost-deterministic computation, where metastable memories are transformed at low error rate \( \epsilon \rightarrow 0 \).
Consider the physical memory apparatus of a computer that approximately implements a deterministic computation $C: \mathcal{M} \rightarrow \mathcal{M}$, mapping memory states to memory states. Then, the memory-state transition probabilities appearing in Eq. (15) are strongly constrained by the computation’s desired reliability. That is, the probability that the nonequilibrium thermodynamic information processing takes a memory state $m$ to anywhere besides $C(m)$ must be no greater than the error tolerance $\epsilon$, with $0 < \epsilon \ll 1$. Reliability requires choosing a time-symmetric drive protocol $x_{0r}$ which guarantees that $P_{x_{0r}}(\mathcal{M}_1 = m' | \mathcal{M}_0 = m) \leq \epsilon$ for all $m, m' \in \mathcal{M}$ such that $m' \neq C(m)$. The thermodynamic implications for a reliable computation then follow from Eq. (15) with

$$p(m \rightarrow m') \geq 1 - \epsilon \quad \text{if} \quad C(m) = m' \quad \text{and} \quad C(m) \neq m''.$$ 

Evaluating Eq. (15) requires addressing four cases, depending on whether $C(m) = m'$ or $C(m) \neq m'$ and on whether $C(m') = m'$ or $C(m') \neq m'$. Any given implementation results in an actual probability of error for each of the intended transitions: $\epsilon_m = 1 - p(m \rightarrow C(m))$. We adopt the design constraint that $\epsilon_m \leq \epsilon$ for all possible initial memories $m$. Then, given an implementation $x_{0r}$, the probability of an accidental memory transition is $\epsilon_{m \rightarrow m'} = p(m \rightarrow m')$ for $m', m' \neq C(m)$. Since $\sum_{m'} \epsilon_{m \rightarrow m'} = \sum_{m'} \epsilon_m = \epsilon$, we must have that $0 \leq \epsilon_{m \rightarrow m'} \leq \epsilon_m \leq \epsilon$.

To simplify, we temporarily restrict to the case of time-reversible memories, where $m = m'$. The four cases then depend on whether $C(m) = m'$ or $C(m) \neq m'$, and on whether $C(m') = m$ or $C(m') \neq m$. Later, we quote the general result that does not require $m = m'$.

Reference [59], part C, uses the error constraints to evaluate the weighted nonreciprocity:

$$d(m, m') = p(m \rightarrow m') \ln \frac{p(m \rightarrow m')}{p(m' \rightarrow m)},$$

for the four possible cases, finding

1. $C(m) = m'; C(m') = m$: 
   $$-\epsilon \leq d^{(1)}(m, m') \leq \epsilon + \frac{1}{2} \epsilon^2 + \mathcal{O}(\epsilon^3).$$

2. $C(m) = m'; C(m') \neq m$: 
   $$\ln(\epsilon^{-1}) \approx d^{(2)}(m, m') \approx \ln(\epsilon_{m \rightarrow m}'^{-1}).$$

3. $C(m) \neq m'; C(m') = m$: 
   $$\epsilon_{m \rightarrow m} \ln(\epsilon_{m \rightarrow m'}) < d^{(3)}(m, m') < 0.$$

4. $C(m) \neq m'; C(m') \neq m$: 
   $$-\epsilon / \epsilon \leq d^{(4)}(m, m') = \epsilon_{m \rightarrow m} \ln(\epsilon_{m \rightarrow m'}).$$

In the high-reliability limit $\epsilon \rightarrow 0$, most $d^{(i)}(m, m')$ terms are of order $\epsilon$ and so tend to zero. That is, except for $d^{(2)}(m, m')$ in case 2 whose contribution to the dissipation diverges as $\epsilon \rightarrow 0$ since $d^{(2)}(m, m') \approx \ln(\epsilon^{-1})$.

Case 4 is somewhat more delicate than cases 1 and 3, due to the ratio of errors. When $\epsilon_{m \rightarrow m} \geq \epsilon_{m' \rightarrow m'}$, then $-\epsilon / \epsilon \leq d^{(4)}(m, m') \leq 0$, so that $d^{(4)}(m, m')$ vanishes as $\epsilon \rightarrow 0$. However, when $\epsilon_{m \rightarrow m} > \epsilon_{m' \rightarrow m'}$, there is a chance for slightly more dissipation. Nevertheless, $d^{(2)}(m, m')$ is still only on the order of $\epsilon$, so long as the relative error rates $\epsilon_{m \rightarrow m'}$ and $\epsilon_{m' \rightarrow m}$ are within several orders of magnitude of each other. If $\epsilon_{m \rightarrow m'} \gg \epsilon_{m' \rightarrow m}$, then extra dissipation will be incurred due to the stringent reliability of the forbidden $m \rightarrow m'$ transition, beyond the design constraint.

The upper bound in case 2 also implies extra dissipation when $\epsilon_{m' \rightarrow m}$ is more reliable than the reliability design constraint. And, this extra dissipation need not be small. Generically, though, for a given error tolerance, we expect that minimal dissipation can be achieved by allowing all error rates to be as close to uniform as possible, meeting but not significantly exceeding the overall reliability constraint. Still, due to case 2, even the minimal dissipation diverges with increasing reliability.

Thus, case 2 provides the only contribution of the $d(m, m')$ terms to Eq. (10) in the low-error limit. Since the target computation is deterministic—$C$ maps each memory state $m$ to one memory state $C(m)$—there can be at most one contribution of $d^{(2)}(m, m')$ for each $m$, coming from $m' = C(m)$, and only if $C(m') = m$. The total contribution from each $m$ is

$$[C(C(m)) \neq m] \ln(\epsilon^{-1}_{C(m) \rightarrow m}),$$

where $[\ ]$ is the Iverson bracket, which returns 1 when its argument is true and 0 otherwise. In this case, $[C(m) \neq m] = 1 - \delta_{m, C(m)}$.

Applying the low-$\epsilon$ contributions to Eq. (14) yields an approximate bound for the time-symmetric work. If the memories have the same local-equilibrium free energy, the bound is

$$\beta (W)_{\text{approx}} \approx \sum_{m \in \mathcal{M}} \mu_0(m) [C(C(m)) \neq m] \ln(\epsilon^{-1}_{C(m) \rightarrow m})$$

$$\geq \ln(\epsilon^{-1}) \sum_{m \in \mathcal{M}} \mu_0(m) [C(C(m)) \neq m]$$

$$= \beta (W)_{\text{approx}} \approx \ln[\mathcal{C}(\mathcal{C}(0)) \neq \mathcal{M}_0] \mathcal{L}_0 \ln(\epsilon^{-1}) + \Delta H(\mathcal{M}_i).$$

Since the change $\Delta H(\mathcal{M}_i)$ in entropy is finite as $\epsilon \rightarrow 0$, the divergent nonreciprocity contribution to the dissipation dominates. Moreover, the simple scaling of the reliability-dissipation tradeoff is the same as that for the reliability-work tradeoff. In the low-error limit, the minimal nonreciprocity depends only on the computation:

$$\langle \mathcal{C}(\mathcal{M}_0 \rightarrow \mathcal{M}_i) \rangle \geq [C(C(m)) \neq \mathcal{M}_0] \mathcal{L}_0 \ln(\epsilon^{-1}).$$

The reciprocity coefficient

$$\langle [C(C(m)) \neq \mathcal{M}_0] \mathcal{L}_0 \rangle = \sum_{m \in \mathcal{M}} \mu_0(m) [C(C(m)) \neq m]$$

033524-9

BALANCING ERROR AND DISSIPATION IN COMPUTING

PHYSICAL REVIEW RESEARCH 2, 033524 (2020)
is the probability that the memory makes a transition that is not reciprocated if the output becomes the input to the computation. In short, computations with nonreciprocity require significant dissipation when implemented with a time-symmetric protocol.

A sequel explores the dynamical mechanism for the dissipation in this case. With time-symmetric protocols, state-space compression occurs only via a loss of stability and subsequent dissipation. This corresponds to a topological restriction on the bifurcation structure of local attractors—a restriction imposed by time-symmetric control.

One also concludes that strictly reciprocal transformations, such as the identity (do nothing), swap, and bit flip, where $C(C(m)) = m$ for all $m \in \mathcal{M}$, are exempt from this dissipation. One can efficiently implement them with time-symmetric protocols. For all other computations, time-asymmetric control is the only possible recourse to avoid this dissipation. Figure 4 gives examples of reciprocated and nonreciprocated memory transitions.

The main result Eq. (28) generalizes to the case where $m, m' \in \mathcal{M}$ without the requirement that $m = m'$. Then, dissipation diverges with increasing reliability whenever $m \rightarrow m'$ transitions are made such that $C(m) = m'$ but $C(m') \neq m'$. For uniform local free energies, Eq. (28) becomes

$$\beta (W_{\text{min}})_{\text{approx}} = \ln(e^{-1}) \sum_{m \in \mathcal{M}} p_0(m) [C(C(m)) \neq m^\dagger].$$

Notably, this allows one to quantify error–work and error–dissipation tradeoffs when computing with magnetic memory systems, for example. For a single bistable magnetic device, $m$ could represent the 1 memory of having an “up” magnetic moment or “clockwise” in the case of toroidal magnetic core memory elements. And, $m^\dagger$ would then represent the “down” or “counterclockwise” memory 0, as shown on the right side of Fig. 3. This implies that memories with certain types of symmetry are better suited for certain types of logical operation, in that they minimize nonreciprocity and so dissipation.

To emphasize, the dissipations here are distinct from the heat associated with logically irreversible transformations, as discussed by Landauer and Bennett [1,19], which arises as a compensation to microscopic state-space contraction. There are two important distinctions.

First, the heat devolved with logical irreversibility—e.g., the minimal heat of $k_B T \ln 2$ released to the environment upon erasure—is not necessarily irreversibly dissipated [2–5]. It is offset by a change in the nonequilibrium addition to free energy and so can be leveraged later to do an equal amount of useful work. In contrast, nonreciprocity dissipation is energy that is truly dissipated. It is irretrievably lost to the environment, with no compensation via change of the nonequilibrium addition to free energy and so can never be recovered.

Second, logical nonreciprocity is distinct from logical irreversibility. Strictly reciprocal computations, where $C(C(m)) = m$ for all $m \in \mathcal{M}$, are logically reversible, being their own inverses. However, logically reversible permutations of the memory can be completely nonreciprocal, as demonstrated in Fig. 4(d). Reciprocity requires not only that the deterministic logic be invertible, but further that the logical dynamic inverts itself. In short, the Landauer work cost corresponds to logical noninvertibility, while nonreciprocity cost corresponds to logical noninversion. The reciprocity bound on dissipation can therefore be interpreted as the minimal work required to implement a reliable computation with time-symmetric protocols in addition to the well-known Landauer bound.

IV. APPLICATIONS

With the basics of time-symmetric and nonreciprocal computing in hand, we turn to explore the thermodynamic implications for erasure, logic gates, and biological information processing. Our companion work Ref. [37] gives a detailed analysis of erasure and explores several model implementations.

A. Erasure

To directly illustrate how the cost of time-symmetric control differs from Landauer’s bound, consider the classic example of bit erasure. Landauer originally described a method for erasing a bit of information stored in a double-well potential in contact with a heat bath: tilt the potential—moving overdamped stochastic particles to one side—and return the potential to its original orientation, leaving the particles (temporarily) trapped in one well [19]. If implemented naively, via a protocol that raises the energy of one well and then lowers it at the same rate, the time-symmetry results in dissipation significantly above Landauer’s bound, as described by the bounds just developed. Note that protocol time-symmetry does not imply spatial symmetry—one can tilt the potential to the left without needing to tilt it to the right.
Bit erasure $C_{\text{erase}}$ operates on a single bistable element with memory states $\mathcal{M} = \{L, R\}$ that correspond to occupying the Left or Right side of a double-well potential energy landscape. The computation is defined by resetting to the $L$ memory state:

\[ C_{\text{erase}}(L) = L \text{ and } C_{\text{erase}}(R) = L. \]

Hence, $C_{\text{erase}}(C_{\text{erase}}(L)) = L$, while $C_{\text{erase}}(C_{\text{erase}}(R)) \neq R$. Thus, the net transition probabilities $p(m \rightarrow m')$ shown in Fig. 5 characterize any reliable implementation of this computation in terms of the probabilities of errors, $\epsilon_R = \epsilon_{R \rightarrow L}$ and $\epsilon_L = \epsilon_{L \rightarrow R}$, that leave the system in the $R$ state.

From this, the only terms contributing to the bound on time-symmetric work are

\[
d(L, R) = \epsilon_L \ln \frac{\epsilon_L}{1 - \epsilon_R} \text{ and } d(R, L) = (1 - \epsilon_R) \ln \frac{1 - \epsilon_R}{\epsilon_L}.
\]

Allowing for a potentially nonuniform input distribution $\mu_0 = (1 - p_R, p_R)$ over memory states yields the exact bound on time-symmetric work investment:

\[
\beta \langle W \rangle_{\text{sym}} = \langle 1 - p_R \rangle d(L, R) + p_R d(R, L)
\]

\[= (p_R(1 - \epsilon_R) - (1 - p_R)\epsilon_L) \ln \frac{1 - \epsilon_R}{\epsilon_L}.
\]

\[= (p_R - \langle \epsilon \rangle) \ln \frac{1 - \epsilon_R}{\epsilon_L}, \tag{31}\]

where $\langle \epsilon \rangle = p_R \epsilon_R + (1 - p_R)\epsilon_L$ is the average error of the computation.

Figure 6 plots the work requirements for an initially unbiased memory state with $p_R = 1/2$. We see that the work diverges for small values of error $\epsilon_L$. The plot also indicates that the work diverges for high values of error $\epsilon_R$, due to the nonreciprocity of the system between left and right. It is also worth noting from the plot that there are computations, such as bit flips, which this bound suggests may be achievable with time-symmetric control and without energetic cost. However, we are primarily concerned with the lower left portion of the plot, where effective erasures occur. The divergent scaling of the work required to reliably erase overwhelms the rather meager energy requirements given by Landauer’s bound. And, unlike the latter, this excess work requirement must be irreversibly dissipated. Subtracting off the Landauer bound $k_B T \Delta H(M)$ to calculate the minimum dissipation $\langle W_{\text{diss}} \rangle_{\text{sym}}$, Fig. 7 shows that the divergent contribution to the work is attributed to nonreciprocity.

Applying Eq. (28)’s work bounds to erasure, we see that

\[
\langle W \rangle_{\text{sym}} = \langle 1 - p_R \rangle d(L, R) + p_R d(R, L) = \langle p_R(1 - \epsilon_R) - (1 - p_R)\epsilon_L \rangle \ln \frac{1 - \epsilon_R}{\epsilon_L}
\]

\[\approx (p_R - \langle \epsilon \rangle) \ln \frac{1 - \epsilon_R}{\epsilon_L}, \tag{32}\]

where $\langle \epsilon \rangle = p_R \epsilon_R + (1 - p_R)\epsilon_L$ is the average error of the computation.

The initial memory-state entropy is $H_0(p_R) = -p_R \ln p_R - (1 - p_R) \ln (1 - p_R)$, which is called the binary entropy function [24]. Whereas, the final memory-state entropy vanishes as $\epsilon \rightarrow 0$. Hence, from Eq. (29), we immediately find that the small-error dissipation necessary for time-symmetric erasure diverges as $\epsilon \rightarrow 0$:

\[
\langle W_{\text{diss}} \rangle_{\text{sym}} = p_R \ln (\epsilon) k_B T - H_0(p_R) k_B T. \tag{33}\]

Landauer erasure typically assumes a uniform initial distribution where $p_R = 1/2$. This results in $\langle W_{\text{diss}} \rangle_{\text{sym}} \gtrsim \frac{1}{4} \ln (1/4\epsilon) k_B T$. Since the memory-state entropy is bounded by $k_B T \ln 2$, the contribution from Landauer’s bound is negligible compared to the $\ln(\epsilon)$ term for small $\epsilon$. And so, the latter dominates both the work and dissipation for high-fidelity erasure.

FIG. 5. Markov transition matrix for any implementation of erasure gives the error rate $\epsilon_L = \epsilon_{L \rightarrow R}$ from initial memory state $M_0 = L$ and the error rate $\epsilon_R = \epsilon_{R \rightarrow L}$ from the initial memory state $M_0 = R$.

FIG. 6. Average minimum work (units of $k_B T$) required for a 1-bit operation diverges for low $\epsilon_L$ or high $\epsilon_R$, corresponding to erasure to L or erasure to R, respectively. The state-transition diagrams at the corners show the basic computations associated with each extremal point in computation parameter space. The identity map (upper left), the bit-flip (lower right), and every operation interpolated between require no work according to the time-symmetric control bound. The required work diverges at perfect erasure to Left (bottom-left corner) and perfect erasure to Right (top-right corner), where the red coloring covers all work values from 1.4 to $\infty k_B T$. Where effective erasures occur. The divergent scaling of the work required to reliably erase overwhelms the rather meager energy requirements given by Landauer’s bound. And, unlike the latter, this excess work requirement must be irreversibly dissipated. Subtracting off the Landauer bound $k_B T \Delta H(M)$ to calculate the minimum dissipation $\langle W_{\text{diss}} \rangle_{\text{sym}}$, Fig. 7 shows that the divergent contribution to the work is attributed to nonreciprocity.

Applying Eq. (28)’s work bounds to erasure, we see that

\[
\langle W \rangle_{\text{sym}} = \langle 1 - p_R \rangle d(L, R) + p_R d(R, L) = \langle p_R(1 - \epsilon_R) - (1 - p_R)\epsilon_L \rangle \ln \frac{1 - \epsilon_R}{\epsilon_L}
\]

\[\approx (p_R - \langle \epsilon \rangle) \ln \frac{1 - \epsilon_R}{\epsilon_L}, \tag{32}\]

where $\langle \epsilon \rangle = p_R \epsilon_R + (1 - p_R)\epsilon_L$ is the average error of the computation.

The initial memory-state entropy is $H_0(p_R) = -p_R \ln p_R - (1 - p_R) \ln (1 - p_R)$, which is called the binary entropy function [24]. Whereas, the final memory-state entropy vanishes as $\epsilon \rightarrow 0$. Hence, from Eq. (29), we immediately find that the small-error dissipation necessary for time-symmetric erasure diverges as $\epsilon \rightarrow 0$:

\[
\langle W_{\text{diss}} \rangle_{\text{sym}} = p_R \ln (\epsilon) k_B T - H_0(p_R) k_B T. \tag{33}\]

Landauer erasure typically assumes a uniform initial distribution where $p_R = 1/2$. This results in $\langle W_{\text{diss}} \rangle_{\text{sym}} \gtrsim \frac{1}{4} \ln (1/4\epsilon) k_B T$. Since the memory-state entropy is bounded by $k_B T \ln 2$, the contribution from Landauer’s bound is negligible compared to the $\ln(\epsilon)$ term for small $\epsilon$. And so, the latter dominates both the work and dissipation for high-fidelity erasure.
FIG. 7. Dissipated work—that required beyond the Landauer bound—behaves much the same as the minimum work over possible bit computations, diverging for high values of $\varepsilon_R$ and low values of $\varepsilon_L$ and vanishing for the identity map and bit flip. Dissipation diverges at the bottom-left and top-right corners, where the red coloring covers all values of dissipated work from 1.4 to $\infty k_BT$.

To test the strength of these bounds and their approximations, Ref. [37] considers two different implementations of time-symmetric erasure—rate equations for a two-level system and one-dimensional Langevin dynamics in a controlled double-well potential. The example implementations not only validate our bounds, but moreover show that the bounds are tight.

B. Logic gates

We next consider the minimal dissipation required to implement conventional two-input one-output logic gates that serve as the basis for modern digital computing: AND, NAND, OR, NOR, XOR, and the like. The typical implementation requires two bistable memory elements ($M^{(in)}_t$) and $M^{(in)}_t$ for the input and another bistable memory element ($M^{(out)}_t$) to robustly store the output. Each memory element can take on one of two memory states $M^{(i)} \in \{0, 1\}$. Each logic gate is a computation $C$ on the set of composite memory states $\mathcal{M} = M^{(in)}_t \times M^{(in)}_t \times M^{(out)}_t = \{000, 001, 010, \ldots, 111\}$.

In particular, since networks of NAND gates are sufficient for universal computation, the NAND gate is worthy of immediate investigation. Reference [59], part D, analyzes the NAND gate, showing that: if the memory elements are initiated statistically independent of each other, then the minimal time-symmetric-implement dissipation is comparable to that expected for bit erasure of the output bit, depending on how often the output bit is overwritten. However, when the memory system is initialized with correlation among the memory elements (perhaps as the correlated output of previous upstream computations), then the change in memory entropy is $\Delta H(M_t) \approx -H(M^{(out)}_t|M^{(in)}_t; M^{(in)}_t)$ in the small $\epsilon$ limit, and the reciprocity coefficient is

$$\langle [C_{NAND}(C_{NAND}(M_0)) \neq M_0] \rangle_{M_0}$$

$$= \mu_0(000) + \mu_0(010) + \mu_0(100) + \mu_0(111). \quad (34)$$

As a consequence, the initial correlation among the memory elements has a profound impact on the coefficient of minimal dissipation. The overall dissipation still diverges $\sim \ln(\epsilon^{-1})$, though, with increasing reliability as long as the implementation is time-symmetric.

This example highlights a previously unsuspected source of unnecessary and preventable dissipation in digital computers. Indeed, computations are currently implemented in hardware, whether via periodic clocking or static gate voltages, through time-symmetric driving. Error rates in today’s reliable digital computers are extremely low. The so-called soft error rate (also known as single-event upsets) for 50 nm gate technology has been estimated to be on the order of $10^{-4}$ errors per $10^9$ hours of operation of a CMOS logic circuit [38]. Moreover, these errors are overwhelmingly due to cosmic rays rather than conventional thermal fluctuations. If the computer is properly shielded, the error rate is significantly lower. Using the aforementioned numbers for a conservative estimate and assuming a 3 GHz clock rate, this extreme reliability of $\epsilon \lesssim 10^{-20}$ errors/instruction would imply at least 30 $k_BT$ is dissipated in nearly every elementary operation in modern circuitry directly due to time-symmetric implementation. This is well beyond the Landauer bound that, for NAND and any other traditional two-input-one-output logic gate, is given by

$$-\Delta H(M_t) \approx H(M^{(out)}_t|M^{(in)}_t; M^{(in)}_t) \leq \ln 2.$$

By way of contrast, the universal Fredkin gate—swap the state of the first two memory elements only if the third memory element is in state 1—is not only logically reversible, but is also a strictly reciprocal gate [39]. As a strictly reciprocal gate, it can implement universal computation efficiently using time-symmetric control. This makes Fredkin gates an especially tempting basis for energetically efficient future computing technologies, assuming they can be made to transform metastable memories without the exponential sensitivity to initial conditions that condemned their initially proposed mechanical instantiation [1,40].

C. Dissipation in biological processing

Beyond logical computation, the reliability–energy-efficiency tradeoff also applies to information processing in far-from-equilibrium biological systems. For this, it is useful to reformulate Eq. (3) to apply when $W_{\text{disc}}$ is the work dissipated in controlling (through $x_{0,1}$) a system already maintained out of equilibrium by a given housekeeping entropy production $\Sigma_{hk}$. This can happen, for example, via ATP hydrolysis or the maintenance of a chemical potential gradient. Then, according to Eq. (27) of Ref. [41], the more general starting point is

$$\beta W_{\text{disc}} + k_B^{-1} \Sigma_{hk} = \ln \frac{P_{x_{0,1}}(s_{0,1}) = s_0) = s_0 \sim \mu_0}{P_{x_{0,1}}(s_{0,1}) = s_0) = s_0 \sim \mu_1).} \quad (35)$$
In steady-state biological processes, there is no external control. The controllable parameters can thus be considered as held constant, such that the driving protocol is trivially time-symmetric $\mathbf{M}(x_{0:t}) = x_{0} = x_{0} x_{0} \ldots x_{0}$ and $W_{\text{det}} = 0$. That is, the system is driven solely by the housekeeping entropy production—power supplied from a stationary stochastic influence—that maintains the steady state. For example, molecular machines can sustain time-asymmetric functionality within their time-symmetric ambient chemical environment due to the entropy they produce during chemical catalysis [15, 18]. In these nonequilibrium steady states, the expected microstate distribution of the system is the time-invariant steady-state distribution:

$$\mu_{0} = \mu_{\tau} = \pi_{\text{s.s.}}. \quad (36)$$

Moreover, due to the low Reynolds numbers of the microbiological realm, momentum is heavily damped and the relevant memory states are time-reversal-invariant: $m^\prime = m$ for all $m \in \mathcal{M}$. We also have $\pi_{\text{s.s.}}^\prime = \pi_{\text{s.s.}}$.

Integrating these generic features of steady-state biological transformations, Eq. (35) simplifies to

$$k_{B}^{-1} \Sigma_{\text{hs}} = \text{ln} \left( \frac{\text{Pr}(S_{0:\tau} = s_{0:\tau} | S_{0} = s_{0}) \pi_{\text{s.s.}}(s_{0})}{\text{Pr}(S_{0:\tau} = \mathbf{M}(s_{0:\tau}) | S_{0} = s_{\tau}) \pi_{\text{s.s.}}(s_{\tau})} \right). \quad (37)$$

The single-timestep version of this is well known:

$$\delta \Sigma_{\text{hs}} = k_{B} \text{ln} \left( \frac{\text{Pr}(S_{\text{det}} = s_{\text{det}} | S_{0} = s_{0}) \pi_{\text{s.s.}}(s_{0})}{\text{Pr}(S_{0:\tau} = s_{0} | S_{0} = s_{0}) \pi_{\text{s.s.}}(s_{0})} \right).$$

As $\delta t \to 0$, the relative probabilities reduce to the relative transition rates $\{r_{m \to m^\prime}\}$:

$$\delta \Sigma_{\text{hs}} = k_{B} \text{ln} \left( \frac{\pi_{\text{s.s.}}(s_{0}) r_{s_{0} \to s_{m}}}{\pi_{\text{s.s.}}(s_{m}) r_{s_{m} \to s_{0}}} \right). \quad (38)$$

(Metastability of mesoscopic conformations allows Eq. (38) to hold approximately for transitions between mesoscopic states as well.) For example, Eq. (38) was the starting point for the recently derived thermodynamic uncertainty relations that received attention [42].

However, much can be learned from approaching finite-duration transformations directly, by coarse-graining Eq. (37) according to initial and final functionally relevant states. Notably, it allows simple yet powerful analysis of the minimal dissipation required for biological functionality, regardless of how complicated the finite-duration biological implementations appear. And, it implies a generic error–dissipation tradeoff that we expect to apply broadly to the reliable performance of microbiological systems.

Working this out explicitly leads to a pleasantly simple picture of steady-state biological information processing and its requisite dissipation:

$$k_{B}^{-1} \langle \Sigma_{\text{hs}} \rangle \geq D_{\text{KL}}[\text{Pr}(M_{0} = m, M_{\tau} = m') | | \text{Pr}(M_{\tau} = m', M_{2\tau} = m)]$$

$$= \Delta H(M_{\tau}) + \sum_{m, m' \in \mathcal{M}} \pi_{\text{s.s.}}(m) p(m \to m') \ln \frac{p(m \to m')}{p(m' \to m)}$$

$$= \langle \Psi(M \to M') \rangle_{\pi_{\text{s.s.}}(M) p(M \to M')} \cdot (39)$$

Steady-state computation quite simply becomes enforcing a Markov model for memory transitions $\{p(m \to m')\}$. The minimal dissipation achievable via any biological implementation then equals or exceeds the expected nonreciprocity. It should be noted that the steady-state distribution $\pi_{\text{s.s.}}$ over memory states is easily found as the stationary eigenstate of the Markov model’s transition matrix.

In equilibrium, where $\pi_{\text{s.s.}} = \pi_{\text{eq}}$, the average nonreciprocity vanishes since detailed balance demands $\pi_{\text{eq}}(m) p_{\text{eq}}(m \to m') = \pi_{\text{eq}}(m') p_{\text{eq}}(m' \to m)$ while, as always, $\Psi(m \to m') = -\Psi(m' \to m)$. As necessary, this allows zero dissipation in equilibrium. In contrast, nonequilibrium steady-state computations must dissipate since they bias nonreciprocated transitions. These computations enforce nondetailed-balanced dynamics.

Equation (39) shows that dissipation in steady-state biological processes is bounded by the average nonreciprocity.

Remarkably, this bound depends only on the net transformation and is independent of intermediary steps and biochemical details.

Thus, as in the preceding analysis, one expects a generic error–dissipation tradeoff:

$$d \Sigma_{\text{hs}} / dt \propto \ln(1/\epsilon)$$

for reliable steady-state biological transformations of genetic information—whenever certain transitions need to be strongly enforced with only a small probability of failure $\epsilon$. The generic error–dissipation tradeoff holds, as an important example, for processes such as chemical proofreading necessary for reliable DNA replication. In this way, our contribution dovetails with more model-specific results on the latter by Refs. [43–46]. Since codons are nonreciprocally corrected to satisfy Watson-Crick base pairing, each correction requires significant dissipation.

The $d \Sigma_{\text{hs}} / dt \propto \ln(1/\epsilon)$ error–dissipation tradeoff should arise in other reliable biological transformations as well. For example, Ref. [47] finds exactly this minimum dissipation rate to reliably maintain a target nonequilibrium state. This
is relevant to protein folding, as well as other biological functionalities.

Our results also support and update the perspective of Ref. [48], where steady-state biochemical copying networks (e.g., for sensing) were found to have an unavoidable error-dissipation tradeoff due to time-invariant driving. Our results more broadly suggest error–dissipation tradeoffs for any biochemical task with nonreciprocity. It should now be clear that autonomous biochemical systems in steady state can never reach the Landauer bound on efficiency due to the time-symmetric control penalty.

V. RELATED RESULTS

Let’s consider how the tradeoffs between accuracy and energetic-efficiency under time-symmetric protocols compare to related approaches to thermodynamic control, both historically and currently.

In Ref. [49], Brillouin defines the reliability as 1/ε, where ε is the probability of reading out the wrong value due to thermal agitation. He found that the entropy production necessary for some methods of reliable readout is $k_B \ln(1/\epsilon)$. It is interesting that Brillouin’s readout result mirrors the same error–dissipation scaling as found in our analysis, despite the fact that detailed fluctuation theorems, the foundations of our analysis, were still decades away. Brillouin’s readout penalty is distinct from our result, but it hints at a more general error–dissipation tradeoff (under particular physical constraints) that could unite the results.

Brillouin’s readout penalty also appeared in disguise in Ref. [50], which again, superficially, appears to propose a similar error–dissipation tradeoff to what we have identified. As in Ref. [51], Ref. [50] defines a ‘logic operation’ specifically as bit inversion, 0 → 1 and 1 → 0—a strictly reciprocal computation. Reference [50] thus defines the “minimal work per logic operation” as the minimal work for bit inversion. Despite similarities in appearance, it is clearly different from our result since it predicts divergent dissipation for reliable implementation of a strictly reciprocal computation. In this setting, if the voltage drifts from one input to the other (a readout error), then the wrong input will be inverted. While some computational implementations may require further inefficiency to avoid this source of error, it is distinct and perhaps additive to our nonreciprocity dissipation.

A discussion of reliable computing would be incomplete without mention of von Neumann’s well-known analysis of error correction [52]. Indeed, it might seem plausible that less reliable components can be harnessed to avoid the error–dissipation tradeoff while nevertheless achieving the same net reliability through error correction. However, applying the analysis above to account for the dissipation from all components, one finds that error correction via redundancy leads to a worse error–dissipation tradeoff. We report on this elsewhere.

THERMODYNAMICALLY INSPIRED STUDIES OF CLASSICAL REVERSIBLE COMPUTING, E.G., BY BENNETT, FREDKIN, AND TOFOLLI [1,40], INSPIRED MUCH EARLY WORK ON QUANTUM COMPUTATION—E.G., IN REF. [53–55]—WHERE LOGICAL REVERSIBILITY OF THE UNITARY TRANSFORMATIONS IS MANIFEST. HOWEVER, NONRECIPROCITY BRINGS DISSIPATION BACK TO THE TABLE, EVEN FOR REVERSIBLE LOGIC. WE HOPE IT IS NOW CLEAR THERE IS A NECESSARY RESEARCH PROGRAM TO INVESTIGATE NONRECIPROCITY DISSIPATION IN QUANTUM COMPUTING.

VI. DISCUSSION: TIME ASYMMETRY?

Another insight from the preceding is exploring time-asymmetric control when designing energetically efficient computation. However, is time-asymmetric free? Often in thermodynamic control, one posits an external signal that can be produced in any sequence, reversible or not, at no cost beyond the work-energy imparted to the driven system. Inexpensive time asymmetry can be generated, for example, from inertial degrees of freedom [58].

However, in biological systems and in many engineered nanoscale systems, inertial degrees of freedom are not available. Noninertial batteries reliably transforming noninertial systems will drain according to our time-symmetric-control bounds on dissipation.

Time-symmetric control might also seem to apply, rather broadly, if one removes the boundaries separating a driven system from its driver. Including the driver as part of an enlarged system gives the appearance of trivially time-symmetric driving of the composite system. Do our time-symmetric-control bounds on dissipation then apply to any reliable computations that occur within this enlarged system? Generically, our bounds will not apply to such enlarged systems because the enlarged microstate distributions $\mu_0$ and $\mu_\tau$ will no longer be metastable.

This suggests that time-asymmetry and instability can both lead to energetic advantages. Future efforts explore which features of time-asymmetry matter for efficiency. This helps identify those qualitative features of time-symmetric control that must be overcome for improved efficiency.

For time-symmetric protocols, we now ask: How does $\epsilon$ scale with $\tau$ or with the modularity of control? Generically, we do not expect a one-to-one relationship between the error and duration. Rather, generally, there will be a high-dimensional space of tradeoff scalings that includes speed, error, modularity of control, robustness of information storage, energetic efficiency, and many other computationally relevant design restrictions.

033524-14
VII. CONCLUSION

We discovered that time symmetry and metastability together imply a general error–dissipation tradeoff. The minimal work expected for a computation $C$ is the average nonreciprocity. In the low-error limit—where the probability of error must be less than $\epsilon$—the minimum work diverges according to

$$\beta W_{\text{min}}^{\text{approx}} = \langle [C(M_0)] \rangle \approx M_0 \ln(\epsilon^{-1}).$$

Of all of this work, only the meager Landauer cost $\Delta H_0(M_{\epsilon})$, which saturates to some finite value as $\epsilon \to 0$, can be retrieved in principle. Thus, irretrievable dissipation scales as $\ln(\epsilon^{-1})$. The reciprocity coefficient $\langle [C(M_0)] \rangle \approx M_0 \ln(\epsilon^{-1})$ depends only on the deterministic computation to be approximated. This, in turn, identifies sources of energy inefficiency in current implementations of reliable computation. It also implies that time-asymmetric control can allow for more efficient computation—but only when time-asymmetric signals are a free resource.

Restricting to time-symmetric driving may seem unusual. However, with the time-symmetric clock signal that drives modern microprocessors, it is the norm rather than the exception in the realm of computational controls. Sections IV B and IV C discussed the naturalness of time-symmetry in digital computers and in genetic processing, respectively. To get there, though, required developing several prerequisite thermodynamic results.

After developing the general thermodynamics, we analyzed the limit of highly reliable computing. As important examples, we then specialized to dissipation in erasure, logic gates, and reliable biological transformations.

Overall, the results elevate time-asymmetric control to a design principle for efficient thermodynamic computing. This then must be added to the growing list of recently discovered principles, including tracking system modularity [5,33] and logic gates whose protocols adapt to their input [5]. Further progress will turn on how these lessons are incorporated as constraints in the principled design and search for near-optimal finite-time protocols. An opportunity presents itself to adapt these lessons to developing fast hyper-efficient computers in the not-so-distant future.

ACKNOWLEDGMENTS

The authors thank Gavin Crooks, Mike DeWeese, and Chris Jarzynski for helpful discussions and the Telluride Science Research Center for hospitality during visits and the participants of the Information Engines Workshops there. J.P.C. acknowledges the kind hospitality of the Santa Fe Institute, Institute for Advanced Study at the University of Amsterdam, and California Institute of Technology for their hospitality during visits. This material is based upon work supported by, or in part by, Foundational Questions Institute Grant number FQXi-RFP-IPW-1902, Templeton World Charity Foundation Power of Information fellowship TWCFO337, and U.S. Army Research Laboratory and the U.S. Army Research Office under contracts W911NF-13-1-0390 and W911NF-18-1-0028.

[1] C. H. Bennett, Thermodynamics of computation—A review, Int. J. Theor. Phys. 21, 905 (1982).
[2] O. J. E. Maroney, Generalizing Landauer’s principle, Phys. Rev. E 79, 031105 (2009).
[3] T. Sagawa, Thermodynamic and logical reversibilities revisited, J. Stat. Mech. (2014) P03025.
[4] J. M. R. Parrondo, J. M. Horowitz, and T. Sagawa, Thermodynamics of information, Nat. Phys. 11, 131 (2015).
[5] P. M. Riechers, Transforming metastable memories: The nonequilibrium thermodynamics of computation, in The Energetics of Computing in Life and Machines, edited by D. Wolpert, C. Kempes, P. Stadler, and J. Grochow (SFI Press, Santa Fe, New Mexico, 2019).
[6] P. Salamon and R. S. Berry, Thermodynamic Length and Dissipated Availability, Phys. Rev. Lett. 51, 1127 (1983).
[7] D. A. Sivak and G. E. Crooks, Thermodynamic Metrics and Optimal Paths, Phys. Rev. Lett. 108, 190602 (2012).
[8] P. R. Zulkowski and M. R. DeWeese, Optimal finite-time erasure of a classical bit, Phys. Rev. E 89, 052140 (2014).
[9] M. V. S. Bonança and S. Deffner, Optimal driving of isothermal processes close to equilibrium, J. Chem. Phys. 140, 244119 (2014).
[10] D. Mandal and C. Jarzynski, Analysis of slow transitions between nonequilibrium steady states, J. Stat. Mech. (2016) 063204.
[11] A. B. Boyd, A. Patra, C. Jarzynski, and J. P. Crutchfield, Shortcuts to thermodynamic computing: The cost of fast and faithful information processing, arXiv:1812.11241.
[12] A. P. Chandrakasan, S. Sheng, and R. W. Brodersen, Low-power CMOS digital design, IEEE J. Solid-State Circ. 27, 473 (1992).
[13] A. Iyer and D. Marculescu, Power efficiency of voltage scaling in multiple clock, multiple voltage cores, in Proc. 2002 IEEE/ACM Intl. Cong. Computer-aided Design (ACM Press, San Jose, California, 2002), pp. 379–386.
[14] J. S. Seldenthuis, F. Prins, J. M. Thijssen, and H. S. J. van der Zant, An all-electric single-molecule motor, ACS Nano 4, 6681 (2010).
[15] R. D. Astumian, How molecular motors work–insights from the molecular machinist’s toolbox: The Nobel Prize in Chemistry 2016, Chem. Sci. 8, 840 (2017).
[16] A. I. Brown and D. A. Sivak, Toward the design principles of molecular machines, arXiv:1701.04868.
[17] L. Zhang, V. Marcos, and D. A. Leigh, Molecular machines with bio-inspired mechanisms, Proc. Natl. Acad. Sci. USA 115, 9397 (2018).
[18] E. H. Feng and G. E. Crooks, Length of Time’s Arrow, Phys. Rev. Lett. 101, 090602 (2008).
[19] R. Landauer, Irreversibility and heat generation in the computing process, IBM J. Res. Develop. 5, 183 (1961).
[20] G. E. Crooks, Entropy production fluctuation theorem and the nonequilibrium work relation for free energy differences, Phys. Rev. E 60, 2721 (1999).
[21] C. Jarzynski, Hamiltonian derivation of a detailed fluctuation theorem, J. Stat. Phys. 98, 77 (2000).
[22] U. Seifert, Entropy Production Along a Stochastic Trajectory and an Integral Fluctuation Theorem, Phys. Rev. Lett. 95, 040602 (2005).
[23] M. Esposito and C. Van den Broeck, Second law and Landauer principle far from equilibrium, Europhys. Lett. 95, 4 (2011).
[24] T. M. Cover and J. A. Thomas, Elements of Information Theory, 2nd ed. (Wiley-Interscience, New York, 2006).
[25] M. Esposito, Stochastic thermodynamics under coarse graining, Phys. Rev. E 85, 041125 (2012).
[26] C. Jarzynski, Rare events and the convergence of exponentially averaged work values, Phys. Rev. Lett. 73, 046105 (2006).
[27] R. Kawai, J. M. R. Parrondo, and C. Van den Broeck, Dissipation: The Phase-Space Perspective, Phys. Rev. Lett. 105, 150607 (2010).
[28] A. Gomez-Marín, J. M. R. Parrondo, and C. Van den Broeck, Lower bounds on dissipation upon coarse graining, Phys. Rev. E 78, 011107 (2008).
[29] E. Roldan and J. M. R. Parrondo, Estimating Dissipation from Single Stationary Trajectories, Phys. Rev. Lett. 98, 080602 (2007).
[30] A. Gomez-Marín, J. M. R. Parrondo, and C. Van den Broeck, Lower bounds on dissipation upon coarse graining, Phys. Rev. E 78, 011107 (2008).
[31] E. Roldan and J. M. R. Parrondo, Estimating dissipation from single stationary trajectories, PhD thesis, Universidad Complutense de Madrid, Madrid, Spain, 2010.
[32] Y. Jun, M. Gavrilov, and J. Bechhoefer, Probability density functions for the initial distribution over states, J. Stat. Mech. (2017) 083202.
[33] J. L. England, Statistical physics of self-replication, J. Chem. Phys. 139, 121923 (2013).
[34] G. Wimsatt, O.-P. Saira, A. B. Boyd, M. H. Matheny, S. Han, M. L. Roukes, and J. P. Crutchfield, Harnessing fluctuations in thermodynamic computing via time-reversal symmetries, arXiv:1906.11973.
[35] G. W. Wimsatt, A. B. Boyd, P. M. Riechers, and J. P. Crutchfield, Balancing error and dissipation when erasing information (unpublished).
[36] P. Shivakumar, M. Kistler, S. W. Keckler, D. Burger, and L. Alvisi, Modeling the effect of technology trends on the soft error rate of combinational logic, in Dependable Systems and Networks, Dependable Systems and Networks, Proceedings International Conference on DSN 2002 (IEEE, Bethesda, Maryland, 2002), pp. 389–398.
[37] E. Fredkin and T. Toffoli, Conservative logic, Int. J. Theor. Phys. 21, 219 (1982).
[38] E. Fredkin and T. Toffoli, Conservative logic, in Collision-Based Computing (Springer, Heidelberg, 2002), pp. 47–81.
[39] P. M. Riechers and J. P. Crutchfield, Fluctuations when driving between nonequilibrium steady states, J. Stat. Phys. 168, 873 (2017).
[40] T. R. Gingrich, J. M. Horowitz, N. Perunov, and J. L. England, Dissipation Bounds all Steady-State Current Fluctuations, Phys. Rev. Lett. 116, 120601 (2016).
[41] J. J. Hopfield, Kinetic proofreading: A new mechanism for reducing errors in biosynthetic processes requiring high specificity, Proc. Natl. Acad. Sci. USA 71, 4135 (1974).
[42] Charles H. Bennett, Dissipation-error tradeoff in proofreading, Biosystems 11, 85 (1979).
[43] J. M. Horowitz, K. Zhou, and J. L. England, Minimum energetic cost to maintain a target nonequilibrium state, Phys. Rev. E 95, 042102 (2017).
[44] T. E. Ouldridge, C. C. Govern, and P. R. ten Wolde, Thermodynamics of Computational Copying in Biochemical Systems, Phys. Rev. X 7, 021004 (2017).
[45] L. Brillouin, Science and Information Theory (Academic Press, New York, 1956).
[46] A. Murugan, D. A. Huse, and S. Leibler, Speed, dissipation, and error in kinetic proofreading, Proc. Natl. Acad. Sci. USA 109, 12034 (2012).
[47] E. Sartori and S. Pigolotti, Thermodynamics of Error Correction, Phys. Rev. X 5, 041039 (2015).
[48] J. M. Horowitz, K. Zhou, and J. L. England, Minimum energetic cost to maintain a target nonequilibrium state, Phys. Rev. E 95, 042102 (2017).
[49] T. E. Ouldridge, C. C. Govern, and P. R. ten Wolde, Thermodynamics of Computational Copying in Biochemical Systems, Phys. Rev. X 7, 021004 (2017).
[50] K.-L. Stein, Noise-induced error rate as limiting factor for energy per operation in digital ICs, IEEE J. Solid-State Circ. SC-12, 527 (1977).
[51] R. Muller, H. J. Pfeiferderer, and K. U. Stein, Energy per logic operation in integrated circuits: Definition and determination, IEEE J. Solid-State Circ. 11, 657 (1976).
[52] J. von Neumann, Probabilistic logics and the synthesis of reliable organisms from unreliable components, in Automata Studies, edited by C. E. Shannon and J. McCarthy, Annals of Mathematical Studies No. 34 (Princeton University Press, Princeton, New Jersey, 1956), pp. 329–378.
[53] P. A. Benioff, Quantum mechanical Hamiltonian models of discrete processes that erase their own histories: Application to Turing machines, Intl. J. Theo. Phys. 21, 177 (1982).
[54] W. H. Zurek, Reversibility and Stability of Information Processing Systems, Phys. Rev. Lett. 53, 391 (1984).
[55] A. Peres, Reversible logic and quantum computers, Phys. Rev. A 32, 3266 (1985).
[56] G. Lan, P. Sartori, S. Neumann, V. Sourjik, and Y. Tu, The energy–speed–accuracy trade-off in sensory adaptation, Nat. Phys. 8, 422 (2012).
[57] S. Lahiri, J. Sohl-Dickstein, and S. Ganguli, A universal tradeoff between power, precision and speed in physical communication, arXiv:1603.07758.
[58] S. Defner and C. Jarzynski, Information Processing and the Second Law of Thermodynamics: An Inclusive, Hamiltonian Approach, Phys. Rev. X 3, 041003 (2013).
[59] See Supplemental Material at http://link.aps.org/supplemental/10.1103/PhysRevResearch.2.033524 for a general bound on dissipated work in thermodynamic computing, a transition-specific fluctuation theorem, detailed error analysis for the four cases of memory transitions, and the minimal dissipation in the time-symmetric universal NAND gate.