Minimized state complexity of quantum-encoded cryptic processes

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(Received 28 October 2015; published 16 May 2016)

The predictive information required for proper trajectory sampling of a stochastic process can be more efficiently transmitted via a quantum channel than a classical one. This recent discovery allows quantum information processing to drastically reduce the memory necessary to simulate complex classical stochastic processes. It also points to a new perspective on the intrinsic complexity that nature must employ in generating the processes we observe. The quantum advantage increases with codeword length: the length of process sequences used in constructing the quantum communication scheme. In analogy with the classical complexity measure, statistical complexity, we use this reduced communication cost as an entropic measure of state complexity in the quantum representation. Previously difficult to compute, the quantum advantage is expressed here in closed form using spectral decomposition. This allows for efficient numerical computation of the quantum-reduced state complexity at all encoding lengths, including infinite. Additionally, it makes clear how finite-codeword reduction in state complexity is controlled by the classical process's cryptic order, and it allows asymptotic analysis of infinite-cryptic-order processes.

DOI: 10.1103/PhysRevA.93.052317

I. INTRODUCTION

To efficiently synchronize predictions of a given process over a classical communication channel two observers, call them Alice and Bob, must know the process' internal structure and communicate the relevant history. In particular, leveraging common knowledge of the process' dynamic, what is the minimal amount of information that Alice must communicate to Bob so that he can make the same probabilistic prediction as Alice? The answer is given by the process' internal state information or *statistical complexity* C_{μ} [1].

A closely related question immediately suggests itself: is it more efficient to synchronize via a quantum communication channel that transmits qubits instead of bits? Extending early answers [2,3], a sequence of constructions (*q-machines*) was recently introduced that offers substantial message-size reduction below C_{μ} [4]. In these constructions, each codeword length L yields a quantum communication cost $C_q(L) \leqslant C_{\mu}$ that decreases with increasing L. Moreover, the maximum-compression complexity, $C_q(\infty) = C_q(k)$, is achieved at a codeword length called the *cryptic order k* [5,6]: a recently discovered classical, topological property that is a cousin to the Markov order familiar from stochastic process theory.

Reference [4] pointed out that the new efficiency in synchronization comes with a tradeoff. Bob can only make predictions that are more specialized than Alice's: those consistent with Alice's but also consistent with a probabilistically generated extension of the codewords Alice uses to construct the qubits she sends. These constraints lead to a seemingly odd way for Alice and Bob to synchronize, but there is no way around this. The constraints of this tradeoff are more apparent if we consider the related scenario of "Alice Past" synchronizing

to "Alice Future" (aka Bob) as she generates a realization of the process and updates her state. To *generate* a process the *future* possibilities must be synchronized with the *past* in just such a way that information shared between past and future is channeled through the *present* without violating the process' causality or time order.

One consequence is that the quantum communication cost $C_q(L)$ demands a more refined interpretation: it is the average state information that must be remembered to *generate* the process. Another is that $C_q(L)$ decreases with increasing L since codewords merge, yielding increasingly coincident predictions. The conclusion is that a process's correlational structure controls the degree of quantum compression.

There are both theoretical and practical implications. On the one hand, the theory of minimized quantum-state complexity greatly broadens our notions of the structural complexity inherent in processes; for example, allowing us to quantitatively compare classical- and quantum-state memories [7]. In an applied setting, on the other, it identifies significantly reduced memory requirements for simulating complex classical stochastic processes via a quantum device.

Reduced memory requirements for stochastic simulation were recognized previously for Markov order-1 processes, whose quantum advantage saturates at $C_q(1)$ [2]. For example, it was shown that the classical nearest-neighbor one-dimensional Ising model has a less complex quantum representation [8]. Recently, the quantum advantage of reduced state complexity was experimentally demonstrated for a simple Markovian dynamic [9].

The increasing quantum advantage discovered in Ref. [4], as encapsulated by $C_q(L)$, was challenging to calculate, analytically and numerically. This was unfortunate since for most complex processes, the optimal state complexity reduction is only achieved asymptotically as codeword length $L \to \infty$. Moreover, without a comprehensive theory, few conclusions could be rigorously drawn about $C_q(L)$'s convergence and limits. The following removes the roadblocks. It delivers closed-form expressions, yielding both numerical efficiency and analytic insight.

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Our first contribution is the introduction of the *quantum* pairwise-merger machine (QPMM). The QPMM contains, in a compact form, all of the information required for efficient calculation of the signal-state overlaps used in the q-machine encoding. In particular, we derive closed-form expressions for overlaps in terms of the QPMM's spectrum and projection operators.

This leads to our second contribution: a decomposition of the quantum state complexity $C_q(L)$ into two qualitatively distinct parts. The first part is present for codeword lengths only up to a finite-horizon equal to the index of the QPMM dynamic which, for the case of finite cryptic order, is equal to the process' cryptic order. This provides a nearly complete understanding of $C_q(L)$ for finite-cryptic-order processes. The second part asymptotically decays with an infinite-horizon and is present only in infinite-cryptic order processes. Moreover, we show that $C_q(L)$ oscillates under an exponentially decaying envelope and explain the relevant rates and frequencies in terms of the QPMM's spectral decomposition.

Our third contribution comes in analyzing how computing $C_q(L)$ requires efficiently manipulating quantum-state overlaps. The technique for this presented in Ref. [4] required constructing a new density matrix that respects overlaps. However, it is known that overlaps may be monitored much more directly via a Gram matrix. Here we adapt this to improve calculational efficiency and theoretical simplicity, and we improve matters further by introducing a new form of the Gram matrix.

Our final contribution follows from casting $C_q(L)$'s calculation in its spectral form. This has the distinct advantage that the limit of the overlaps, and thus $C_q(\infty)$, can be calculated analytically. Illustrative examples are placed throughout to ground the development.

II. TWO REPRESENTATIONS OF A PROCESS

The objects of interest are discrete-valued, stationary, stochastic processes. A *process* consists of a bi-infinite sequence $X_{-\infty:\infty} = \dots X_{-2}X_{-1}X_0X_1X_2\dots$ of random variables X_t that take on one or another value in a discrete alphabet: $x_t \in \mathcal{A}$. For each time t and subsequent contiguous block length L, a process assigns a particular probability $\Pr(w)$ to each length-L word $w = x_t \dots x_{t+L-1}$. For *stationary* processes, these probabilities are independent of t. A stationary process' *language* is that set of words $w = x_0 \dots x_{L-1}$ of any length L generated with positive probability.

In particular, we consider processes that can be generated by finite hidden Markov models (HMMs). For edge-output HMMs (i.e., Mealy HMMs), introduced more formally in Appendix A, the observed symbol is generated on transitions between states.

We next consider two representations of a given process, first a canonical classical representation and then a newer quantum representation. Each utilizes the concept of a process' causal states, which are equivalence classes of histories that yield the same conditional probability distributions over future trajectories. Specifically, causal states are the equivalence classes induced by the predictive equivalence relation \sim_{ϵ}

applied to observable histories $x_{-\infty:0}$:

$$\overleftarrow{x} \sim_{\epsilon} \overleftarrow{x'} \Leftrightarrow \operatorname{Pr}(X_{0:L}|X_{-\infty:0} = \overleftarrow{x}) = \operatorname{Pr}(X_{0:L}|X_{-\infty:0} = \overleftarrow{x'}),$$

for all $L \in \{1,2,\ldots\}$. Said another way, causal states are the minimal sufficient statistic of the *past* $X_{-\infty:0}$ for predicting the *future* $X_{0:\infty}$. (We use indexing $X_{a:b}$ that is left inclusive, but right exclusive.)

A. ϵ -Machine: optimal, minimal predictor

While a given process generally has many alternative HMM representations, there exists a unique, canonical form: the process's ϵ -machine [1], which is a process's minimal optimal predictor. Causal states, which are by definition predictive equivalence classes of histories, are the latent states of the ϵ -machine.

Definition 1. A process's ϵ -machine is the 4-tuple $\{S, A, \{T^{(x)}\}_{x \in A}, \pi\}$, where S is the set $\{\sigma_0, \sigma_1, \ldots\}$ of the process' causal states, A is the set of output symbols x, $\{T^{(x)}: T_{i,j}^{(x)} = \Pr(\sigma_j, x | \sigma_i)\}_{x \in A}$ consists of the symbol-labeled state transition matrices, and π is the stationary distribution over states.

The probability that a word $w = x_0 x_1 \dots x_{L-1}$ is generated by an ϵ -machine is given in terms of the labeled transition matrices and the initial state distribution:

$$Pr(w) = \pi \prod_{i=0}^{L-1} T^{(x_i)} \mathbf{1},$$

where $\mathbf{1} = [1, \dots, 1]^{\mathsf{T}}$. These probabilities are constructed to agree with those of the words in a given process language.

The ensemble temporal evolution of internal state probability $\mu = (\mu_0, \dots, \mu_{|S|-1})$, with $\mu_i = \Pr(\sigma_i)$, is given by

$$\mu(t+1) = \mu(t)T,$$

where the transition matrix T is the sum over all output symbols:

$$T \equiv \sum_{x \in A} T^{(x)}.$$

Transition probabilities are normalized. That is, the transition matrix T is *row-stochastic*:

$$\sum_{j=1}^{|\mathcal{S}|} T_{i,j} = \sum_{j=1}^{|\mathcal{S}|} \sum_{x \in \mathcal{A}} \Pr(\sigma_j, x | \sigma_i) = 1.$$

Its component matrices $T_{ij}^{(x)}$ are said to be *substochastic*. Under suitable conditions on the transition matrix, $\lim_{t\to\infty} \mu(t) = \pi$.

Unifilarity, a property derived from the ϵ -machine equivalence relation [1], means that for each state σ_i , each symbol x may lead to at most one successor state σ_j [10]. In terms of the labeled transition matrices, for each row i and each symbol x the row $T_{ij}^{(x)}$ has at most one nonzero entry. We also will have occasion to speak of a *counifilar* HMM, which is the analogous requirement of unique labeling on transitions coming *into* each state.

One of the most important informational properties of a process, directly calculable from its ϵ -machine, is its statistical

complexity C_{μ} [1]. Used in a variety of contexts, it quantifies the size of a process's minimal description.

Definition 2. A process's statistical complexity C_{μ} is the Shannon entropy of the stationary distribution over its causal states:

$$C_{\mu} = \mathbf{H}(\boldsymbol{\pi})$$

$$= -\sum_{i=1}^{|\mathcal{S}|} \pi_i \log_2 \pi_i.$$

The statistical complexity has several operational meanings. For example, it is the average amount of information one gains upon learning a process' current causal state. It is also the minimal amount of information about the past that must be stored to predict the future as well as could be predicted if the entire past were stored. Most pertinent to our purposes, though, it also quantifies the communication cost of synchronizing two predicting agents through a classical channel [4].

B. q-Machine

The *q-machine* is a quantum representation of a classical stochastic process. Introduced in Ref. [4], it offers the largest reduction in state complexity known so far among quantum models capable of generating classical processes.

A process' q-machine is constructed by first selecting a codeword length L. The q-machine (at L) consists of a set $\{|\eta_i(L)\rangle\}_{i=1}^{|\mathcal{S}|}$ of pure quantum signal states that are in one-to-one correspondence with the classical causal states $\sigma_i \in \mathcal{S}$. Each signal state $|\eta_i(L)\rangle$ encodes the set of length-L words $\{w : \Pr(w|\sigma_i) > 0\}$ that may follow causal state σ_i , as well as the corresponding conditional probability:

$$|\eta_i(L)\rangle \equiv \sum_{w \in \mathcal{A}^L} \sum_{\sigma_j \in \mathcal{S}} \sqrt{\Pr(w, \sigma_j | \sigma_i)} |w\rangle |\sigma_j\rangle,$$
 (1)

where $\{|w\rangle\}_{w\in\mathcal{A}^L}$ denotes an orthonormal basis in the "word" Hilbert space with one dimension for each possible word w of length L. Similarly, $\{|\sigma_j\rangle\}_{j=1}^{|\mathcal{S}|}$ denotes an orthonormal basis in the "state" Hilbert space with one dimension for each classical causal state. The ensemble of length-L quantum signal states is then described by the density matrix:

$$\rho(L) = \sum_{i=1}^{|\mathcal{S}|} \pi_i |\eta_i(L)\rangle \langle \eta_i(L)|. \tag{2}$$

The ensemble's von Neumann entropy is defined in terms of its density matrix: $S(\rho) = -\text{tr}[\rho \log_2(\rho)]$, where $\text{tr}[\cdot]$ is the trace of its argument. Paralleling the classical statistical complexity, the quantity

$$C_q(L) \equiv S(\rho(L))$$

$$= -\text{tr}[\rho(L)\log_2(\rho(L))]$$
(3)

has the analogous operational meaning of the communication cost to send signal states over a quantum channel. Von Neumann entropy decreases with increasing signal-state overlap. It is generically smaller than the classical cost [4]: $C_q(L) \leq C_\mu$. In fact, $C_\mu = C_q$ if and only if the process' ϵ -machine is counifilar: there are no states with (at least) two similarly labeled incoming edges [2]. Notably, as we

increase state number, processes with counifilar ϵ -machines represent a vanishing proportion of all possible processes [11]. The consequence is that almost all classical processes can be more compactly represented using quantum mechanics. This presents an opportunity to use quantum encoding to more efficiently represent processes.

Quantifying a process' quantum-reduced state complexity via the von Neumann entropy of Eq. (3) is rooted in the existence of optimal quantum compression algorithms, such as Schumacher compression [12]. The advantage of smaller state complexity with larger L, though, is not a consequence of the well developed theory of quantum compression. Rather it derives from carefully harnessing a model's coincident predictions by constructing a process's nonorthogonal quantum signal states. This is a new kind of quantum information processing. Notably, this quantum reduction, of requisite state memory in the simulation of a classical stochastic process, was recently experimentally verified using nonorthogonal photon polarization signal states [9], though only for codeword length L=1. Leveraging both technological and theoretical advancements, the significant reduction in memory requirements quantified by $C_q(L)$ should enable efficient simulation of important complex systems whose dynamics were previously prohibitively memory intensive.

Calculating a process's quantum cost function $C_q(L)$ is challenging, however. The following shows how to circumvent the difficulties. Beyond practical calculational concerns, the theory leads to a deeper appreciation of quantum structural complexity.

III. QUANTUM OVERLAPS

Reference [4] showed that the reduction $C_{\mu} - C_q(L)$ in state complexity is determined by quantum overlaps between signal states in the q-machine. Accordingly, calculation of these overlaps is a primary task. Intuitively, nonorthogonal signal states correspond to causal states that yield "similar" predictions, in a sense to be explained. More rigorously, the overlap between nonorthogonal signal states is determined by words whose causal-state paths merge.

To illustrate, we compute several overlaps for the (R-k)-Golden Mean Process, showing how they depend on L. (See Fig. 1 for its ϵ -machine state transition diagram.) This process was designed to have tuneable Markov order R and cryptic order k; here we choose R=4 and k=3. (Refer to Ref. [11] for more on this process and a detailed discussion of Markov and cryptic orders.)

At length L=0, each signal state is simply the basis state corresponding to its causal state: $|\eta_i(0)\rangle = |\sigma_i\rangle$. Since the ϵ -machine is minimal, there are no overlaps in the state vectors.

At length L=1 codewords, we find the first nontrivial overlap. This corresponds to paths $A \xrightarrow{1} A$ and $G \xrightarrow{1} A$ merging at state A, and we have

$$|\eta_A(1)\rangle = \sqrt{p}|1A\rangle + \sqrt{1-p}|0B\rangle$$
 and $|\eta_G(1)\rangle = |1A\rangle$.

This yields the overlap:

$$\langle \eta_A(1)|\eta_G(1)\rangle = \sqrt{p}.$$

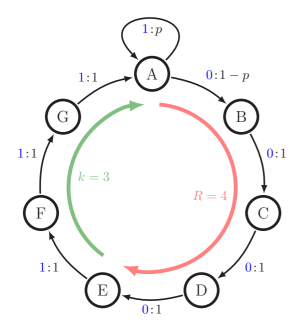


FIG. 1. ϵ -Machine for the (4–3)-Golden Mean Process: The cycle's red segment (labeled R=4) indicates the "Markov" portion, and the green (labeled k=3) the "cryptic" portion. The length scales R and k are tuned by changing the lengths of these two components, respectively. Edges labeled x:p denote taking the state-to-state transition with probability p while emitting symbol $x \in \mathcal{A}$.

Going on to length L=2 codewords, more overlaps arise from mergings of more state paths. The three quantum signal states

$$\begin{split} |\eta_A(2)\rangle &= p|11A\rangle + \sqrt{p(1-p)}|10B\rangle + \sqrt{(1-p)}|00C\rangle, \\ |\eta_F(2)\rangle &= |11A\rangle, \quad \text{and} \\ |\eta_G(2)\rangle &= \sqrt{p}|11A\rangle + \sqrt{1-p}|10B\rangle \end{split}$$

interact to yield the overlaps:

$$\langle \eta_A(2)|\eta_F(2)\rangle = p,$$

 $\langle \eta_F(2)|\eta_G(2)\rangle = \sqrt{p},$ and
 $\langle \eta_A(2)|\eta_G(2)\rangle = p\sqrt{p} + (1-p)\sqrt{p} = \sqrt{p}.$

The overlaps between (A,F) and (F,G) are new. The (A,G) overlap has the same value as that for (F,G); however, its calculation at L=2 involved two terms instead of one. This is because no new merger has occurred; the L=1 merger, effected by symbol 1, was simply propagated forward along two different state paths having prefix 1. There are two redundant paths: $A \xrightarrow{10} B$ overlaps $G \xrightarrow{11} A$ and $A \xrightarrow{11} A$ overlaps $A \xrightarrow{11} A$ overlaps $A \xrightarrow{11} A$. A naive calculation of overlaps must contend with this type of redundancy.

IV. QUANTUM PAIRWISE-MERGER MACHINE

To calculate signal-state overlaps, we introduce the quantum pairwise-merger machine, a transient graph structure that efficiently encapsulates the organization of state paths. As we saw in the example, calculation of overlaps amounts to tracking

state path mergers. It is important that we do this in a systematic manner to avoid redundancies. The new machine does just this.

We begin by first constructing the *pairwise-merger machine* (PMM), previously introduced to compute overlaps [4]. There, probabilities were computed for each word found by scanning through the PMM. This method significantly reduced the number of words from the typically exponentially large number in a process' language and also gave a stopping criterion for PMMs with cycles. This was a vast improvement over naive constructions of the signal-state ensemble (just illustrated) and over von Neumann entropy calculation via diagonalization of an ever-growing Hilbert space.

Appropriately weighting PMM transitions yields the *quantum PMM* (QPMM), which then not only captures which states merge given which words, but also the contribution each merger makes to a quantum overlap. The QPMM has one obvious advantage over the PMM. The *particular* word that produces an overlap is ultimately unimportant; only the *amount* of overlap generated is important. Therefore, summing over symbols in the QPMM to obtain its internal state transitions removes this combinatorial factor. There are additional significant advantages to this matrix-based approach. Appreciating this requires more development.

To build the QPMM from a given process' ϵ -machine:

- (1) Construct the set of (unordered) pairs of (distinct) ϵ -machine states: (σ_i, σ_j) . We call these *pair states*. To this set, add a special state called *SINK* (short for "sink of synchronization") which is the terminal state.
- (2) For each pair state (σ_i, σ_j) and each symbol $x \in \mathcal{A}$, there are three cases to address:
 - (a) If at least one of the two ϵ -machine states σ_i or σ_j has no outgoing transition on symbol x, then do nothing.
 - (b) If both ϵ -machine states σ_i and σ_j have a transition on symbol x to the same state σ_m , then connect pair state (σ_i, σ_j) to SINK with an edge labeled x. This represents a merger.
 - (c) If both ϵ -machine states σ_i and σ_j have a transition on symbol x to two distinct ϵ -machine states σ_m and σ_n where $m \neq n$, then connect pair state (σ_i, σ_j) to pair state (σ_m, σ_n) with an edge labeled x. (There are no further restrictions on m and n.)
- (3) Remove all edges that are not part of a path that leads to SINK.
- (4) Remove all pair states that do not have a path to SINK. This is the PMM. Now, add information about transition probabilities to this topological structure to obtain the QPMM:
- (5) For each pair state (σ_i, σ_j) in the PMM, add to each outgoing edge the weight $\sqrt{\Pr(x|\sigma_i)\Pr(x|\sigma_j)}$, where x is the symbol associated with that edge. Note that two states in QPMM may be connected with multiple edges (for different symbols).

Returning to our example, Fig. 2 gives the QPMM for the (4–3)-Golden Mean Process. Using it, we can easily determine the length at which a contribution is made to a given overlap. We consider codeword lengths $L=1,2,\ldots$ by walking up the QPMM from SINK. For example, pair (A,G) receives a contribution of \sqrt{p} at L=1. Furthermore, (A,G) receives no additional contributions at larger L. Pairs (A,F) and (F,G), though, receive contributions $p=\sqrt{p}\times\sqrt{p}$ and $\sqrt{p}=\sqrt{p}\times 1$ at L=2, respectively.

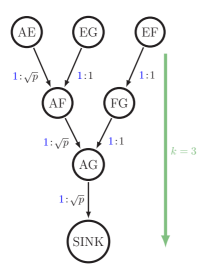


FIG. 2. Quantum pairwise-merger machine for the (4-3)-Golden Mean Process. Its depth is related to the cryptic order k.

The QPMM is *not* a HMM, since the edge weights do not yield a stochastic matrix. However, like a HMM, we can consider its labeled "transition" matrices $\{\zeta^{(x)}\}$, $x \in \mathcal{A}$. Just as for their classical ϵ -machine counterparts, we index these matrices such that $\zeta^{(x)}_{u,v}$ indicates the edge going from pair state u to pair state v. Since the overlap contribution, and not the inducing word, is of interest, the important object is simply the resulting state-to-state substochastic matrix $\zeta = \sum_{x \in \mathcal{A}} \zeta^{(x)}$. The matrix ζ is the heart of our closed-form expressions for quantum coding costs, which follow shortly. As we noted above, it is this step that greatly reduces the combinatorial growth of paths that would otherwise make calculations unwieldy.

To be explicit, our (4–3)-Golden Mean Process has

$$AE \quad EG \quad EF \quad AF \quad FG \quad AG \quad SINK$$

$$AE \quad \begin{pmatrix} 0 & 0 & 0 & \sqrt{p} & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & \sqrt{p} & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ AG \quad SINK \quad \begin{pmatrix} 0 & 0 & 0 & 0 & 0 & 0 & \sqrt{p} \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 \end{pmatrix}$$

V. OVERLAPS FROM THE QPMM

As we saw in the example, overlaps accumulate contributions as "probability amplitude" is pushed through the QPMM down to SINK. Each successive overlap augmentation can thus be expressed in terms of the next iterate of ζ :

$$\langle \eta_i(L)|\eta_i(L)\rangle - \langle \eta_i(L-1)|\eta_i(L-1)\rangle = \langle (\sigma_i,\sigma_i)|\zeta^L|SINK\rangle.$$

The general expression for quantum overlaps follows immediately:

$$\langle \eta_i(L)|\eta_j(L)\rangle = \langle (\sigma_i,\sigma_j)|\sum_{n=0}^L \zeta^n|\text{SINK}\rangle,$$
 (4)

which is true for all processes by design of the QPMM. This form makes clear the cumulative nature of quantum overlaps and the fact that overlap contributions are not labeled.

Note that there are two trivial overlap types. Self-overlaps are always 1; this follows from Eq. (4) since $\langle (\sigma_i, \sigma_i)| = \langle \text{SINK}|$. Overlaps with no corresponding pair state in the QPMM are defined to be zero for all L.

Now, we show that there are two behaviors that contribute to overlaps: a finite-horizon component and an infinite-horizon component. Some processes have only one type or the other, while many have both. We start with the familiar (R-k)-GM, which has only finite-horizon contributions.

A. Finite horizon: (R-k)-Golden Mean Process

Overlap matrices are Hermitian, positive-semidefinite matrices and can therefore be represented as the product $A_L A_L^{\dagger}$. Let us use the general expression Eq. (4) to compute the matrix elements $(A_L A_L^{\dagger})_{i,j} = \langle \eta_i(L) | \eta_j(L) \rangle$ for lengths L=1,2,3,4 for the (R-k)-Golden Mean Process. We highlight in blue (and bold) the matrix elements that have changed from the previous length. All overlaps begin with the identity matrix, here I_7 as we have seven states in the ϵ -machine (Fig. 1). Then, at L=1 we have one overlap. The overlap matrix, with elements $\langle \eta_i(1) | \eta_j(1) \rangle$, is

$$A \quad B \quad C \quad D \quad E \quad F \quad G$$

$$A \quad \begin{pmatrix} 1 & 0 & 0 & 0 & 0 & 0 & \sqrt{\mathbf{p}} \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ E \quad 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ F \quad 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ G \quad \sqrt{\mathbf{p}} \quad 0 & 0 & 0 & 0 & 0 & 1 \end{pmatrix}$$

Next, for L=2 we find two new overlaps. The overlap matrix, with elements $\langle \eta_i(2) | \eta_i(2) \rangle$, is

$$A \quad B \quad C \quad D \quad E \quad F \quad G$$

$$A \quad \begin{pmatrix} 1 & 0 & 0 & 0 & 0 & \mathbf{p} & \sqrt{p} \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ E \quad 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ F \quad \mathbf{p} \quad 0 & 0 & 0 & 0 & 1 & \sqrt{\mathbf{p}} \\ G \quad \sqrt{p} \quad 0 & 0 & 0 & 0 & 0 & \sqrt{\mathbf{p}} \quad 1 \end{pmatrix}$$

For L=3, there are three new overlaps. The overlap matrix, with elements $\langle \eta_i(3) | \eta_j(3) \rangle$, is

Finally, for L=4, we find the same matrix as L=3: $\langle \eta_i(4)|\eta_j(4)\rangle = \langle \eta_i(3)|\eta_j(3)\rangle$ for all i and j. And, in fact, this is true for all $L\geqslant 3$. Therefore, all overlap information has been uncovered at codeword length L=3.

Looking at the QPMM in Fig. 2, we recognize that the saturation of the overlap matrix corresponds to the finite *state depth d* of the directed graph: the number of states in longest path through the QPMM that ends in the SINK state. Equivalently, the depth corresponds to the nilpotency of ζ :

$$d = \min\{n \in \mathbb{N} : \zeta^n = 0\}. \tag{5}$$

Note that the (4-3)-Golden Mean Process QPMM is a tree of state depth d=4.

Whenever the QPMM is a tree or, more generally, a directed-acyclic graph (DAG), the overlaps will similarly have a finite-length horizon equal to the depth d. The nilpotency of ζ for finite-depth DAGs allows for a truncated form of the general overlap expression [Eq. (4)]:

$$\langle \eta_i(L)|\eta_j(L)\rangle = \langle (\sigma_i,\sigma_j)|\sum_{n=0}^{\min(L,d-1)} \zeta^n |\text{SINK}\rangle.$$
 (6)

This form is clearly advantageous for any process whose QPMM is a finite DAG. Naturally then, we are led to ask: What property of a process leads to a finite DAG? To answer this question, we reconsider how overlap is accumulated via the merging of state paths.

Paths through the QPMM represent causal-state path mergers. To make this more precise, we introduce the concept of an *L*-merge, which is most intuitively understood through Fig. 3:

Definition 3. An L-merge consists of a length-L word w and two state paths each of length L+1 that each allow the word w to terminate in the same state F. We denote the word $w = x_0 \dots x_{L-1}$ and state paths (a_0, \dots, a_{L-1}, F) and

FIG. 3. *L*-merge: Two causal state paths, (a_0,\ldots,a_{L-1},F) and (b_0,\ldots,b_{L-1},F) where states $a_i\neq b_i$, for all $i\in\{0,\ldots,L-1\}$, generate the same word $w=x_0\,x_1\ldots x_{L-1}$ and merge only on the last output symbol x_{L-1} into a common final state F.

 $(b_0, \ldots, b_{L-1}, F)$ where states $a_i \neq b_i$, for all $i \in \{0, \ldots, L-1\}$ and, trivially, F = F, the final state in which the paths end.

Immediately, we see that every labeled path of length-L through the QPMM that ends in SINK is precisely an L-merge.

Such causal state path merging not only contributes to quantum overlap, but also contributes to a process's crypticity. Let S_L denote the random variable for the particular causal state $\sigma \in S$ at time L. Then the *crypticity* of a process, the average uncertainty about the present causal state S_0 given perfect observation of the entire infinite future $x_{0:\infty}$, but not knowing the history of observations prior to the present moment, can be written as $H[S_0|X_{0:\infty}]$, which is accumulated at all lengths up to the cryptic order [13].

Definition 4. A process' *cryptic order k* is the minimum length *L* for which $H[S_L|X_{0:\infty}] = 0$.

That is, given knowledge of the entire infinite future of observations, the cryptic order quantifies how far back into the past one must remember to always know the present causal state.

By way of comparison, a process's Markov order is

$$R = \min\{L : \mathbf{H}[\mathcal{S}_L | X_{0:L}] = 0\}.$$

That is, given knowledge (e.g., the ϵ -machine) of which process is being observed but without knowing future observations, the Markov order quantifies how far back into the past one must remember to always know the present causal state. A more familiar length scale characterizing historical correlation, R depends on both path merging and path termination due to disallowed transitions. The cryptic order, in contrast, effectively ignores the termination events and is therefore upper-bounded by the Markov order: $k \leq R$. This bound is also easy to see given the extra conditional variable $X_{L:\infty}$ in the definition of crypticity ($X_{0:\infty} = X_{0:L}X_{L:\infty}$) [5,6].

The following lemma states a helpful relation between cryptic order and L-merges.

Lemma 1. Given an ϵ -machine with cryptic order k: for $L \leq k$, there exists an L-merge; for L > k, there exists no L-merge.

Proof 1. See Appendix B.

Each L-merge corresponds with a real, positive contribution to some quantum overlap. By Lemma 1, for a cryptic-order k process there is at least one L-merge at each length $L \in \{1, \ldots, k\}$ and none beyond k. Therefore, at least one overlap receives a real, positive contribution at each length up until k, where there are no further contributions. This leads to our result for overlap accumulation and saturation in terms of the cryptic order.

Theorem 1. Given a process with cryptic order k, for each $L \in \{0, ..., k\}$, each quantum overlap is a nondecreasing function of L:

$$\langle \eta_i(L+1)|\eta_i(L+1)\rangle \geqslant \langle \eta_i(L)|\eta_i(L)\rangle.$$

Furthermore, for each $L \in \{1, ..., k\}$, there exists at least one overlap that is increased as a result of a corresponding L-merge. For all remaining $L \ge k$, each overlap takes the constant value $\langle \eta_i(k) | \eta_j(k) \rangle$.

Proof 2. See Appendix B.

Evidently, the cryptic order is an important length scale not only for classical processes, but also when building efficient quantum encoders. As an important corollary, this theorem also establishes the relation between a process's cryptic order and the depth of its QPMM:

$$d = k + 1. (7)$$

Thus, we have discovered that the process property corresponding to a finite DAG QPMM is finite cryptic order. Moreover, the cryptic order corresponds to a topological feature of the QPMM, the depth d, responsible for saturation of the overlaps.

This leads to rephrasing the truncated form of the overlaps sum in Eq. (4):

$$\langle \eta_i(L)|\eta_j(L)\rangle = \langle (\sigma_i, \sigma_j)| \sum_{n=0}^{\min(L,k)} \zeta^n |\text{SINK}\rangle.$$
 (8)

This form is advantageous for any process that is finite cryptic order. This, of course, includes all finite Markov-order processes: processes used quite commonly in a variety of disciplines.

Since the quantum-reduced state complexity $C_q(L)$ is a function only of π and quantum overlaps, the preceding development also gives a direct lesson about the $C_q(L)$ saturation.

Corollary 1. $C_q(L)$ has constant value $C_q(k)$ for $L \ge k$.

Proof 1. The entropy of an ensemble of pure signal states $\{p_i, |\psi_i\rangle\}$ is a function of only probabilities p_i and overlaps $\{\langle \psi_i | \psi_i \rangle\}$. The result then follows directly from Theorem 1.

Having established connections among depth, cryptic order, and saturation, we seem to be done analyzing quantum overlap, at least for the finite-cryptic case. To prepare for going beyond finite horizons, however, we should reflect on the spectral origin of the nilpotency of ζ .

A nilpotent matrix, such as ζ in the finite-cryptic case, has only the eigenvalue zero. This can perhaps be most easily seen if the pair states are ordered according to their distance from SINK, so that ζ is triangular with only zeros along the diagonal.

Notably, for finite DAGs with depth d>1, the standard eigenvalue-eigenvector decomposition is insufficient to form a complete basis: the corresponding ζ is necessarily nondiagonalizable due to the geometric multiplicity of the zero eigenvalue being less than its algebraic multiplicity. *Generalized eigenvectors* must be invoked to form a complete basis [14]. Intuitively, this type of nondiagonalizability can be understood as the intrinsic interdependence among pair states in propagating probability amplitude through a branch of the DAG. When ζ is rendered into Jordan block form via a similarity transformation, the size of the largest Jordan block associated with the zero eigenvalue is called the *index* ν_0 of the zero eigenvalue. It turns out to be equal to the depth for finite DAGs.

Summarizing, the finite-horizon case is characterized by several related features: (i) the QPMM is a DAG (of finite depth), (ii) the depth of the QPMM is one greater than the cryptic order, (iii) the matrix ζ has only the eigenvalue zero, and (iv) the depth is equal to the index of this zero eigenvalue, meaning that ζ has at least k generalized eigenvectors. More generally, ζ can have nonzero eigenvalues and this corresponds to richer structure that we explore next.

B. Infinite horizon: Lollipop Process

Now we ask, what happens when the QPMM is not a directed acyclic graph? That is, what happens when it contains *cycles*?

It is clear that the depth d diverges, implying that the cryptic order is infinite. Therefore, the sum in Eq. (4) may no longer be truncated. We also know that infinite-cryptic processes become ubiquitous as ϵ -machine state size increases [11]. Have we lost our calculational efficiencies? No, in fact, there are greater advantages yet to be gained.

We first observe that a QPMM's ζ breaks into two pieces. One has a finite horizon reminiscent of the finite cryptic order just analyzed, and the other has an infinite horizon, but is, as we now show, analytically quite tractable.

In general, a linear operator *A* may be decomposed using the *Dunford decomposition* [15] (also known as the Jordan-Chevalley decomposition) into

$$A = \mathcal{D} + \mathcal{N},\tag{9}$$

where \mathcal{D} is diagonalizable, \mathcal{N} is nilpotent, and \mathcal{D} and \mathcal{N} commute. In the current setting, \mathcal{N} makes the familiar finite-horizon contribution, whereas the new \mathcal{D} term has an infinite horizon: $\mathcal{D}^n \neq 0$, for all $n < \infty$. In the context of infinite cryptic processes, the finite horizon associated with \mathcal{N} is no longer simply related to QPMM depth nor, therefore, the cryptic order which is infinite.

The systematic way to address the new diagonalizable part is via a spectral decomposition [16], where the persistent leaky features of the QPMM state probability evolution are understood as independently acting modes. It is clear that ζ always has a nilpotent component associated with a zero eigenvalue, due to the SINK state. Assuming that the remaining eigenspaces are diagonalizable, the form of the overlaps becomes

$$\langle \eta_{i}(L)|\eta_{j}(L)\rangle = \sum_{\xi \in \Lambda_{\zeta} \setminus \{0\}} \frac{1 - \xi^{L+1}}{1 - \xi} \langle (\sigma_{i}, \sigma_{j})|\zeta_{\xi}|\text{SINK}\rangle$$

$$+ \sum_{m=0}^{\min\{L, \nu_{0}-1\}} \langle (\sigma_{i}, \sigma_{j})|\zeta^{m}\zeta_{0}|\text{SINK}\rangle, \quad (10)$$

where Λ_{ζ} is the set of the eigenvalues of ζ , ζ_{ξ} are the projection operators corresponding to each eigenvalue, and ν_0 is the index of the zero eigenvalue, which is the size of its largest Jordan block. We refer to this as the *almost-diagonalizable* case since all eigenspaces, besides possibly the zero-eigenvalue space, are diagonalizable. This case covers all processes with generic parameters. Here ν_0 is still responsible for the length of the finite-horizon component, but is no longer directly related to QPMM depth or process cryptic order.

Note that in the finite-cryptic order case, the only projector ζ_0 is necessarily the identity. Therefore, Eq. (10) reduces to the previous form in Eq. (8).

The spectral decomposition yields a new level of tractability for the infinite-cryptic case. The infinite-horizon piece makes contributions at all lengths, but in a regular way. This allows for direct calculation of its total contribution at any particular L, including $L \to \infty$.

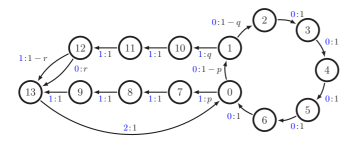


FIG. 4. ϵ -Machine for the (7–4)-Lollipop Process. The cycle of 0s on the right leads to infinite Markov and cryptic orders.

To highlight this behavior, consider the (7-4)-Lollipop Process, whose ϵ -machine is shown in Fig. 4. It is named for the shape of its QPMM; see Fig. 5. This process is a simple example of one where the cryptic order is infinite and the finite-horizon length of the nilpotent contribution is tunable. Roughly speaking, the diagonalizable component comes from the "head" of the lollipop (the cycle), and the nilpotent part comes from the "stick."

It is straightforward to construct the general QPMM and thereby derive ζ for the (N-M)-Lollipop Process. Its QPMM has N pair states in a cyclic head. The M remaining pair states constitute a finite-horizon "stick." We find

$$\det(\zeta - \xi I) = (-\xi)^M [(-\xi)^N - (1-p)(1-q)],$$

yielding

$$\Lambda_{\zeta} = \{0, [(1-p)(1-q)]^{1/N} e^{in2\pi/N} \}_{n=0}^{N-1}, \tag{11}$$

with $v_0 = M$.

For concreteness, consider the (7–4)-Lollipop Process with transition parameters p=q=1/2 and $r\in(0,1)$. It has eigenvalues $\Lambda_{\zeta}=\{0,ae^{in\theta}\}$ and $\nu_0=4$, where $a=(1/4)^{1/7}$, $\theta=2\pi/7$, and $n\in\{0,1,2,3,4,5,6\}$.

Each $\xi = ae^{in\theta}$ eigenvalue has algebraic multiplicity 1 and associated left eigenvector,

$$\langle \xi | = [2\sqrt{2}\xi^6, \sqrt{2}\xi^5, \xi^4, \xi^3, \xi^2, \xi^1, \xi^0, \sqrt{2}\xi^5, \sqrt{2}\xi^4, \sqrt{2}\xi^3, \sqrt{2}(1-r)\xi^2],$$

and right eigenvector,

$$|\xi\rangle = \left[\frac{1}{2\xi}, 1, \sqrt{2\xi}, \sqrt{2\xi^2}, \sqrt{2\xi^3}, \sqrt{2\xi^4}, \sqrt{2\xi^5}, 0, 0, 0, 0\right]^{\top}.$$

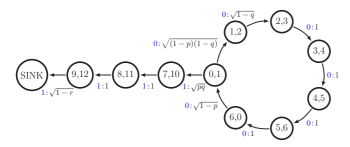


FIG. 5. Quantum pairwise-merger machine for the (7–4)-Lollipop Process.

The general relationship among left and right eigenvectors, left and right generalized eigenvectors, and projection operators, and their reduction in special cases is discussed in Ref. [17]. In the present case, notice that, since ζ is not a normal operator, the right eigenvectors are not simply the conjugate transpose of their left counterparts. (*Normal operators* by definition commute with their conjugate transpose; e.g., Hermitian operators.) The left and right eigenvectors are fundamentally different, with the differences expressing the QPMM's directed causal architecture.

Since each of these eigenvalues has algebraic multiplicity 1, the corresponding projection operators are defined in terms of right and left eigenvectors,

$$\zeta_{\xi} = \frac{|\xi\rangle\langle\xi|}{\langle\xi|\xi\rangle}.$$

The zero eigenvalue has algebraic multiplicity 4 and geometric multiplicity 1, meaning that while there is only one eigenvector there are three generalized eigenvectors. The left and right eigenvectors are

$$[0,0,0,0,0,0,0,0,0,0,1]$$
 and $[0,1,0,0,0,0,0,0,-1,0,0,0]^{\top}$.

The three generalized left eigenvectors are

$$[0,0,0,0,0,0,0,1,0,0,0],$$

 $[0,0,0,0,0,0,0,0,0,1,0,0],$ and
 $[0,0,0,0,0,0,0,0,0,0,1,0];$

and the three generalized right eigenvectors are

$$[0,0,\sqrt{2},0,0,0,0,0,-1,0,0]^{\top},$$

 $[0,0,0,\sqrt{2},0,0,0,0,0,-1,0]^{\top},$ and
 $[0,0,0,0,\sqrt{2(1-r)},0,0,0,0,0,-1]^{\top}.$

Since the index of the zero eigenvalue is larger than 1 (ν_0 = 4), the projection operator ζ_0 for the zero eigenvalue includes the contributions from both its standard and generalized eigenvectors:

$$\zeta_0 = \sum_{n=0}^{3} \frac{|0_n\rangle\langle 0_n|}{\langle 0_n|0_n\rangle},\tag{12}$$

where $|0_0\rangle$ is the standard eigenvector and $|0_n\rangle$ is the *n*th generalized eigenvector for $n \ge 1$. More generally, when the geometric multiplicity is greater than one, this sum goes over all standard and all generalized eigenvectors of the zero eigenvalue.

Since all projection operators must sum to the identity, the projection operator for the zero eigenvalue can be obtained alternatively from

$$\zeta_0 = I - \sum_{\xi \in \Lambda_{\mathcal{E}} \setminus 0} \zeta_{\xi}, \tag{13}$$

which is often useful during calculations.

This very efficient procedure allows us to easily probe the form of quantum advantage for any process described by a finite ϵ -machine.

Finally, we jump directly to the asymptotic overlap using the following expression:

$$\langle \eta_i(\infty) | \eta_j(\infty) \rangle = \langle (\sigma_i, \sigma_j) | \left(\sum_{n=0}^{\infty} \zeta^n \right) | \text{SINK} \rangle$$
$$= \langle (\sigma_i, \sigma_j) | (I - \zeta)^{-1} | \text{SINK} \rangle. \tag{14}$$

Note that $I - \zeta$ is invertible, since ζ is substochastic. Hence, its spectral radius is less than unity, $1 \notin \Lambda_{\zeta}$, and so $\det(1I - \zeta) \neq 0$. Moreover, $(I - \zeta)^{-1}$ is *equal* to the convergent Neumann series $\sum_{n=0}^{\infty} \zeta^n$ by Theorem 3 of Ref. [18, Ch. VIII, Sec. 2].

Yielding an important calculational efficiency, the form of Eq. (14) does not require spectral decomposition of ζ and so immediately provides the asymptotic quantum-reduction of state complexity. Finally, this form does not depend on the previous assumption of ζ being almost-diagonalizable.

VI. QUANTUM REDUCED STATE COMPLEXITY

The preceding development focused on computing overlaps between quantum signal states for q-machine representations of a given process. Let us not forget that the original goal was to compute the von Neumann entropy of this ensemble: the quantum-reduced state complexity $C_q(L)$, which is the memory that must be transferred about the state of the process to synchronize compatible predictions.

The naive approach to calculating $C_q(L)$ constructs the signal states directly and so does not make use of overlap computation. This involves working with a Hilbert space of increasing dimension, exponential in codeword length L. This quickly becomes intractable, for all but the simplest processes.

The second approach, introduced in Ref. [4], made use of the PMM to compute overlaps. These overlaps were then used to construct a density operator with those same overlaps, but in a Hilbert space of fixed size |S|, essentially obviating the high-dimensional embedding of the naive approach. And, we just showed how to calculate overlaps in closed form. The elements of the resulting density matrix, however, are nonlinear functions of the overlaps. Besides the computational burden this entails, it makes it difficult to use the overlap matrix to theoretically infer much about the general behavior of $C_q(L)$.

Here we present two markedly improved approaches that circumvent these barriers. We are ultimately interested in the von Neumann entropy, which depends only on the spectrum of the density operator. It has been pointed out that the Gram matrix of an ensemble shares the same spectrum [19]. The *Gram matrix* for our ensemble of pure quantum signal states is

$$G = \begin{bmatrix} \sqrt{\pi_1 \pi_1} \langle \eta_1 | \eta_1 \rangle & \cdots & \sqrt{\pi_1 \pi_{|\mathcal{S}|}} \langle \eta_1 | \eta_{|\mathcal{S}|} \rangle \\ \vdots & \ddots & \vdots \\ \sqrt{\pi_{|\mathcal{S}|} \pi_1} \langle \eta_{|\mathcal{S}|} | \eta_1 \rangle & \cdots & \sqrt{\pi_{|\mathcal{S}|} \pi_{|\mathcal{S}|}} \langle \eta_{|\mathcal{S}|} | \eta_{|\mathcal{S}|} \rangle \end{bmatrix}.$$
(15)

If we define $D_{\pi} \equiv \mathrm{diag}(\pi)$, then $G = D_{\pi}^{1/2} A A^{\dagger} D_{\pi}^{1/2}$.

Given that it is only a small step from the overlap matrix AA^{\dagger} to the Gram matrix G, we see the usefulness of the thoroughgoing overlap analysis above. The spectrum of G is

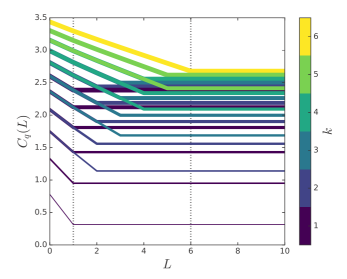


FIG. 6. Quantum costs $C_q(L)$ for the (R-k)-Golden Mean Process with $R \in \{1, \ldots, 6\}$ and $k \in \{1, \ldots, R\}$. R and k are indicated with line width and color, respectively. The probability of the self-loop is p = 0.7. $C_q(L)$ roughly linearly decreases until L = k where it is then constant. Note that (R-k)-GM agrees exactly with ((R+1)-(k-1))-GM for $L \le k$, as explained in Appendix D.

then computed using standard methods, either symbolically or numerically.

There is another surrogate matrix that shares the spectrum but is simpler, yet again, for some calculations. We call this matrix \widetilde{G} the *left-consolidated Gram matrix*:

$$\widetilde{G} = \begin{bmatrix} \pi_1 \langle \eta_1 | \eta_1 \rangle & \cdots & \pi_1 \langle \eta_1 | \eta_{|\mathcal{S}|} \rangle \\ \vdots & \ddots & \vdots \\ \pi_{|\mathcal{S}|} \langle \eta_{|\mathcal{S}|} | \eta_1 \rangle & \cdots & \pi_{|\mathcal{S}|} \langle \eta_{|\mathcal{S}|} | \eta_{|\mathcal{S}|} \rangle \end{bmatrix}.$$
(16)

Note that $\widetilde{G} = D_{\pi} A A^{\dagger}$ —i.e., D_{π} has been consolidated on the left. A right-consolidated Gram matrix would work just as well for the calculation of $C_q(L)$.

Since the spectra are identical, we can calculate $C_q(L)$ directly from the density matrix $\rho(L)$, Gram matrix $G^{(L)}$, or consolidated Gram matrix $\widetilde{G}^{(L)}$:

$$\begin{split} C_q(L) &= -\sum_{\lambda \in \Lambda_{\rho(L)}} \lambda \log_2 \lambda \\ &= -\sum_{\lambda \in \Lambda_{G(L)}} \lambda \log_2 \lambda \\ &= -\sum_{\lambda \in \Lambda_{S(L)}} \lambda \log_2 \lambda. \end{split}$$

For further discussion, see Appendix C.

Using the Gram matrix as described, we illustrate the behavior of $C_q(L)$ for the (R-k)-Golden Mean (Fig. 6) and (N-M)-Lollipop (Fig. 7). For each of the two process families, we compute several instances by varying R and k and by varying N and M while holding fixed their transition parameters. Comparing the two figures, we qualitatively confirm the difference between a process with only a finite-horizon contribution and one with an infinite-horizon contribution. The (R-k)-Golden Mean reaches its encoding saturation at L=k

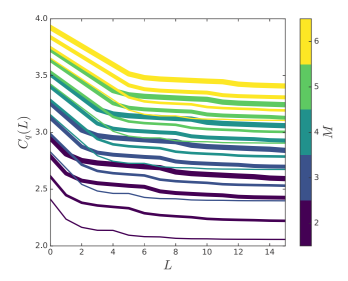


FIG. 7. Quantum costs $C_q(L)$ for the Lollipop process for $N \in$ $\{3,4,5,6\}, M \in \{2,3,4,5,6\}, p = q = 0.5, \text{ and } r = 0.1. N \text{ and } M \text{ are } M = 0.1. N \text{ and } M \text{ are } M = 0.1. N \text{ and } M = 0.1. N \text$ indicated with line width and color, respectively. After a fast initial decrease, these curves approach their asymptotic values more slowly.

the cryptic order. The (N-M)-Lollipop only approaches this limit asymptotically.

In contrast to the customary approach in quantum compression [12], in which an entire message is to be compressed with perfect fidelity, the compression advantage here is obtained by throwing away information that is not relevant for simulating a process, with the goal of correctly sampling from a conditional future distribution.

Recall that the quantum-reduced state complexity $C_q(L)$ quantifies a communication cost. Specifically, it is the amount of memory about a process's state that must be queried to move the system forward in time. However, to avoid misinterpretation, we note that this cost does not have a simple relationship to the "quantum communication cost" as the phrase is sometimes used in the distributed computing setting of communication complexity theory [20].

To supplement the details already given, annotated analytic derivations of several example processes are given in Appendix D. These examples serve as a pedagogical resource, with comparison and discussion of various analytical techniques.

VII. COSTS USING LONG CODEWORDS

The preceding discussed quantum state overlaps extensively. We found that the behavior of the overlaps with L is completely described through the spectral decomposition of ζ . And, we showed that, for any L, the von Neumann entropy $C_a(L)$ can be found from the eigenvalues of the Gram matrix: a direct transformation of the overlap matrix. This is all well and good and key progress. But, can we use this machinery to directly analyze the behavior of $C_q(L)$ as a function of L? For infinite-cryptic processes, the answer is an especially pleasing affirmative.

This section derives the asymptotic behavior of $C_q(L)$ for large L; viz., $v_0 < L \le k = \infty$. We show that a periodic pattern, exponentially decaying at the rate of the largest

 ζ -eigenvalue magnitude, dominates the deviation of $C_q(L)$ from $C_q(\infty)$ for large L. In particular, we show two things: First, the asymptotic behavior of $C_q(L) - C_q(\infty)$ is, to first order, exponentially decreasing as r_1^L , where r_1 is the spectral radius of ζ . Second, this exponential defines an envelope for a Ψ -periodic asymptotic structure, where Ψ is the least common multiple of slowest-decaying QPMM cycle lengths.

Recall that the minimal known upper bound on state complexity is given by the asymptotic von Neumann entropy:

$$C_q(\infty) = -\sum_{\lambda^{(\infty)} \in \Lambda_{G(\infty)}} \lambda^{(\infty)} \log_2(\lambda^{(\infty)}).$$

We will show that when L is large, $(\delta G)^{(L)} \equiv G^{(L)} - G^{(\infty)}$ can be treated as a perturbation to $G^{(\infty)}$. From the corresponding small variations $\{(\delta\lambda)^{(L)}\}_{\lambda\in\Lambda_G}$, direct calculation of the first differential yields the approximate change in the von Neumann entropy:

$$(\delta S)^{(L)} = -\sum_{\lambda \in \Lambda_G} [\log_2(\lambda^{(\infty)}) + 1] (\delta \lambda)^{(L)}, \qquad (17)$$

so long as no zero eigenvalues of $G^{(\infty)}$ prematurely vanish at finite L. Our task, therefore, is to find $(\delta \lambda)^{(L)}$ from $(\delta G)^{(L)}$ in terms of the spectral properties of ζ .

For easy reference, we first highlight our notation:

- (1) $G^{(L)}$ is a Gram matrix at length L corresponding to $\rho(L)$.
 - (2) $\lambda^{(L)} \in \Lambda_{G^{(L)}}$ is any one of its eigenvalues.
- (3) $|\lambda^{(L)}\rangle$ and $\langle\lambda^{(L)}|$ are the right and left eigenvectors of
- $G^{(L)}$ corresponding to $\lambda^{(L)}$, respectively. (4) $(\delta G)^{(L)} \equiv G^{(L)} G^{(\infty)}$ is the perturbation to $G^{(\infty)}$ investigated here.
- (5) $\xi \in \Lambda_{\zeta}$ is an eigenvalue of the QPMM transition dynamic ζ .

If using G's symmetric version, the right and left eigenvectors are simply transposes of each other: $\langle \lambda^{(L)} | = (|\lambda^{(L)} \rangle)^{\top}$. For simplicity of the proofs, we assume nondegeneracy of $G^{(L)}$'s eigenvalues, so that the projection operator associated with $\lambda^{(L)}$ is $|\lambda^{(L)}\rangle\langle\lambda^{(L)}|/\langle\lambda^{(L)}|\lambda^{(L)}\rangle$, where the denominator assures normalization. Nevertheless, the eigenbasis of $G^{(L)}$ is always complete and the final result, Theorem 3, retains general validity.

Here, we show that the matrix elements of $(\delta G)^{(L)}$ are arbitrarily small for large enough L, such that first-order perturbation is appropriate for large L, and give the exact form of $(\delta G)^{(L)}$ for use in the calculation of $(\delta \lambda)^{(L)}$.

Proposition 1. For $L \ge v_0$, the exact change in Gram matrix is

$$(\delta G)^{(L)} = -\sum_{\xi \in \Lambda_{\xi} \setminus 0} \frac{\xi^{L+1}}{1 - \xi} C_{\xi},$$

where C_{ξ} is independent of L and has matrix elements:

$$(C_{\xi})_{i,j} = \sqrt{\pi_i \pi_j} \langle (\sigma_i, \sigma_j) | \zeta_{\xi} | SINK \rangle.$$

Proof 1. We calculate

$$\begin{split} (\delta G)_{i,j}^{(L)} &= G_{i,j}^{(L)} - G_{i,j}^{(\infty)} \\ &= \sqrt{\pi_i \pi_j} \left(\left| \eta_i^{(L)} \right| \eta_j^{(L)} \right) - \left| \eta_i^{(\infty)} \right| \eta_j^{(\infty)} \right) \right) \\ &= -\sqrt{\pi_i \pi_j} \left\langle (\sigma_i, \sigma_j) | \zeta^{L+1} (1 - \zeta)^{-1} | \text{SINK} \right\rangle. \end{split}$$

If we assume that all nonzero eigenvalues of ζ correspond to diagonalizable subspaces, then for $L \geqslant \nu_0$, the elements of $(\delta G)^{(L)}$ have the spectral decomposition:

$$(\delta G)_{i,j}^{(L)} = -\sum_{\xi \in \Lambda_c \setminus 0} \frac{\xi^{L+1}}{1 - \xi} \sqrt{\pi_i \pi_j} \langle (\sigma_i, \sigma_j) | \zeta_{\xi} | \text{SINK} \rangle.$$

Since this decomposition is common to all matrix elements, we can factor out $\left\{\frac{\xi^{L+1}}{1-\xi}\right\}_{\xi}$, leaving the L-independent set of matrices:

$$\{C_{\xi}: (C_{\xi})_{i,j} = \sqrt{\pi_i \pi_j} \langle (\sigma_i, \sigma_j) | \zeta_{\xi} | SINK \rangle \}_{\xi \in \Lambda_{\xi}},$$

such that

$$(\delta G)^{(L)} = -\sum_{\xi \in \Lambda_1 \setminus 0} \frac{\xi^{L+1}}{1 - \xi} C_{\xi}.$$

Proposition 2. At large L, the first-order correction to $\lambda^{(\infty)}$ is

$$(\delta\lambda)^{(L)} = -\sum_{\xi \in \Lambda_{+}\setminus 0} \frac{\xi^{L+1}}{1-\xi} \frac{\langle \lambda^{(\infty)} | C_{\xi} | \lambda^{(\infty)} \rangle}{\langle \lambda^{(\infty)} | \lambda^{(\infty)} \rangle}.$$
(18)

Proof 2. Perturbing $G^{(\infty)}$ to $G^{(\infty)} + (\delta G)^{(L)}$, the first-order change in its eigenvalues is given by

$$(\delta\lambda)^{(L)} = \frac{\langle \lambda^{(\infty)} | (\delta G)^{(L)} | \lambda^{(\infty)} \rangle}{\langle \lambda^{(\infty)} | \lambda^{(\infty)} \rangle}, \tag{19}$$

which is standard first-order nondegenerate perturbation theory familiar in quantum mechanics, with the allowance for unnormalized bras and kets. Proposition 2 then follows directly from Eq. (19) and Proposition 1.

Theorem 2. At large L, such that $v_0 < L \leqslant k = \infty$, the first-order correction to $C_q(\infty)$ is

$$C_{q}(L) - C_{q}(\infty) \approx (\delta S)^{(L)}$$

$$= \sum_{\xi \in \Lambda_{\xi} \setminus 0} \frac{\xi^{L+1}}{1 - \xi} \sum_{\lambda^{(\infty)} \in \Lambda_{G}(\infty)} \langle C_{\xi} \rangle [\log_{2}(\lambda^{(\infty)}) + 1], \quad (20)$$

where

$$\langle C_{\xi} \rangle \equiv rac{\langle \lambda^{(\infty)} | C_{\xi} | \lambda^{(\infty)} \rangle}{\langle \lambda^{(\infty)} | \lambda^{(\infty)} \rangle}.$$

Proof 1. This follows directly from Eq. (17) and Proposition 2.

The large-L behavior of $C_q(L) - C_q(\infty)$ is a sum of decaying complex exponentials. And, to first order, we can even calculate the coefficient of each of these contributions.

Notice that the only L-dependence in Proposition 2 and Theorem 2 comes in the form of exponentiating eigenvalues of the QPMM transition dynamic ζ . For very large L, the dominant structure implied by Proposition 2 and Theorem 2 can be teased out by looking at the relative contributions from the first- and second-largest magnitude sets of eigenvalues of ζ .

Let r_1 be the spectral radius of ζ , shared by the largest eigenvalues $\Lambda(r_1)$: $r_1 \equiv \max\{|\xi| : \xi \in \Lambda_{\zeta}\}$. And let $\Lambda(r_1) \equiv \arg\max\{|\xi| : \xi \in \Lambda_{\zeta}\}$. Then let r_2 be the second-largest magnitude of all of the eigenvalues of ζ that differs from r_1 : $r_2 \equiv \max\{|\xi| : \xi \in \Lambda_{\zeta} \setminus \Lambda(r_1)\}$. And let $\Lambda(r_2) \equiv \arg\max\{|\xi| : \xi \in \Lambda_{\zeta} \setminus \Lambda(r_1)\}$.

 $\Lambda_{\zeta} \setminus \Lambda(r_1)$ }. Multiple eigenvalues can belong to $\Lambda(r_1)$. Similarly, multiple eigenvalues can belong to $\Lambda(r_2)$.

Then, $0 \le (r_2/r_1) < 1$, if ζ has at least one nonzero eigenvalue. This is the case of interest here since we are addressing those infinite-horizon processes with $k = \infty > \nu_0$. Hence, as L becomes large, $(r_2/r_1)^L$ vanishes exponentially, if it is not already zero. This leads to a corollary of Proposition 2.

Corollary 2. For $L \geqslant \nu_0$, the leading deviation from $\lambda^{(\infty)}$ is

$$(\delta \lambda)^{(L)} = -r_1^{L+1} \sum_{\xi \in \Lambda(r_1)} \frac{(\xi/|\xi|)^{L+1}}{1-\xi} \langle C_{\xi} \rangle \left\{ 1 + O\left[\left(\frac{r_2}{r_1} \right)^L \right] \right\}.$$

Notice that $\xi/|\xi|$ lies on the unit circle in the complex plane. Due to their origin in cyclic graph structure, we expect each $\xi \in \Lambda(r_1)$ to have a phase in the complex plane that is a rational fraction of 2π . Hence, there is some n for which $(\xi/|\xi|)^n = 1$, for all $\xi \in \Lambda(r_1)$. The minimal such n, call it Ψ , will be of special importance:

$$\Psi \equiv \min\{n \in \mathbb{N} : (\xi/|\xi|)^n = 1 \text{ for all } \xi \in \Lambda(r_1)\}.$$
 (21)

Since all $\xi \in \Lambda(r_1)$ originate from cycles in ζ 's graph, we have the result that Ψ is equal to the least common multiple of the cycle lengths implicated in $\Lambda(r_1)$.

For example, if all $\xi \in \Lambda(r_1)$ come from the same cycle in the graph of ζ , then $\Psi = |\Lambda(r_1)|$ and

$$\Lambda(r_1) = \{ \xi_m = r_1 e^{im2\pi/|\Lambda(r_1)|} \}_{m=1}^{|\Lambda(r_1)|}.$$

That is, $\{\xi_m/|\xi_m|\}_{m=1}^{|\Lambda(r_1)|}$ are the $|\Lambda(r_1)|^{\text{th}}$ roots of unity, uniformly distributed along the unit circle. If, however, $\Lambda(r_1)$ comes from multiple cycles in the graph of ζ , then the least common multiple of the cycle lengths should be used in place of $|\Lambda(r_1)|$.

Recognizing the Ψ -periodic structure of $(\xi/|\xi|)^n$ yields a more informative corollary of Proposition 2:

Corollary 3. For $L \geqslant \nu_0$, the leading deviation from $\lambda^{(\infty)}$

$$(\delta \lambda)^{(L)} = -r_1^{L+1} \sum_{\xi \in \Lambda(r_1)} \frac{(\xi/|\xi|)^{\mod(L+1,\Psi)}}{1-\xi} \langle C_{\xi} \rangle$$
$$\times \{1 + O[(r_2/r_1)^L]\}.$$

Hence:

$$(\delta\lambda)^{(L+\Psi)} \approx r_1^{\Psi} (\delta\lambda)^{(L)}. \tag{22}$$

We conclude that asymptotically a pattern, of changes in the density-matrix eigenvalues (with period Ψ), decays exponentially with decay rate of r_1^{Ψ} per period. There are immediate implications for the pattern of asymptotic changes in $C_q(L)$ at large L.

Corollary 4. For $L \geqslant \nu_0$, the leading deviation from $C_q(\infty)$ is

$$\begin{split} C_q(L) - C_q(\infty) &\approx (\delta S)^{(L)} \\ &= r_1^{L+1} \sum_{\xi \in \Lambda(r_1)} \frac{(\xi/|\xi|)^{\mod(L+1,\,\Psi)}}{1 - \xi} \\ &\times \sum_{\lambda^{(\infty)} \in \Lambda_{C(\infty)}} \langle C_\xi \rangle \log_2(\lambda^{(\infty)}) \{1 + O[(r_2/r_1)^L]\}. \end{split}$$

The most profound implication of this detailed analysis can be summarized succinctly.

Theorem 3. For sufficiently large L,

$$\frac{C_q(L+\Psi) - C_q(\infty)}{C_q(L) - C_q(\infty)} \approx r_1^{\Psi}.$$
 (23)

That is, asymptotically a pattern, of changes in $C_q(L) - C_q(\infty)$ (with period Ψ), decays exponentially with decay rate of r_1^{Ψ} per period [21].

While the first-order perturbation allowed us to identify both the roles and values of r_1 and Ψ for any process and Corollary 4 would imply Theorem 3, Theorem 3 actually transcends the limitations of the first-order approximation.

Proof 2. Expanding $\log_2 G^{(L)}$ in powers of $(G^{(L)} - I)$, then multiplying by $-G^{(L)}$, shows that $C_q(L) = -\text{tr}[G^{(L)}\log_2 G^{(L)}]$ can be written as

$$C_q(L) = -\sum_{n=0}^{\infty} a_n \operatorname{tr}[(G^{(L)})^n],$$
 (24)

for proper $a_n \in \mathbb{R}$. Using

$$G^{(L)} = \sum_{\xi \in \Lambda_{\xi} \setminus 0} \frac{1 - \xi^{L+1}}{1 - \xi} C_{\xi} + \sum_{m=0}^{\min\{L, \nu_{0} - 1\}} C_{0,m}, \qquad (25)$$

with appropriate constant matrices $C_{0,m}$, together with Eqs. (21) and (24), yields Theorem 3 with general validity.

In the simplest case, when ζ has only one largest eigenvalue, then $\Psi = |\Lambda(r_1)| = 1$ and so $C_q(L) - C_q(\infty)$ is dominated by a simple exponential decay at large L.

For the case of multiple largest eigenvalues originating from the same cycle in the graph of ζ , then $\Psi = |\Lambda(r_1)| > 1$. And so, the asymptotic behavior of $C_q(L) - C_q(\infty)$ is dominated by a decaying pattern of length $|\Lambda(r_1)|$.

For example, the Lollipop processes have an exponentially decaying pattern of length-N that dominates $C_q(L) - C_q(\infty)$ for $L > \nu_0 = M$:

$$\Psi = |\Lambda(r_1)| = N. \tag{26}$$

This periodic behavior is apparent in the semilog plots of Figs. 8 and 10 and is especially emphasized in Fig. 9, which shows that $\Psi = N$ for various N. The figures demonstrate excellent agreement with our qualitative expectations from the above approximations.

Showing the effect of different v_0 , Fig. 10 emphasizes that the initial rolloff of $C_q(L)-C_q(\infty)$ is due to $L\leqslant v_0=M$. The dominant asymptotic behavior is reached soon after $L=v_0$ in this case since the remaining (i.e., nonzero) eigenvalues of the QPMM transition dynamic ζ are all in the largest-magnitude set $\Lambda(r_1)$. In other words, Theorem 2's Eq. (20) is not only approximated by but, in this case, also equal to the simpler expression in Corollary 4, since $r_2=0$.

The slope r_1 indicated in Figs. 8 and 10 corresponds to the asymptotic decay rate of the envelope for $C_q(L) - C_q(\infty)$. This asymptotic decay rate is a function of both N and p, since for Lollipop:

$$r_1 = [(1-p)(1-q)]^{1/N}.$$
 (27)

Figure 8 shows that we have indeed identified the correct slope for different *p*.

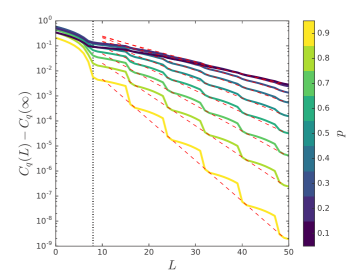


FIG. 8. (8,8)-Lollipop with transition parameters $p \in [0.1,0.9]$, q = 0.5, and r = 0.1. $C_q(L) - C_q(\infty)$ on semilog plot illustrates asymptotically exponential behavior. Red dashed lines, r_1^L where r_1 (no relation to r) is the spectral radius of ζ , quantify the exponential rate of decay. The height of each red dashed line is set equal to $C_q(49)$; we can see that the decay is very close to exponential even as early as $L \simeq 15$. Vertical dashed line at L = M = 8 shows change in behavior after the length of the "stick."

The central asymptotic features of the quantum advantage $C_q(L) - C_q(\infty)$ of reduced state complexity are all captured succinctly by Theorem 3: First, the asymptotic behavior of $C_q(L) - C_q(\infty)$ is exponentially decreasing at rate r_1 , which is the spectral radius of ζ . Second, this exponential envelope is modulated by an asymptotic Ψ -periodic structure, where Ψ is the least common multiple of slowest-decaying QPMM cycle lengths.

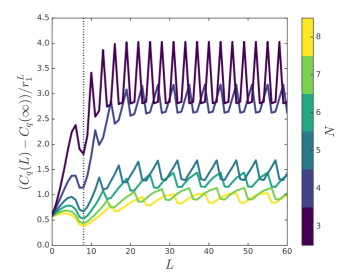


FIG. 9. Lollipop with $N \in \{3,4,5,6,7,8\}$ and M=8, and transition parameters p=q=0.5 and r=0.1. $(C_q(L)-C_q(\infty))/r_1^L$ demonstrates the periodicity of asymptotic behavior. Removing the exponential envelope makes periodicity of the remaining deviation more apparent. For Lollipop, the periodicity $\Psi = |\Lambda(r_1)| = N$.

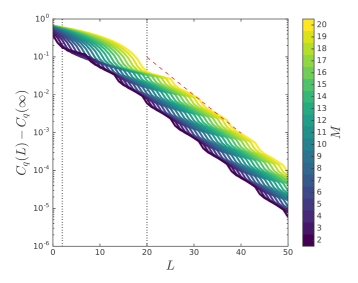


FIG. 10. $C_q(L)-C_q(\infty)$ on a semilog plot for Lollipop with N=6 and $M\in\{2,\ldots,20\}$ and transition parameters p=q=0.5 and r=0.1. M determines the finite-horizon length, where the nilpotent part of ζ vanishes. Vertical (dashed) lines indicate L=2 and 20, the shortest and longest such length in this group.

These results summarize the expected behavior of the L-dependent quantum reduction of state complexity for all classical processes that can be described by a finite-state ϵ -machine. Using codeword length of at least the finite-horizon length ν_0 of the process's QPMM seems advisable for significant reduction of memory costs in simulations that utilize the advantage of quantum signal states discussed here. The cost-benefit analysis of further increasing encoding length for infinite-cryptic processes will be application-specific, but now has theoretical grounding in the above results.

VIII. CONCLUSION

We developed a detailed analytical theory of how to maximally reduce the state complexity of a classical, stationary finite-memory stochastic process using a quantum channel. This required using the new quantum state machine representation (q-machines) [4], carefully constructing its codewords and quantitatively monitoring their overlaps (via the quantum pairwise-merger machine), and utilizing a new matrix formulation of the overlap density matrix (consolidated Gram matrix). Applying spectral decomposition then lead directly to closed-form expressions for the quantum coding costs at any codeword length, including infinite length.

The theoretical advances give an efficient way to probe the behavior of quantum-reduced state complexity with increasing codeword length, both analytically and, when symbolic calculation become arduous, numerically. The efficient numerical algorithm (linear in L) improves on previous exponential algorithms; moreover, the infinite-L limit can now be obtained directly in finite time. Analyzing selected example processes illustrated the required calculations and also the range of phenomena that occur when compressing memoryful processes. We expect the results to aid understanding complex classical stochastic systems of biological and technological importance

via efficient simulations now possible due to the quantum reduction in memory requirements.

Particular phenomena we reported here included (i) details of how a process's cryptic order determines its quantum reduction in state complexity, (ii) transient and persistent contributions to reduced state complexity, (iii) exponential convergence to optimum compression, and (iv) oscillations in the convergence that reveal how a process gives up its crypticity with increasing codeword length. Our results apply to both finite and infinite Markov- and cryptic-order processes.

The overall result appears as a rather complete quantitative toolkit for analyzing quantum state compressibility of classical processes, including finite and infinite codeword closed-form expressions. That said, many issues remain, both technical and philosophical. We believe, however, that the approach's mathematical grounding and analytical and numerical efficiency will go some distance to solving them in the near future.

For example, one of the abiding questions is the meaning of process crypticity $\chi = C_{\mu} - \mathbf{E}$: the difference between a process's predictable information or excess entropy E and its stored state information or statistical complexity C_{μ} [22,23]. Most directly, χ measures how much state information (C_{μ}) is hidden from observation (E). Cryptic processes and even those with infinite cryptic order dominate the space of classical processes [11]. This means that generically we can compress C_{μ} down to $C_{q}(L)$. However, this begs the question of what crypticity is in the quantum domain. Now that we can work analytically in the infinite-length limit, we can explore the quantum crypticity $\chi_q = C_q(\infty) - \mathbf{E}$. From our studies, some not reported here, it appears that one cannot compress the state information all the way down to the excess entropy. Why? Why do not quantum models exist of "size" E bits? Does this point to a future, even more parsimonious physical theory? Or, to a fundamental limitation of communication that even nature must endure, as it channels the past through the present to the future?

For another, are we really justified in comparing Shannon bits (C_{μ}) to qubits (C_{q}) ? This is certainly not a new or recent puzzle. However, the results on compression bring it to the fore anew. And, whatever the outcome, the answer will change our view of what physical pattern and structure are. Likely, the answer will have a profound effect. Assuming the comparison is valid, why is there a perceived level of classical reality that is more structurally complex when, as we demonstrated and now can calculate, processes might be more compactly represented quantum mechanically?

ACKNOWLEDGMENTS

We thank Ryan James for helpful conversations and two anonymous reviewers for helpful suggestions. This material is based upon work supported by, or in part by, the John Templeton Foundation and U.S. Army Research Laboratory and the U.S. Army Research Office under Contract W911NF-13-1-0390.

APPENDIX A: MEALY HMMs

Edge-emitting hidden Markov models (HMMs) are called *Mealy HMMs*. This should be contrasted with the state-emitting HMMs, called *Moore HMMs*. Mealy and Moore HMMs are different representations, but all processes that

can be generated by finite-state Moore HMMs can also be generated by Mealy HMMs, and vice versa; i.e., Mealy and Moore HMMs are class equivalent. The causal equivalence relation, however, implies that the minimal classical model of a process, the ϵ -machine, is a Mealy HMM.

Definition 5. A Mealy HMM \mathcal{M} is the 4-tuple $\{\mathcal{R}, \mathcal{A}, \{T^{(x)}\}_{x \in \mathcal{A}}, \boldsymbol{\mu}_0\}$, where \mathcal{R} is the set $\{\rho_0, \rho_1, \ldots\}$ of latent states, \mathcal{A} is the set of output symbols x, $\{T^{(x)}: T_{i,j}^{(x)} = \Pr(\rho_j, x | \rho_i)\}_{x \in \mathcal{A}}$ consists of the symbol-labeled state transition matrices, and $\boldsymbol{\mu}_0$ is the initial distribution over latent states.

Certain nonstationary processes can be generated by Mealy HMMs when the initial distribution over latent states is not the stationary distribution. In this work, we consider only stationary processes, so $\mu_0 = \pi$. In such cases, time-independent word probabilities can be calculated as

$$\Pr(w) = \pi \prod_{i=0}^{L-1} T^{(x_i)} \mathbf{1},$$

where $w = x_0 \dots x_{L-1}$ and $\mathbf{1} = [1, \dots, 1]^{\top}$. When these probabilities are constructed to agree with those of the words in a given process language, the HMM is said to be a *presentation* of the process.

The ϵ -machine is the Mealy HMM presentation of a process, whose latent states are the process' causal states: $\mathcal{R} = \mathcal{S}$. The ϵ -machine is provably the minimal classical unifilar generator of a process, minimal both in the number of states and the entropy over states [1].

APPENDIX B: QUANTUM OVERLAPS AND CRYPTIC ORDER

Lemma 1. Given an ϵ -machine with cryptic order k: for $L \leq k$, there exists an L-merge; for L > k, there exists no L-merge.

Proof 3. By definition of cryptic order *k*:

$$H[S_k|X_{0:\infty}]=0.$$

This means that for any given x_0 : there exists a unique σ_k . Since k is the minimum such length, for L=k-1 there exists some word $x_{0:\infty}$ that leaves uncertainty in causal state \mathcal{S}_{k-1} . Call two of these uncertain \mathcal{S}_{k-1} states A and B ($A \neq B$). Tracing $x_{0:\infty}$ backwards from A and B, we produce two state paths. These state paths must be distinct at each step due to ϵ -machine unifilarity. If they were not distinct at some step, they would remain so for all states going forward, particularly at \mathcal{S}_{k-1} . The next symbol x_k must take A and B to the same next state F or violate the assumption of cryptic order k. These two state paths and the word $x_{0:k}$ and the final state F make up a k-merger, meaning that cryptic order k implies the existence of a k-merger.

By removing states from the left side of this k-merger, it is easy to see that a k-merger implies the existence of all shorter L-mergers.

By unifilarity again, $H[S_k|X_{0:\infty}] = 0 \Rightarrow H[S_L|X_{0:\infty}] = 0$, for all $L \geqslant k$. Assume there exists an L-merger for L > k with word w. By definition of L-merger, there is then uncertainty in the state S_{L-1} . This uncertainty exists for any word with w as the prefix—a set with nonzero probability. This contradicts the definition of cryptic order.

Theorem 1. Given a process with cryptic order k, for each $L \in \{0, \ldots, k\}$, each quantum overlap $\langle \eta_i(L) | \eta_j(L) \rangle$ is a nondecreasing function of L. Furthermore, for each $L \in \{1, \ldots, k\}$, there exists at least one overlap that is increased (as a result of a corresponding L-merge). For all remaining $L \geqslant k$, each overlap takes a constant value $\langle \eta_i(k) | \eta_j(k) \rangle$.

Proof 4. We directly calculate

$$\begin{split} \langle \eta_a(L) | \eta_b(L) \rangle &= \sum_{\substack{w,w' \in \mathcal{A}^L \\ j_L, l_L \in \{i\}_{i=1}^M}} \sqrt{T_{al_L}^{(w)}} \sqrt{T_{bj_L}^{(w')}} \langle w | w' \rangle \langle \sigma_{l_L} | \sigma_{j_L} \rangle \\ &= \sum_{\substack{w,j_I}} \sqrt{T_{aj_L}^{(w)}} \sqrt{T_{bj_L}^{(w)}}. \end{split}$$

So we have

$$\begin{split} &\langle \eta_{a}(L+1) | \eta_{b}(L+1) \rangle \\ &= \sum_{w' \in \mathcal{A}^{L+1}} \sqrt{T_{aj_{L+1}}^{(w')}} \sqrt{T_{bj_{L+1}}^{(w')}} \\ &= \sum_{w \in \mathcal{A}^{L}, s \in \mathcal{A}} \sqrt{T_{aj_{n}}^{(w)}} \sqrt{T_{j_{n}j_{L+1}}^{(s)}} \sqrt{T_{bl_{L}}^{(w)}} \sqrt{T_{l_{L}j_{L+1}}^{(s)}} \\ &= \sum_{w \in \mathcal{A}^{L}, s \in \mathcal{A}} \sqrt{T_{aj_{L}}^{(w)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \sqrt{T_{bj_{L}}^{(w)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \\ &+ \sum_{w \in \mathcal{A}^{L}, s \in \mathcal{A}} \sqrt{T_{aj_{L}}^{(w)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \sqrt{T_{bl_{L}}^{(w)}} \sqrt{T_{l_{L}j_{L+1}}^{(s)}}. \end{split}$$

The first sum represents the overlaps obtained already at length L. To see this, we split the sum to two parts, where the first contains

$$\begin{split} \sum_{w \in \mathcal{A}^{L}, s \in \mathcal{A}} \sqrt{T_{aj_{L}}^{(w)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \sqrt{T_{bj_{L}}^{(w)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \\ &= \sum_{w \in \mathcal{A}^{L}} \sqrt{T_{aj_{L}}^{(w)}} \sqrt{T_{bj_{L}}^{(w)}} \left(\sum_{\substack{s \in \mathcal{A} \\ j_{L+1}}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \right) \\ &= \sum_{w \in \mathcal{A}^{L}} \sqrt{T_{aj_{L}}^{(w)}} \sqrt{T_{bj_{L}}^{(w)}} = \langle \eta_{a}(L) | \eta_{b}(L) \rangle. \end{split}$$

We use Lemma 1 to analyze the second sum, which represents the change in the overlaps, finding that

$$\sum_{\substack{w \in \mathcal{A}^{L}, s \in \mathcal{A} \\ l_{1}, \dots, l_{L}, l_{L}, \dots}} \sqrt{T_{aj_{L}}^{(w)}} \sqrt{T_{j_{L}j_{L+1}}^{(s)}} \sqrt{T_{bl_{L}}^{(w)}} \sqrt{T_{l_{L}j_{L+1}}^{(s)}} \geqslant 0,$$

with equality when $L \geqslant k$. Summarizing,

$$\langle \eta_a(L+1)|\eta_b(L+1)\rangle \geqslant \langle \eta_a(L)|\eta_b(L)\rangle,$$

with equality for $L \ge k$.

Note that while the set of overlaps continues to be augmented at each length up until the cryptic order, we do not currently have a corresponding statement about the nontrivial change in $C_q(L)$ or its monotonicity. Although a proof has been elusive, it would be an important extension of our work. Nevertheless, the asymptotic analysis of Sec. VII shows an overall decay of $C_q(L)$ for infinite cryptic processes. Moreover, extensive numerical exploration suggests that $C_q(L)$ is indeed monotonic at all scales for all orders of crypticity.

APPENDIX C: MATRICES AND THEIR ENTROPY

1. Density matrix

The density matrix can now be expressed using a fixed $|\mathcal{S}|$ -by- $|\mathcal{S}|$ matrix, valid for all L. Using the Gram-Schmidt procedure one can choose a new orthonormal basis. Let

$$\begin{aligned} |\eta_{1}(L)\rangle &= \left| e_{1}^{(L)} \right\rangle, \\ |\eta_{2}(L)\rangle &= a_{21}^{(L)} \left| e_{1}^{(L)} \right\rangle + a_{22}^{(L)} \left| e_{2}^{(L)} \right\rangle, \\ |\eta_{3}(L)\rangle &= a_{31}^{(L)} \left| e_{1}^{(L)} \right\rangle + a_{32}^{(L)} \left| e_{2}^{(L)} \right\rangle + a_{33}^{(L)} \left| e_{3}^{(L)} \right\rangle, \\ &\vdots \end{aligned}$$

and so on. Then

$$\begin{aligned} a_{21}^{(L)} &= \langle \eta_1(L) | \eta_2(L) \rangle \\ &= \langle (\sigma_1, \sigma_2) | \left(\sum_{n=0}^L \zeta^n \right) | \text{SINK} \rangle, \\ a_{22}^{(L)} &= (1 - |\langle \eta_1(L) | \eta_2(L) \rangle|^2)^{1/2}, \\ a_{31}^{(L)} &= \langle \eta_1(L) | \eta_3(L) \rangle \\ &= \langle (\sigma_1, \sigma_3) | \left(\sum_{n=0}^L \zeta^n \right) | \text{SINK} \rangle, \end{aligned}$$

and so on. Now, it is useful to rewrite what we can in matrix form:

$$\begin{bmatrix} \langle \eta_{1}(L) | \\ \langle \eta_{2}(L) | \\ \langle \eta_{3}(L) | \\ \vdots \\ \langle \eta_{|\mathcal{S}|}(L) | \end{bmatrix}$$

$$= \underbrace{ \begin{bmatrix} 1 & & & & 0 \\ a_{21}^{(L)} & a_{22}^{(L)} & & & \\ a_{31}^{(L)} & a_{32}^{(L)} & a_{33}^{(L)} & & \\ \vdots & & \ddots & & \\ a_{|\mathcal{S}|1}^{(L)} & \cdots & & a_{|\mathcal{S}||\mathcal{S}|}^{(L)} \end{bmatrix} }_{\mathbf{A}} \begin{bmatrix} \langle e_{1}^{(L)} | \\ \langle e_{2}^{(L)} | \\ \langle e_{3}^{(L)} | \\ \vdots \\ \langle e_{|\mathcal{S}|}^{(L)} | \end{bmatrix},$$

which defines the lower-triangular matrix A_L . Note that the rightmost matrix of orthonormal basis vectors is simply the identity matrix, since we are working in that basis.

In this new basis, we construct the |S|-by-|S| density matrix as

$$\rho(L) = \sum_{i=1}^{|\mathcal{S}|} \pi_i |\eta_i(L)\rangle \langle \eta_i(L)|$$

$$= [|\eta_1(L)\rangle \quad \cdots \quad |\eta_{|\mathcal{S}|}(L)\rangle]$$

$$\times \begin{bmatrix} \pi_1 & 0 \\ & \ddots \\ 0 & \pi_{|\mathcal{S}|} \end{bmatrix} \begin{bmatrix} \langle \eta_1(L)| \\ \langle \eta_2(L)| \\ \langle \eta_3(L)| \\ \vdots \\ \langle \eta_{|\mathcal{S}|}(L)| \end{bmatrix}$$

$$= A^{\dagger} D A$$

Since all entries are real, the conjugate transpose is the transpose. This more general framework may be useful, however, if we want to consider the effect of adding phase to the quantum states.

2. Von Neumann entropy

The quantum coding cost is

$$\begin{split} C_q(L) &= -\mathrm{tr}[\rho(L)\log_2\rho(L)] \\ &= -\mathrm{tr}[A_L^\dagger D_\pi A_L \log_2(A_L^\dagger D_\pi A_L)] \\ &= -\sum_{\lambda \in \Lambda_{A_L^\dagger D_\pi A_L}} \lambda \log_2 \lambda. \end{split}$$

This is relatively easy to evaluate since the density matrix $\rho(L)$ is only a $|\mathcal{S}|$ -by- $|\mathcal{S}|$ function of L. Thus, we calculate $C_q(L)$ analytically from the spectrum of ρ . This, in a curious way, was already folded into the spectrum of ζ .

3. Gram matrix

The A_L matrix is burdensome due to nonlinear dependence on the overlap of the quantum states. We show how to avoid this nonlinearity and instead obtain the von Neumann entropy from a transformation that yields a linear relationship with overlaps.

The *Gram matrix*, with elements $G_{mn}^{(L)} = \sqrt{\pi_m \pi_n} \langle \eta_m(L) | \eta_n(L) \rangle$, can be used instead of $\rho(L)$ to evaluate the von Neumann entropy [19]. In particular, $G^{(L)}$ has the same spectrum as $\rho(L)$, even with the same multiplicities: $\Lambda_{G^{(L)}} = \Lambda_{\rho(L)}$, while a_λ , g_λ , and ν_λ remain unchanged for all λ in the spectrum. (This is a slightly stronger statement than in Ref. [19], but is justified since $\rho(L)$ and $G^{(L)}$ are both $|\mathcal{S}|$ -by- $|\mathcal{S}|$ dimensional.)

Here we briefly explore the relationship between $\rho(L)$ and $G^{(L)}$ and, then, focus on the closed-form expression for $G^{(L)}$. The result is more elegant than $\rho(L)$, allowing us to calculate and understand $C_q(L)$ more directly.

Earlier we found that the density matrix can be written as

$$\rho(L) = A_L^{\dagger} D_{\pi} A_L,$$

which can be rewritten as

$$\rho(L) = A_L^{\dagger} D_{\pi}^{1/2} D_{\pi}^{1/2} A_L$$
$$= \left(D_{\pi}^{1/2} A_L \right)^{\dagger} D_{\pi}^{1/2} A_L.$$

It is easy to show that

$$\begin{aligned} \text{tr} \big[\big(D_{\pi}^{1/2} A_L \big)^{\dagger} D_{\pi}^{1/2} A_L \big] &= \text{tr} \big[D_{\pi}^{1/2} A_L \big(D_{\pi}^{1/2} A_L \big)^{\dagger} \big] \\ &= \text{tr} \big[D_{\pi}^{1/2} A_L A_L^{\dagger} D_{\pi}^{1/2} \big]. \end{aligned}$$

This means that the sum of the eigenvalues is conserved in transforming from $A_L^{\dagger}D_{\pi}A_L$ to $D_{\pi}^{1/2}A_LA_L^{\dagger}D_{\pi}^{1/2}$. It is less obvious that the spectrum is also conserved, but this is also true and even easy to prove. [Observe that $AB\vec{v} = \lambda\vec{v} \Longrightarrow BAB\vec{v} = \lambda B\vec{v} \Longrightarrow BA(B\vec{v}) = \lambda(B\vec{v})$.] Interestingly, the new object turns out to be exactly the Gram matrix, which was previously introduced, although without this explicit relationship to the density matrix. We now see that

$$\begin{split} D_{\pi}^{1/2}A_{L}A_{L}^{\dagger}D_{\pi}^{1/2} &= D_{\pi}^{1/2}\begin{bmatrix} \langle \eta_{1}(L)| \\ \vdots \\ \langle \eta_{|\mathcal{S}|}(L)| \end{bmatrix} [|\eta_{1}(L)\rangle \quad \cdots \quad |\eta_{|\mathcal{S}|}(L)\rangle] D_{\pi}^{1/2} \\ &= \begin{bmatrix} \sqrt{\pi_{1}}\langle \eta_{1}(L)| \\ \vdots \\ \sqrt{\pi_{|\mathcal{S}|}}\langle \eta_{|\mathcal{S}|}(L)| \end{bmatrix} [\sqrt{\pi_{1}}|\eta_{1}(L)\rangle \quad \cdots \quad \sqrt{\pi_{|\mathcal{S}|}}|\eta_{|\mathcal{S}|}(L)\rangle] \\ &= \begin{bmatrix} \sqrt{\pi_{1}\pi_{1}}\langle \eta_{1}(L)|\eta_{1}(L)\rangle & \cdots & \sqrt{\pi_{1}\pi_{|\mathcal{S}|}}\langle \eta_{1}(L)|\eta_{|\mathcal{S}|}(L)\rangle \\ \vdots & \ddots & \vdots \\ \sqrt{\pi_{|\mathcal{S}|}\pi_{1}}\langle \eta_{|\mathcal{S}|}(L)|\eta_{1}(L)\rangle & \cdots & \sqrt{\pi_{1}\pi_{|\mathcal{S}|}}\langle \eta_{|\mathcal{S}|}(L)|\eta_{|\mathcal{S}|}(L)\rangle \end{bmatrix} \\ &= G^{(L)} \end{split}$$

Since the spectrum is preserved, we can use the Gram matrix directly to compute the von Neumann entropy:

$$C_q(L) = -\sum_{\boldsymbol{\lambda} \in \Lambda_{G^{(L)}}} \boldsymbol{\lambda} \log_2 \boldsymbol{\lambda} = -\mathrm{tr}[G^{(L)} \log_2 G^{(L)}].$$

4. Consolidated Gram matrix

Transforming to the Gram matrix suggests a similar and even more helpful simplification that can be made while preserving the spectrum. Define the *left-consolidated Gram matrix* to be

$$\begin{split} \widetilde{G}^{(L)} &\equiv D_{\pi} A_{L} A_{L}^{\dagger} \\ &= D_{\pi} \begin{bmatrix} \langle \eta_{1}(L) | \\ \vdots \\ \langle \eta_{|\mathcal{S}|}(L) | \end{bmatrix} [|\eta_{1}(L)\rangle & \cdots & |\eta_{|\mathcal{S}|}(L)\rangle] \\ &= \begin{bmatrix} \pi_{1} \langle \eta_{1}(L) | \eta_{1}(L)\rangle & \cdots & \pi_{1} \langle \eta_{1}(L) | \eta_{|\mathcal{S}|}(L)\rangle \\ \vdots & \ddots & \vdots \\ \pi_{|\mathcal{S}|} \langle \eta_{|\mathcal{S}|}(L) | \eta_{1}(L)\rangle & \cdots & \pi_{|\mathcal{S}|} \langle \eta_{|\mathcal{S}|}(L) | \eta_{|\mathcal{S}|}(L)\rangle \end{bmatrix}. \end{split}$$

Clearly, this preserves the same trace as the density matrix and previous Gram matrix. It also preserves the spectrum, and it has the advantage of not using square-roots of two different state probabilities in each element. Rather it has a single probability attached to each element. The same is true for the right-consolidated Gram matrix $A_L A_L^{\dagger} D_{\pi}$.

Since the spectrum is preserved, we can use the consolidated Gram matrix to compute the von Neumann entropy:

$$C_q(L) = -\sum_{\lambda \in \Lambda_{\widetilde{O}(L)}} \lambda \log_2 \lambda \tag{C1}$$

$$= -\text{tr}[\widetilde{G}^{(L)}\log_2\widetilde{G}^{(L)}]. \tag{C2}$$

APPENDIX D: EXAMPLES

Exploring several more examples will help to illustrate the methods and lead to additional observations.

1. Biased Coins Process

The Biased Coins Process provides a first, simple case that realizes a nontrivial quantum state entropy [2]. There are two biased coins, named A and B. The first generates 1 with probability q; the second, 0 with probability p. A coin is picked and flipped, generating outputs 0 or 1. With probability q the other coin is used next similarly with different probability. Its two causal-state ϵ -machine is shown in Fig. 11.

After constructing the QPMM for the Biased Coins Process, as outlined in Figs. 11 and 12, we observe

$$\boldsymbol{\zeta}^{(0)} = \begin{bmatrix} 0 & \sqrt{p(1-q)} \\ 0 & 0 \end{bmatrix},$$

$$\boldsymbol{\zeta}^{(1)} = \begin{bmatrix} 0 & \sqrt{q(1-p)} \\ 0 & 0 \end{bmatrix},$$

and so:

$$\zeta = \begin{bmatrix} 0 & \beta \\ 0 & 0 \end{bmatrix},$$

where we defined $\beta \equiv \sqrt{p(1-q)} + \sqrt{q(1-p)}$. Let us also define the suggestive quantity $\gamma \equiv (1-\beta^2)^{-1/2}$.

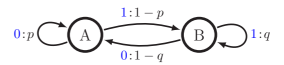


FIG. 11. ϵ -Machine for the Biased Coins Process.

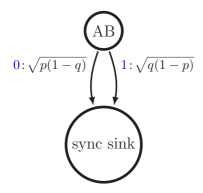


FIG. 12. QPMM for the Biased Coins Process.

The only overlap to consider is $\langle \eta_A(L) | \eta_B(L) \rangle$. For this, we note that $\langle (A,B) | = \begin{bmatrix} 1 & 0 \end{bmatrix}$. Also, $|\text{SINK}\rangle = \begin{bmatrix} 0 & 1 \end{bmatrix}^{\top}$.

Spectrally, ζ here is a nilpotent matrix with only a zero eigenvalue with index two: $\Lambda_{\zeta} = \{0\}$ and $\nu_0 = 2$. Since the projection operators must sum to the identity, we have $\zeta_0 = I$.

 ζ^L is the null matrix for L > 1, so either by Eq. (6) or by Eq. (8), we have

$$\langle \eta_A(L)|\eta_B(L)\rangle = \sum_{m=1}^{\min\{L, 1\}} \langle (A, B)|\zeta^m|\text{SINK}\rangle;$$

that is,

$$\langle \eta_A(L) | \eta_B(L) \rangle = \begin{cases} 0 & \text{if } L = 0, \\ \beta & \text{if } L \geqslant 1. \end{cases}$$

(a) Entropy from the density matrix

For the density matrix, we turn to the L-dependent orthonormal basis $\{|e_1^{(L)}\rangle,|e_2^{(L)}\rangle\}$ and use the stationary distribution over \mathcal{S} : $\pi=[p/(p+q)-q/(p+q)]$.

Apparently for L=0 we have $|\eta_A(0)\rangle = |e_1^{(0)}\rangle$ and $|\eta_B(0)\rangle = |e_2^{(0)}\rangle$. Hence, $\rho(0) = D_\pi$ and $C_q(0) = H_2(p/(p+q)) = C_\mu$ qubits.

For $L \ge 1$ we have: $|\eta_A(L)\rangle = |e_1^{(L)}\rangle$ and $|\eta_A(L)\rangle = a_{21}^{(L)}|e_1^{(L)}\rangle + a_{22}^{(L)}|e_2^{(L)}\rangle$, where $a_{21}^{(L)} = \langle \eta_A(L)|\eta_B(L)\rangle = \beta$ and $a_{22}^{(L)} = (1 - \beta^2)^{1/2} = \gamma^{-1}$ for $L \ge 1$. We find that

$$A_L = \begin{bmatrix} 1 & 0 \\ \beta & \gamma^{-1} \end{bmatrix}$$
, for $L \geqslant 1$.

Hence, for $L \ge 1$ the density matrix is

$$\rho(L) = A_L^{\dagger} D_{\pi} A_L$$

$$= \begin{bmatrix} 1 & \beta \\ 0 & \gamma^{-1} \end{bmatrix} \begin{bmatrix} \frac{p}{p+q} & 0 \\ 0 & \frac{q}{p+q} \end{bmatrix} \begin{bmatrix} 1 & 0 \\ \beta & \gamma^{-1} \end{bmatrix}$$

$$= \frac{1}{p+q} \begin{bmatrix} p & q\beta \\ 0 & q\gamma^{-1} \end{bmatrix} \begin{bmatrix} 1 & 0 \\ \beta & \gamma^{-1} \end{bmatrix}$$

$$= \frac{q}{p+q} \begin{bmatrix} \frac{p}{q} + \beta^2 & \beta/\gamma \\ \beta/\gamma & 1 - \beta^2 \end{bmatrix}.$$

Since

$$\det(\rho(L) - \lambda I) = \lambda^2 - \lambda + \frac{pq}{(p+q)^2} (1 - \beta^2),$$

we find the eigenvalues of $\rho(L)$ eigenvalues to be

$$\Lambda_{\rho(L)} = \left\{ \frac{1}{2} \pm \frac{1}{2(p+q)} \sqrt{4pq\beta^2 + (p-q)^2} \right\},$$

which yields the von Neumann entropy for $L \geqslant 1$:

$$C_q(L) = -\sum_{\lambda \in \Lambda_{q(I)}} \lambda \log_2 \lambda.$$

(b) Entropy from the consolidated Gram matrix

The left-consolidated Gram matrix for the Biased Coins Process is

$$\widetilde{G}^{(L)} = D_{\pi} \begin{bmatrix} \langle \eta_A(L) | \eta_A(L) \rangle & \langle \eta_A(L) | \eta_B(L) \rangle \\ \langle \eta_B(L) | \eta_A(L) \rangle & \langle \eta_B(L) | \eta_B(L) \rangle \end{bmatrix}.$$

Specifically, we have for L=0

$$\widetilde{G}^{(0)} = \frac{1}{p+q} \begin{bmatrix} p & 0 \\ 0 & q \end{bmatrix} \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$$
$$= \frac{1}{p+q} \begin{bmatrix} p & 0 \\ 0 & q \end{bmatrix},$$

and $L \geqslant 1$:

$$\widetilde{G}^{(L)} = \frac{1}{p+q} \begin{bmatrix} p & 0 \\ 0 & q \end{bmatrix} \begin{bmatrix} 1 & \beta \\ \beta & 1 \end{bmatrix}$$
$$= \frac{1}{p+q} \begin{bmatrix} p & p\beta \\ q\beta & q \end{bmatrix}.$$

 $\widetilde{G}^{(0)}$'s eigenvalues are simply its diagonal entries. So, $C_q(0)=H_2(p/(p+q))$ qubits. For $L\geqslant 1$,

$$\det(\widetilde{G}^{(L)} - \lambda I) = \lambda^2 - \lambda + \frac{pq}{(p+q)^2} (1 - \beta^2),$$

which gives the same values for eigenvalues and entropy as we found earlier using the density matrix approach.

As the new method illustrates, there is no need to construct the density matrix. Instead, one uses the consolidated Gram matrix, which can be easily calculated from quantum overlaps. Clearly, the consolidated Gram matrix method is more elegant for our purposes. This is evident even at $|\mathcal{S}| = 2$. This is even more critical for more complex processes since A_L grows as $|\mathcal{S}|$ grows.

2. (R-k)-Golden Mean Process

The (R-k)-Golden Mean Process is constructed to have Markov-order R and cryptic-order k. Its ϵ -machine is shown in Fig. 13. The 0th state σ_0 has probability $\pi_0 = 1/[R+k-p(R+k-1)]$ while all other states σ_i have probability $\pi_i = (1-p)\pi_0$.

Its QPMM is strictly tree-like with depth d = k + 1 and maximal width k. All edges have a unit weight except for those edges leaving A-paired states. The latter edges, numbering k in total, have an associated weight of \sqrt{p} .

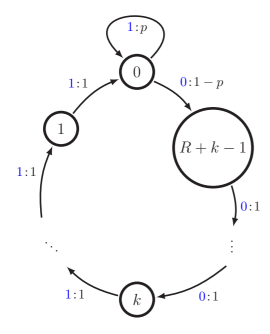


FIG. 13. ϵ -Machine for the (R-k)-Golden Mean Process.

The eigenvalues of the consolidated Gram matrix can be obtained from $\det(\widetilde{G}^{(L)} - \lambda I) = 0$, where

$$\det(\widetilde{G}^{(L)} - \lambda I)$$

$$= (\pi_1 - \lambda)^{R+k-\min(L,k)-1}$$

$$\times \begin{vmatrix} \pi_0 - \lambda & \pi_0 \sqrt{p} & \cdots & \pi_0 \sqrt{p}^{\min(L,k)} \\ \pi_1 \sqrt{p} & \pi_1 - \lambda & \pi_1 \sqrt{p}^{\min(L,k)-1} \\ \vdots & \ddots & \vdots \\ \pi_1 \sqrt{p}^{\min(L,k)} & \pi_1 - \lambda \end{vmatrix}$$

This directly yields the von Neumann entropy. Note that although the $C_q(L)$ is *not* actually linear in L, it appears approximately linear.

We observe that π is invariant under the simultaneous change of

$$R' = R + m, \text{ while } k' = k - m, \tag{D1}$$

for any $m \in \mathbb{Z}$. Although we insist on maintaining $R' \geqslant k' \geqslant 0$ for preservation of their functional roles. Furthermore, this

transformation preserves $\det(\widetilde{G}^{(L)} - \lambda I)$ for $L \leqslant k$ and $L \leqslant k'$. Hence, $C_q(L)$ is invariant to the simultaneous transformation of Eq. (D1) for $L \leqslant k$ and $L \leqslant k'$. This explains the agreement noted in the caption of Fig. 6—that $C_q(L)$ for (R-k)-GM is the same as $C_q(L)$ for ((R+1)-(k-1))-GM for $L \leqslant k$.

To give an explicit example, let us consider the (4–3)-GM Process of Fig. 1. State A has probability $\pi_A = 1/(7 - 6p)$ while all other states have probability $\pi_i = (1 - p)/(7 - 6p)$. Let us calculate the following:

(1) For L = 0:

$$\det(\widetilde{G}^{(0)} - \lambda I) = (\pi_B - \lambda)^6 (\pi_A - \lambda),$$

yielding $\Lambda_{\widetilde{G}^{(0)}} = \{\pi_B, \pi_A\}$ (with $a_{\pi_B} = 6$) and

$$C_a(0) = -6\pi_B \log_2 \pi_B - \pi_A \log_2 \pi_A.$$

(2) For L = 1:

$$\det(\widetilde{G}^{(1)} - \lambda I)$$

$$= (\pi_B - \lambda)^5 [\lambda^2 - (\pi_A + \pi_B)\lambda + \pi_A \pi_B (1 - p)],$$

yielding $\Lambda_{\widetilde{G}^{(1)}} = \{\pi_B, c_+, c_-\}$ with $c_{\pm} = \frac{1}{2}(\pi_A + \pi_B) \pm \frac{1}{2}[(\pi_A + \pi_B)^2 - 4\pi_A\pi_B(1 - p)]^{1/2}$ (and with $a_{\pi_B} = 5$), and

$$C_a(1) = -5\pi_B \log_2 \pi_B - c_+ \log_2 c_+ - c_- \log_2 c_-.$$

(3) For L = 2:

$$\det(\widetilde{G}^{(2)} - \lambda I)$$

$$= (\pi_B - \lambda)^4 \begin{vmatrix} \pi_A - \lambda & \pi_A p^{1/2} & \pi_A p \\ \pi_B p^{1/2} & \pi_B - \lambda & \pi_B p^{1/2} \\ \pi_B p & \pi_B p^{1/2} & \pi_B - \lambda \end{vmatrix}.$$

(4) For $L \geqslant 3$:

$$\det(\widetilde{G}^{(L)} - \lambda I)$$

$$= \det(\widetilde{G}^{(3)} - \lambda I)$$

$$= (\pi_B - \lambda)^3 \begin{vmatrix} \pi_A - \lambda & \pi_A p^{1/2} & \pi_A p & \pi_A p^{3/2} \\ \pi_B p^{1/2} & \pi_B - \lambda & \pi_B p^{1/2} & \pi_B p \\ \pi_B p & \pi_B p^{1/2} & \pi_B - \lambda & \pi_B p^{1/2} \\ \pi_B p^{3/2} & \pi_B p & \pi_B p^{1/2} & \pi_B - \lambda \end{vmatrix}.$$

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