Non-Markovian Momentum Computing: Thermodynamically Efficient and Computation Universal

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Practical, useful computations are implemented via physical processes. Information must be stored and updated within a system’s configurations, whose energetics determine a computation’s cost. To describe thermodynamic and biological information processing, a growing body of results embraces rate equations as the underlying mechanics of computation. Strictly applying these continuous-time stochastic Markov dynamics, however, precludes a universe of natural computing. Within this framework, operations as simple as a NOT gate and flipping a bit are inaccessible. We show that expanding the toolset to continuous-time hidden Markov dynamics substantially removes the constraints, by allowing information to be stored in a system’s latent states. We demonstrate this by simulating computations that are impossible to implement without hidden states. We design and analyze a thermodynamically-costless bit flip, providing a counterexample to rate-equation modeling. We generalize this to a costless Fredkin gate—a key operation in reversible computing that is Turing complete (computational universal). Going beyond rate-equation dynamics is not only possible, but necessary if stochastic thermodynamics is to become part of the paradigm for physical information processing. The increased analytical challenges are readily addressed with recently-introduced spectral decomposition methods for nondiagonalizable dynamics.

Keywords: rate equations, stochastic process, hidden Markov model, information processing, logical circuits, ion channel, entropy production, reversibility

The burgeoning field of thermodynamic computing leverages recent progress in nonequilibrium thermodynamics and information and computation theories [1–5] to establish a new paradigm for physical information processing that promises to increase computational power and efficiency and to reduce energy dissipation in a next generation of computers [6]. Thermodynamic computing is distinguished from alternative paradigms by its focus on an information-processing device’s physical embedding; specifically, by constructively working with $k_B T$-scale fluctuations that a thermal environment generates. More broadly, a general framework rooted in thermodynamics, as thermodynamic computing is, will provide the tools to understand the physics of computation in all its many forms. The following illustrates its breadth by introducing non-Markovian, momentum-based computing—a paradigm that is both computation universal and thermodynamically efficient.

Physically-embedded computation can be described as a stochastic mapping within a system’s set $\mathcal{M}$ of memory states—Landauer’s information-bearing degrees of freedom (IDoF) [7]. Carried out over time interval $t \in (0, \tau)$, the mapping is the conditional probability $\mathbf{p}$ of transitioning from an initial memory state $m(0) \in \mathcal{M}$ to a final state $m(\tau) \in \mathcal{M}$:

$$\mathbf{p}_{m(0) \rightarrow m(\tau)} = \Pr [m(\tau) | m(0)] .$$

The mapping $\mathbf{p}$ determines the probability of the final memory state given the initial memory state, and so updates the state distribution $\tilde{p}(\tau) = \mathbf{p} \tilde{p}(0)$. This describes the dynamics underlying a computation, but what of its thermodynamic consequences? To address this, we must first identify the constraints on dynamics that can implement computations.

Computing with Continuous-Time Markov Chains To date, proposed frameworks for the required mappings in thermodynamic computing assume that the memory state $m$ obeys stochastic Markovian dynamics [8, 9, and references therein]. Taking time to be continuous, the dynamics are continuous-time Markov chains (CTMCs), where the state distribution changes continuously as a function of itself $\tilde{p}(t) = f(\tilde{p}, t)$. The resulting dynamics is necessarily represented by a master equation over the memory-state distribution $\tilde{p}(t) = \mathbf{A}(t) \tilde{p}(t)$ [8, 9]; that is, by rate equations. This is a powerful framework for stochastic thermodynamics [2, 10, 11] that yields insight into physical realizations of computations such as bit erasure and measurement [12].

The constraint that the computation $\mathbf{p}$ is generated by integrating continuous-time master equations comes at a substantial compromise—it limits the range of possible computations. For example, only input-output mappings whose determinants are positive are allowed when memory-state dynamics are restricted to obey CTMCs [9]. This eliminates many common and useful computations,
including flipping a single bit of information. Reference [8] takes these restrictions as delineating the possible physically realizable computations. Given that any computation we can observe—a bit flip, to take one example—is necessarily physical, one must instead interpret the restrictions as a limitation of the CTMC framework, rather than of the physical world.

Understanding both the merits and limits of the CTMC framework requires closely examining the physical mechanisms that underpin it. As a first cut, it is natural to say that the memory states $\mathcal{M}$ are microstates of a physical memory system $\mathcal{S}$, evolving under Hamiltonian dynamics. However, this strongly limits computations to be deterministic and reversible. More realistically (and fortunately) physical computational devices are coupled to an environment (heat bath) that introduces randomness and irreversibility and acts a source of heat to the memory system evolution. It is the system and environment altogether that evolve deterministically according to reversible Liouville dynamics.

That said, our concern is primarily with the performance of the memory system. If a large weakly-coupled heat bath has degrees of freedom that relax sufficiently quickly, it retains no memory of the memory system’s microstates. And so, the effective dynamics for the degrees of freedom of $\mathcal{S}$ considered alone must not just be stochastic, but also Markovian and therefore CTMC [2, 13–16]. In essence, coarse-graining the thermal environment allows for accurate, probabilistic predictions about the memory system, while avoiding the complicated task of tracking the full Hamiltonian dynamics of the joint system and bath.

This justification of the Markov evolution of a memory system recognizes an important fact: $\mathcal{S}$’s states are not themselves full descriptions of physical degrees of freedom. Instead, they are mesostates defined by a coarse-graining over the thermal environment’s microstates. This coarse-graining is appropriate since the environment does not retain information about the past.

Computationally-useful memory states $\mathcal{M}$—Landauer’s IDoF—are mesostates that also coarse-grain over $\mathcal{S}$’s CTMC-evolving states. It is possible, depending on the variables and timescales of interest, that this coarse-graining ignores only rapidly-relaxing subsystems of $\mathcal{S}$ and, then, the IDoF inherit the Markov property of the memory system. The result is a powerful and widely-used framework for thermodynamic computing in which the IDoF also obey CTMC dynamics. This case is typified by IDoF that are positional degrees of freedom and where $\mathcal{S}$ is described by overdamped Langevin dynamics.

**Computing with Continuous-Time Hidden Markov Chains**

However, in the most general case, the IDoF coarse-grain over subsets of $\mathcal{S}$ that carry information relevant for predicting a computation’s performance. That is, the states that $\mathcal{M}$ coarse-grains over are hidden in that they contain dynamically relevant information not determined from instantaneous realizations of the memory. The resulting memory dynamics are non-Markovian, since information is transmitted from past to future without ever appearing in the present memory state [17]. The sobering fact is that a general analytical treatment of partially-observed (and therefore non-Markovian) systems is highly nontrivial [5, 16, 18–20]. No matter, hidden states allow for more general forms of computation [9, 21], since non-Markov dynamics relax the constraints imposed by CTMCs. Following this argument to its conclusion, the following demonstrates that the appropriate setting for thermodynamic computing is continuous-time hidden Markov chains (CTHMCs), in which hidden variables store computationally-relevant information.

Moreover, when memory is stored in positional degrees of freedom, the conjugate momentum variables are particularly useful hidden variables for flexibly designing computations. We demonstrate this first by implementing a thermodynamically-costless bit flip—a simple computation that is explicitly forbidden by CTMCs. We then generalize this to a costless Fredkin gate—a key component in reversible computing that is also impossible to implement with CTMCs. This operation is computation universal (Turing complete), meaning that combinations of the Fredkin gate can implement any logical operation [22]. The implementation of this universal and reversible logic gate via CTHMCs demonstrates that non-Markovian dynamics are essential to thermodynamic computing and that a new class of momentum-based computation is within reach.

**Physically Flipping a Physical Bit** To execute a single bit flip over a time interval $t \in [0, \tau]$, the first step is to store a bit of information. One candidate is a particle with a single position dimension $x \in \mathbb{R}$ and corresponding momentum $p \in \mathbb{R}$ in an even potential energy landscape $V^{\text{store}}(x)$ containing two potential minima at $x = \pm x_0$ with an associated energy barrier between them equal to $\max\{V^{\text{store}}(x), x \in (-x_0, x_0)\}$. The particle’s environment is a thermal bath at temperature $T$. As the height of the potential energy barrier rises relative to the bath energy scale $k_B T$, the probability that the particle transitions between left ($x < 0$) and right ($x \geq 0$) decreases exponentially. In this way, if we assign the left half of the position space to memory state 0 and the right half to memory state 1, the energy landscape is capable of metastably storing a bit $m \in \{0, 1\}$.

To execute a flip operation, we instantaneously reduce the coupling to the thermal reservoir to zero such that the memory system now follows dissipationless Hamiltonian dynamics. Simultaneously, the potential energy landscape changes to a positive quadratic well: $V^{\text{comp}}(x, t = 0^+) = k x^2 / 2$. The resulting particle motion is harmonic oscillation: $x(t) = x^* \cos \left(t \sqrt{k/m + \phi}\right)$, where $x^*$ is the maximum distance from the cycle’s origin and $\phi$ is the phase difference from maximum distance at the time $t = 0^+$. 
Maintaining the decoupled system in the quadratic potential energy landscape for half the oscillation period $t \in \left(0, \pi \sqrt{m/\sqrt{k}}\right)$, the particle’s new position becomes:

$$x\left(\frac{\pi \sqrt{m}}{\sqrt{k}}\right) = x^* \cos(\pi + \phi) = -x^* \cos(\phi) = -x(0).$$

Thus, over the computation interval $\tau = \pi \sqrt{m/\sqrt{k}}$, the position flipped sign so that the memory state has flipped as well: $m(\tau) = 1 - m(0)$. Finally, we instantaneously return the potential energy landscape to $V^{\text{store}}(x)$ and recouple to the thermal bath.

The work involved is the time-integrated rate of potential energy change due to the change in the protocol parameter [23]:

$$W = \int dt \frac{\partial V(x, t)}{\partial x} \big|_{x(t),t}.$$  

(1)

This implies a work cost for the bit flip given by the sum of the changes in potential energy at $t = 0$ and $t = \tau$:

$$W = W_0 + W_\tau$$

(2)

$$W_t = V(x, y, z, t^+) - V(x, y, z, t^-),$$

(3)

where $t^+$ and $t^-$ are times immediately before and after, respectively, $t = 0$ or $t = \tau$. However, since the particle position simply flips sign overall between $t = 0$ and $t = \tau$ and the potential energy landscape is even, zero net work must be generated during this time-symmetric protocol. Not only does this computation go beyond what is physically allowable according to rate-equation dynamics over the memory states, but the states only change while the Hamiltonian control is fixed. Thus, the computation is passive, meaning that it fits the information-ratchet framework introduced by Ref. [24].

**A Physical Fredkin Gate**

The bit-flip implementation may seem obvious in its simplicity. However, sophisticated and functional computing can be built from a similar passive processes. Below we outline an implementation of the Fredkin gate, a reversible and computation universal logical gate [22], using the same strategy. This straightforwardly establishes that CTHMCs give straightforward access to complex and universal Turing thermodynamic computing.

The **Fredkin gate** operates on three bits $\mathcal{M} = \{0, 1\}^3$. That is, we encode the physical substrate as three particle-position variables $(x, y, z)$ that are each separated into negative and positive memory-state regions, as above. This splits the memory states into eight respective octants: $(x < 0, y < 0, z < 0)$ corresponds memory state $m = 000$, $(x < 0, y \geq 0, z < 0)$ to $m = 010$, and so on. The information-storing Hamiltonian is a straightforward sum of bistable, even, one-dimensional storage potentials:

$$V^{\text{store}}(x, y, z) = V^{\text{store}}(x) + V^{\text{store}}(y) + V^{\text{store}}(z).$$

This provides metastable regions corresponding to each memory state $m_x m_y m_z \in \{0, 1\}^3$.

Given this construction, we design physical transformations that implement the Fredkin gate with zero cost in finite time. The Fredkin gate is also known as the **controlled swap gate**, as it exchanges inputs $m_x$ and $m_z$ only if the control $m_y$ is set to 1. In other words, the gate maps all inputs to themselves, excluding 101 and 110 that swap with each other. The implementation uses the bit-flip strategy of decoupling and adding a harmonic potential over the time interval $t \in (0, \tau)$, then recoupling and resetting the original information-storing Hamiltonian. The only difference is that the harmonic potential driving the computation is now embedded in the higher-dimensional space.

To execute the Fredkin gate, first note that the memory-state $x$-index must always be fixed: $m_x(\tau) = m_x(0)$. Moreover, behavior in the $y$–$z$ plane should only depend on $x$ up to whether it is positive or negative. Thus, we first split the potential into two pieces: $V(x, y, z, t) = V^{\text{store}}(x) + V^y(z, y, z, t)$. If $m_y(0) = 0$ then $m_y$ and $m_z$ must also not change. This suggests using the information-storing potential for this region of state space: $V(x < 0, y, z, t) = V^{\text{store}}(x, y, z)$ during the entire computation. For $m_x = 1$, however, we must nontrivially compute on $m_y$ and $m_z$:

$$V^y(x \geq 0, y, z, t \in (0, \tau)) = V^{\text{comp}}(y, z).$$

Here, $V^{\text{comp}}$ determines that part of the Hamiltonian which implements the switch $101 \rightarrow 110$ and $110 \rightarrow 101$ and remains unchanging over $t \in (0, \tau)$. Due to decoupling from the $x$-axis, particle behavior in either the positive or negative $x$ regions can be considered as being purely the result of two-dimensional dynamics.

To swap 101 and 110, while keeping 111 and 100 fixed, consider a new basis for the $yz$-space. Define new variables: $y' = (y - z)/\sqrt{2}$ and $z' = (y + z)/\sqrt{2}$, such that the local equilibrium distributions for states 110 and 101 are centered around $z' = 0$ and those for states 111 and 100 are centered around $y' = 0$. Thus, our goal
We choose the $x$ when waiting half a period $z$ the $y$ contributions from the subspace and remains constant. Thus, there are no work positive, then the work invested also vanishes.

Double-well potential, so instantaneous change, as the system is held in the same single sign for much higher than the vast majority of thermal fluctuations change sign, because the energy barrier between states is only comes from the initial and final instantaneous components of the Fredkin gate.

Their respective potential minima. Thus, the transform $V(x,y,z) = V^{comp}(y,z) = V(y) + V(z')$. Flipping in the $y'$-coordinate employs the same Hamiltonian as for the previous bit-flip protocol: $V(y') = ky'^2/2$. As a result, when waiting half a period $\tau = \pi \sqrt{m/\kappa}$, the $y'$ coordinate changes sign $y'\left(\tau\right) = -y'(0)$, as does its momentum.

We choose the $z'$ coordinate’s potential to be quadratic as well, but with an induced period of oscillation that is half as long: $V(z') = 2kz'^2$. This then undergoes a full cycle after the duration $\tau = \pi \sqrt{m/\kappa}$, returning to its original value $z'(\tau) = z(0)$, as does its momentum.

The resulting full Hamiltonian over the control interval operates piecewise. Figure 1 shows the potential in the $x < 0$ and $x > 0$ regions during the computation interval:

\[
V(x, y', z', t) = \begin{cases} 
V^{store}(x) + V^{yz}(x, y', z', t) & \text{if } x < 0 \\
V^{store}(x) + k \frac{y'^2}{2} + 2kz'^2 & \text{if } x \geq 0
\end{cases}
\]

for $t \in (0, \tau)$. Translating back to the original coordinates $y = (y' + z')/\sqrt{2}$ and $z = (z' - y')/\sqrt{2}$, we find that for $x \geq 0$, this passive Hamiltonian transforms the particle’s state by swapping $y$ and $z$:

\[
(y(\tau), z(\tau)) = \begin{pmatrix} y'(\tau) + z'(\tau) \sqrt{\frac{2}{k}} \\
-\frac{y'(0) + z'(0)}{\sqrt{2}} \end{pmatrix} = \begin{pmatrix} -y'(0) + z'(0) \sqrt{\frac{2}{k}} \\
\frac{y'(0) + z'(0)}{\sqrt{2}} \end{pmatrix}
\]

while it holds the other four quadrants where $m_x = 0$ in their respective potential minima. Thus, the transformation swapped $y$ and $z$ only when $m_x = 1$, implementing the Fredkin gate.

For a particular trajectory $(x, y, z)(t)$, the work invested only comes from the initial and final instantaneous changes in the energy landscape:

\[
W = V(x(0), y(0), z(0), 0^+) - V(x(0), y(0), z(0), 0^-) + V(x(\tau), y(\tau), z(\tau), \tau^-) - V(x(\tau), y(\tau), z(\tau), \tau^-).
\]

Recall the restriction that $x(t)$ is exponentially unlikely to change sign, because the energy barrier between states is much higher than the vast majority of thermal fluctuations can access. Thus, we assume that paths maintain a single sign for $x(t)$. If $x(t)$ is negative, then there is no instantaneous change, as the system is held in the same double-well potential, so $W = 0$. That said, if $x(t)$ is positive, then the work invested also vanishes.

The $z$ subspace of the potential decouples from the $y - z$ subspace and remains constant. Thus, there are no work contributions from the $x$-dependent terms. Additionally, the $y - z$ subspace potential is symmetric with respect to exchange of the $y$ and $z$ coordinates. So, the energy differences above will vanish for the $y$ and $z$ dependent terms as well. (Recall that the action of the potential over our interval is to swap the $y$ and $z$ coordinates so that $(y(\tau), z(\tau)) = ((z(0), y(0)))$. And so, the average work production is nearly zero—only the exponentially suppressed barrier crossing events can contribute work.

Figure 2 demonstrates the evolution of the phase space on an ensemble of initial conditions drawn from the equilibrium distribution of $V_{\text{store}}(x, y, z)$. As shown by the particle coloring, those that start in 110 and 101 swap while all others are fixed. Moreover, none of the particles’ $x$-coordinates change informationally—confirming the effectiveness of the overall transformation.

Langevin Simulation The preceding stipulated that the logical system be isolated from its thermal environment during the swap. It might not seem surprising then, that we are able to accomplish a work-free bit flip, given that other classical implementations of efficient reversible computing—such as ballistic computing with billiards—necessarily operate in a dissipationless environment. However, a key and somewhat surprising point is that the Fredkin gate outlined above tolerates imperfect isolation from its thermal environment. The gate’s robustness to fluctuations separates it from other implementations that are dynamically unstable, such as billiard computing.

To demonstrate this, we investigated how robust the operation is to thermal agitation by using underdamped Langevin dynamics. A simulation was carried out by initializing particles in the equilibrium distribution with a thermal reservoir under a quartic information-storing potential $V^{\text{store}}(x, y, z)$. Next, as described above, we exert work on the system by turning on the computational potential $V^{\text{comp}}$ in the region $x > 0$. However, rather than reducing the thermal coupling to $\lambda = 0$, we drop the coupling coefficient to a nonzero value in the weak coupling regime. This coupling value and potential are held fixed for time $\tau = \pi \sqrt{m/\kappa}$. (The Appendix provides additional detail.)

Thermodynamically Robust Fredkin Gate The particles experience thermal fluctuations as the weak coupling to the bath perturb their trajectories from the otherwise expected harmonic motion. The work gained from shutting off the potential will not generally be the same as the work invested to turn it on (as in the idealized case of zero thermal coupling). In fact, the Second Law guarantees that, generally, positive work is invested for such cyclical transformations, because the net change in equilibrium free energy is zero. Nevertheless, one expects the behavior to approximate the desired Fredkin-gate dynamics if the coupling is sufficiently weak. While the energetic cost of implementing the gate does not remain zero as the coupling approaches zero, Fig. 3 shows that the logical fidelity approaches unity. And, it does so with zero slope, revealing that this Fredkin gate implementation is robust
As a final note, the thermodynamic cost is much more than that predicted by the microscopic detailed-balance dynamics that underlie the Langevin simulation. This suggests the existence of a lower bound on entropy production—one that accounts for the course-graining, as predicted in Ref. [26].

**Conclusion**  Rate-equation dynamics is certainly a venerable and powerful framework, central to reaction kinetics in chemistry [27, 28] and key to the master equations of applied statistical mechanics [2, 10, 11]. Due to the remarkable successes of continuous-time Markov chain predictions of many thermodynamic behaviors, it might seem natural to claim that to be “physically realizable”, thermodynamic computing and biological information processing should only be described and analyzed as rate-equation dynamics [8].

The results here demonstrated that this does not hold generally. And so, it cannot form a complete basis for
thermodynamic computing. Moreover, it levies a heavy penalty, precluding engineering and analyzing Maxwellian information ratchets, which are the physical equivalent of Turing machines [24, 29–31]. The limits are especially draconian, since efficient time-symmetrically controlled general computations consist of involutions [26]—operations that are composed of bit swaps and identity maps in positional memory.

As a constructive alternative, we proposed employing continuous-time hidden Markov chains to realize non-Markovian momentum computing. We demonstrated it provides a more complete framework, using two explicit examples that are forbidden if one is restricted to rate equations to describe the evolution between memory states [8]. Additionally, we introduced explicit mechanisms for implementing both in finite time with zero work, proving them “physically realizable”. However, we did fully acknowledge the increased analytical complexity posed by CTHMC dynamics. Fortunately, requisite tools have been developed that render the behaviors analytically tractable and in closed form [32, 33].

Given that convincing, physically-realizable implementations of the bit flip and Fredkin gate [22, 34, 35] have been known for some time, one can only conclude that computing devices already operate beyond the restrictions imposed by rate-equation dynamics. The examples presented here were intentionally couched in the thermodynamics of information to help bridge an apparent gap in understanding general computing. Most specifically, to fully realize the power and breadth of thermodynamic computing, the conception of memory must be expanded from being the realization of a microscopic physical state to being a mesoscopic coarse-graining, as Landauer emphasized half a century ago. Thus, CTHMCs and the momentum-based computing paradigm they inspire are invaluable tools, required for even the most basic computational tasks.

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Langevin Simulations

We simulated a system undergoing Langevin equations of motion:

\[
dq = v_q dt \\
m \, dv_q = -\lambda v_q dt - \partial_q V dt + \sqrt{2k_B T} \lambda r(t) \sqrt{dt},
\]

over three position coordinates \( q = x, y, \) or \( z \). Here, \( v_q \) is the corresponding velocity, \( m \) is the mass, \( \lambda \) is the damping coefficient, and \( r(t) \) is a memoryless Gaussian random variable with zero mean and unit variance. The one-dimensional storage potential \( V_{\text{store}}(q) = \alpha q^4 - \beta q^2 \) with coefficients \( \alpha \) and \( \beta \).

To simulate the above system, we first nondimensionalized the equations of motion:

\[
\tilde{d}q = \tilde{v}_q d\tilde{t} \\
\tilde{d}\tilde{v}_q = -\gamma \tilde{v}_q d\tilde{t} + \partial_q \tilde{V} d\tilde{t} + \sqrt{2\eta} r(\tilde{t}) \sqrt{d\tilde{t}} \\
V_{\text{store}}(\tilde{q}) = \tilde{\alpha} q^4 - \tilde{\beta} q^2 \\
V_{\text{comp}}(\tilde{y}', \tilde{z}') = \frac{\tilde{k}}{2} y'^2 + 2\tilde{k} z'^2,
\]

where \( \tilde{\cdot} \) denotes a nondimensional quantity, and \( \gamma \) and \( \eta \) are two additional nondimensional parameters.

We conducted simulations over a range of values for \( \gamma \in \{0.0, 0.5\} \), kept \( \eta = \sqrt{\gamma} \), and held the remainder of the nondimensional parameters fixed: \( \tilde{\tau} = \pi, \tilde{\alpha} = 2, \tilde{\beta} = 16, \) and \( \tilde{k} = 1 \), where \( \tilde{\tau} \) is the nondimensional duration of the computation interval.

These choices are equivalent to four relationships between the dimensional parameters. The following three equalities held for all simulations: (i) \( \tau = \pi \sqrt{m/k} \), (ii) \( w = 2\sqrt{k_B T/k} \), and (iii) \( h = 32k_B T \), where \( \tau \) is the dimensional duration of the computation interval, \( w = \sqrt{\beta/2\alpha} \) is the positional distance from the central maximum to the minima in the one-dimensional storage potential \( V_{\text{store}} \), and \( h = \beta^2/4\alpha \) is the energy difference between those points. The final dimensional relationship is an effective sweeping of the dimensional damping coefficient \( \lambda \) in \( 0 \leq \lambda \leq \frac{1}{16} \left( \frac{m}{\tau^2} \right) \).

The simulation employed the fourth-order Runge-Kutta method for the deterministic portion and Euler’s method for the stochastic portion of the integration. (Python NumPy’s Gaussian number generator was used to generate the memoryless Gaussian variable \( r(t) \)).

Figure 3’s plot displays 3\( \sigma \) error bars. The errors, though, are sufficiently small that they do not show up appreciably. Statistical errors were estimated using standard procedures for sample means and proportions.

Figure 3 was generated from simulation using the following procedure. First, an ensemble of 20000 trials were chosen from an approximate equilibrium distribution of \( V_{\text{store}}(x, y, z) \) using the Monte Carlo algorithm. Second, this ensemble was thermalized while coupled to a bath (\( \lambda = m/\tau \)) until the ensemble energy changed by no more than 1 part in 1000 over a unit time interval. Third, this ensemble was then used as the start state for the Fredkin gate operation. We then dropped \( \lambda \) down to a low coupling value and exposed the unit mass particles to the potential in Eq. (4).

Fourth, we measured the work required to change the potential across our ensemble. Fifth, the potential was then held fixed for the computation duration \( \tau \) using an integration step \( dt \approx 0.0005\tau/\pi \). Finally, immediately following the computation interval, we measured the second work contribution—the work that would be harvested by dropping the potential back to \( V_{\text{store}} \). The average net work is the ensemble average difference between the work invested when raising the potential and the work harvested when lowering it.

Figure 2 was generated by starting the particles in the equilibrium distribution described above, and running the simulation with \( \lambda = 0 \), to simulate dissipationless oscillatory dynamics. For clarity, the plot shows a sample of 200 trials, rather than the full 20,000.