Pattern Discovery in Time Series, Part I: Theory, Algorithm, Analysis, and Convergence

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Abstract

We present a new algorithm for discovering patterns in time series and other sequential data. We exhibit a reliable procedure for building the minimal set of hidden, Markovian states that is statistically capable of producing the behavior exhibited in the data — the underlying process's causal states. Unlike conventional methods for fitting hidden Markov models (HMMs) to data, our algorithm makes no assumptions about the process's causal architecture (the number of hidden states and their transition structure), but rather infers it from the data. It starts with assumptions of minimal structure and introduces complexity only when the data demand it. Moreover, the causal states it infers have important predictive optimality properties that conventional HMM states lack. Here, in Part I, we introduce the algorithm, review the theory behind it, prove its asymptotic reliability, and use large deviation theory to estimate its rate of convergence. In the sequel, Part II, we outline the algorithm’s implementation, illustrate its ability to discover even “difficult” patterns, and compare it to various alternative schemes.


Keywords: Pattern discovery, hidden Markov models, causal inference, statistical complexity, computational mechanics, information theory

1. Introduction

Recent years have seen a rising interest in the problem of pattern discovery (Crutchfield, 1994, Hand et al., 2001, Spirtes et al., 2001): Given data produced by a process, how can one extract meaningful, predictive patterns from it, without fixing in advance the kind of patterns one looks for? The problem represents the convergence of several fields: unsupervised learning (Hinton and Sejnowski, 1999), data mining and knowledge discovery (Hastie et al., 2001), causal inference in statistics (Pearl, 2000), and the statistical physics of complex and highly organized forms of matter (Badii and Politi, 1997, Feldman and Crutchfield, 1998, Shalizi and Crutchfield, 2001, Varn et al., 2002). It should be carefully distinguished both from pattern recognition (essentially a matter of learning to classify instances into known categories) and from merely building predictors which, notoriously, may not lead to any genuine understanding of the process (Crutchfield, 1992). We do not want to merely recognize patterns; we want to find the patterns that are there to be recognized. We do not only want to forecast; we want to know the hidden mechanisms that make the forecasts

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work. The point, in other words, is to find the causal, dynamical structure intrinsic to the process we are investigating, ideally to extract all the patterns in it that have any predictive power.

Naturally enough, an account of what “pattern” means (especially in the sequential setting) is crucial to progress in the above fields. We and co-authors have, over several years, elaborated such an account in a series of publications on the computational mechanics of dynamical systems (Crutchfield and Young, 1989, Crutchfield, 1994, Upper, 1997, Feldman and Crutchfield, 1998, Shalizi and Crutchfield, 2001). Here, we cash in these foundational studies to devise a new and practical algorithm — Causal-State Splitting Reconstruction (CSSR) — for the problem of predicting time series and other sequential data. But, to repeat, we are not giving yet another prediction algorithm; not only is the predictor delivered by our algorithm provably optimal, in a straightforward sense that will be made plain below, but it is also a causal model.

The overall development is broken into two parts. Here, in Part I, we first review the basic ideas and results of computational mechanics, limiting ourselves to those we will use more or less directly. With this background in place, we present the CSSR algorithm in considerable detail. We analyze its run-time complexity, prove its convergence using large-deviation arguments, and discuss how to bound the rate of convergence. By way of elucidating CSSR we compare it to earlier computational mechanics algorithms for causal-state reconstruction. Our conclusion summarizes our results and marks out directions for future theoretical work. Thus, Part I focuses on the theoretical aspects and motivating concepts that lead to CSSR. In the sequel, Part II (Shalizi et al., 2002), we show experimentally how well CSSR does at reconstructing the causal architecture of known processes from data and see how various parameters affect the reconstructions. There, we compare it with other well known procedures for inferring Markovian structure in time series.

2. Computational Mechanics

Consider a discrete-time, discrete-valued stochastic process, \( S_{-2}S_{-1}S_0S_1S_2 \ldots \) with \( S_t \) taking values in \( A \), a finite alphabet of \( k \) symbols. At any time \( t \), we can break the sequence of random variables into a past (or history) \( S_t \) and a future \( S_t \), both of which in general extend infinitely far. Assume the process is conditionally stationary; i.e., for all measurable future events \( A \), \( P(S_t \in A \mid S_1 = \hat{s}) \) does not depend on \( t \). We accordingly drop the subscript and simply speak of \( \hat{S} \) and \( \hat{s} \).

When we want to refer only to the first \( L \) symbols of \( S \) we write \( \hat{S} \) and, similarly, \( \hat{s} \) denotes the last \( L \) symbols of \( S \). (Lower-case versions of these random variables denote particular instances.)

The problem of predicting the future is to go from \( \hat{S} \) to a guess at \( \hat{S} \) or, more generally, a guess at the future distribution \( P(\hat{S}) \). There is thus a function \( \eta \) from particular histories \( \hat{s} \) to predictions \( P(\hat{S} \mid \hat{s}) \). Each prediction method therefore imposes a partition on the set \( S \) of histories. The cells of this partition collect all the histories for which the same prediction is made; i.e., \( \hat{s}_1 \) and \( \hat{s}_2 \) are in the same partition cell if and only if \( \eta(\hat{s}_1) = \eta(\hat{s}_2) \). The cells of the partition are the effective states of the process, under the given predictor. We will, by a slight abuse of notation, also use \( \eta \) as the name of the function from histories to effective states. The random variable for the current effective state will be \( \mathcal{R} \) and a particular value of it \( \rho \). The set of states induced by \( \eta \) will be denoted \( \mathcal{R} \).

Clearly, given a partition \( \mathcal{R} \) of \( \hat{S} \), the best prediction to make is whatever the distribution of futures, conditional on that partition, happens to be: \( P(\hat{S} \mid \mathcal{R} = \eta(\hat{s})) \). Getting this conditional

1. Every stationary process is conditionally stationary.
2. It is important to distinguish, here, between the effective states of a given predictor and alternative states used in particular representations of that predictor. In some of the hidden Markov model examples we will discuss in Part II, what are called the (hidden) Markovian states are not the same as the effective states of the predictor, since the former are not equivalence classes of histories. It is always possible, however, to derive the effective states from the HMM states (Upper, 1997).
distribution right is nontrivial, but nonetheless secondary to getting the partition right. The result is that the problem of finding an optimal predictor reduces (largely) to that of finding an optimal partition of $\vec{S}$.

In what sense is one partition, one predictor, better than another? One obvious sense, inspired by information theory, employs the mutual information $I[\vec{S};\mathcal{R}]$ between the process’s future $\vec{S}$ and the effective states $\mathcal{R}$ to quantify “goodness”. This quantity is not necessarily well defined if we consider the entire semi-infinite future $\vec{S}$, since $I$ could be infinite. But it is always well defined if we look at futures $\vec{S}^L$ of finite length. Since $I[\vec{S}^L;\mathcal{R}] = H[\vec{S}^L] - H[\vec{S}^L|\mathcal{R}]$ and the first term is the same for all sets of states, maximizing the mutual information is the same as minimizing the conditional entropy. There is a lower bound on this conditional entropy, namely $H[\vec{S}^L|\mathcal{R}] \geq H[\vec{S}^L|\vec{s}]$ — one can do no better than to remember the whole past. Call a set of states (a partition) that attains this lower bound for all $L$ prescient. We have shown elsewhere (Shalizi and Crutchfield, 2001) that prescient states are sufficient statistics for the process’s future and so, for any reasonable loss function, the optimal prediction strategy can be implemented using prescient states (Blackwell and Girshick, 1954). We will therefore focus exclusively on conditional entropy.

In general, there are many alternative sets of prescient states. To select among them, one can invoke Occam’s Razor and chose the simplest set of prescient states. (In the computational mechanics framework minimality is not assumed — it is a consequence; see below.) Complexity here is also measured information-theoretically, by the Shannon entropy $H$, which is also written $C_p(\mathcal{R})$ in information-theoretical parlance; then, our goal is to find a set of prescient states of minimal statistical complexity.

It turns out that, for each process, there is a unique set of prescient states that minimizes the statistical complexity. These are the causal states.

**Definition 1 (A Process’s Causal States)** The causal states of a process are the the range of a function $\epsilon$ that maps from histories to sets of histories. If $\mu(\vec{S})$ is the collection of all measurable future events, then

$$\epsilon(s) = \{ s’ \in \vec{S} \mid \forall F \in \mu(\vec{S}), P(\vec{s} \in F| \vec{s} = s) = P(\vec{s}’ \in F| \vec{s} = s’) \} ,$$

(1)

Write the $i$th causal state as $\sigma_i$ and the set of all causal states as $\mathcal{S}$; the corresponding random variable is denoted $\mathcal{S}$ and its realization $\sigma$.

A consequence of this definition is that, for any $L$,

$$\epsilon(\vec{s}) = \{ s’ \mid P(\vec{s}^L = s^L | \vec{s} = \vec{s}) = P(\vec{s}^L = s^L | \vec{s} = \vec{s}’) , \forall s^L \in \vec{S}^L , \vec{s}’ \in \vec{S} \} .$$

(2)

Given any other set of prescient states, say $\bar{\mathcal{R}}$, there is a function $f$ such that $\bar{S} = f(\bar{\mathcal{R}})$ almost always. Hence the entropy of the causal states — their statistical complexity — is less than or equal to that of the prescient states. Moreover, if $C_p(\bar{\mathcal{R}}) = C_p(\mathcal{S})$, then $f$ is invertible and the two sets of states are equivalent, up to relabeling (and a possible set of exceptional histories of measure zero). Statistically speaking, the causal states are the minimal sufficient statistic for predicting the process’s future.

Given an initial causal state and the next symbol from the original process, only certain successor causal states are possible. Thus, we may define allowed transitions between causal states and the

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3. For brief definitions of information-theoretic terms and notation, see the appendix. For more details, see Part II of Cover and Thomas (1991).

4. Notably, the equivalence-class definition of causal states leads directly to their being the minimal set. That is, one does not have to invoke Occam’s Razor, it is derived as a consequence of the states being causal.
probabilities of these transitions. Specifically, the probability of moving from state \( \sigma_i \) to state \( \sigma_j \) on symbol \( s \) is

\[
T_{ij}^{(s)} \equiv P(S^{-1} = s, S' = \sigma_j | S = \sigma_i).
\]  

(3)

Note that

\[
\sum_{s \in \mathcal{A}} \sum_{\sigma_i \in \mathcal{S}} T_{ij}^{(s)} = \sum_{s \in \mathcal{A}} P(S^{-1} = s | S = \sigma_i) = 1.
\]

We denote the set of these labeled transition probabilities by \( T = \{ T_{ij}^{(s)} : s \in \mathcal{A}, \sigma_i, \sigma_j \in \mathcal{S} \} \). The combination of states and transitions is called the process’s \( \epsilon \)-machine; it represents the way in which the process stores and transforms information (Crutchfield and Young, 1989). Examples for a number of different kinds of process — including HMMs, cellular automata, and chaotic dynamical systems — are given in (Crutchfield, 1994). An \( \epsilon \)-machine’s entropy rate \( h_\mu(M) \) is the transition uncertainty averaged over all of the causal states.

### 2.1 Properties of Causal States and \( \epsilon \)-Machines

Here we state the main properties of causal states which we use below. See Shalizi and Crutchfield (2001) for proofs.

**Proposition 2** The causal states are homogeneous for future events: That is, all histories belonging to a single causal state \( \sigma \) have the same conditional distribution \( P(S_j | \sigma) \) of future events. The causal states are also the largest sets (partition elements) that are homogeneous for future events: every history with that future conditional distribution is in that state.

**Proposition 3** The process’s future and past are independent

**Proposition 4** The causal states themselves form a Markov p

**Proposition 5** The \( \epsilon \)-machine is deterministic in the sense of automata theory; i.e., given the current state and the next symbol, there is a unique successor state.

Following practice in symbolic dynamics (see, e.g., Lind and Marcus (1995)), we also say that the causal states are future resolving. That is to say, for each state \( \sigma_i \) and symbol \( s \), there is at most one state \( \sigma_j \) such that \( T_{ij}^{(s)} > 0 \).

**Proposition 6** Any prescient set \( \tilde{\mathcal{R}} \) of states is a refinement of the causal states \( \mathcal{S} \).

For more on these properties, and the reasons for calling \( \epsilon \)-machine states causal, see Shalizi and Crutchfield (2001), especially Sections II–IV.

Finally, there is one additional result which is essential to CSSR that we give it together with a summary of its proof.

**Proposition 7** If a set of states is homogeneous for futures of length 1 and deterministic, then it is prescient.

**Proof**: A rough way to see this is to imagine using homogeneity to obtain a prediction for the next symbol, updating the current state using determinism, then getting a prediction for the second symbol from homogeneity again, and so on, as far forward as needed.

More formally, we proceed by induction. Suppose a deterministic set \( \mathcal{R} \) of states has the same distribution for futures of length \( L \) as do the causal states. This implies that they have the same
distribution for futures of shorter lengths, in particular for futures of length 1. We can use that fact, in turn, to show that they must have the distribution for futures of length \( L + 1 \):

\[
P(S_{L+1} = s^L a | \eta(s)) = P(s_{L+1} = a | \eta(s), S = s^L) P(s = s^L | R = \eta(s)) = P(S_{L+1} = a | R = \eta(s), S = s^L) P(S = s^L | R = \eta(s)).
\]

Determinism implies

\[
P(S_{L+1} = a | R = \eta(s), S = s^L) = P(S = a | R = \eta(s s^L)).
\]

So

\[
P(S_{L+1} = s^L a | R = \eta(s)) = P(S = a | R = \eta(s s^L)) P(S = s^L | R = \eta(s)) = P(S = a | R = \eta(s s^L)) P(S = s^L | R = \eta(s)).
\]

\[\square\]

Since the causal states fulfill the hypotheses of the proposition and are minimal among all prescient sets of states, they are also minimal among all sets of states that are deterministic and homogeneous for futures of length 1.

### 2.2 Recurrent, Transient, and Synchronization States

Proposition 4 tells us that the causal states form a Markov process. The states are therefore either recurrent, i.e., returned to infinitely often, or transient, visited only finitely often with positive probability (Grimmett and Stirzaker, 1992). For us, the recurrent states represent the actual causal structure of the process and, as such, they are what we are truly interested in (Upper, 1997). The most important class of transient states, and indeed the only ones encountered in practice, are the synchronization states, which can never be returned to, once a recurrent state has been visited. The synchronization states represent observational histories that are insufficient to fix the process in a definite recurrent state. Given the recurrent states, there is a straightforward algorithm (Feldman and Crutchfield, 1998) which finds the synchronization states. Here, however, we prefer to omit them from our algorithm’s final results altogether. While the algorithm will find them, it will also prune them, reporting only the truly structural, recurrent states.

For a complete taxonomy of causal states, including a discussion of issues of synchronization and reachability, see Upper (1997). For more exploration of synchronization, see Crutchfield and Feldman (2001a,b).

### 3. Causal-State Splitting Reconstruction

Here we introduce the Causal-State Splitting Reconstruction (CSSR) algorithm that estimates an \( \epsilon \)-machine from samples of a process. The algorithm is designed to respect the essential properties of causal states just outlined.

The basic idea of CSSR is straightforward and similar to state-splitting methods for finite-state machines (Lind and Marcus, 1995). It starts out assuming a simple model for the process and elaborates model components (adds states and transitions) only when statistically justified. More specifically, CSSR begins assuming the process is independent, identically distributed (IID) over the alphabet \( \mathcal{A} \). This is equivalent to assuming the process is structurally simple and is as random as possible. One can work through Definition 1 given above for causal states to show that an IID process
has a single causal state. Thus, initially the process is seen as having zero statistical complexity \((C_\mu = H[S] = \log_2 1 = 0)\) and high entropy rate \(h_\mu \leq \log_2 k\). From this starting point, CSSR uses statistical tests to see when it must add states to the model, which increases the estimated complexity, while lowering the estimated entropy rate. The initial model is kept only if the process actually is IID. This leads us to expect that CSSR converges “from below” to the process’s true model, in the sense that the number of states grows from 1 to the true value; likewise the estimated complexity \(C_\mu\) grows from 0 to the true complexity. We justify this view in our analysis of CSSR’s convergence properties below and demonstrate it empirically in Part II.

A key and distinguishing property of CSSR is that it maintains homogeneity of the causal states and determinism of the state-to-state transitions as the model grows. The result is that at each stage the estimated model is an \(\epsilon\)-machine, satisfying the criteria above, for an approximation of the process being modeled. One important consequence is that the degree of unpredictability (the process’s entropy rate \(h_\mu\)) can be directly estimated from the approximate \(\epsilon\)-machine.\(^5\)

### 3.1 The Algorithm

Assume we are given a sequence of length \(N\) over the finite alphabet \(A\). We wish to estimate from this a class \(\hat{S}\) of effective states. Each member \(\hat{S}\) of \(\hat{S}\) is a set of histories. Say that a string \(w\) is a suffix of the history \(s\) if \(s^{-L} = w\) for some \(L\), i.e., if the end of the history matches the string. To emphasize that a given string is a suffix, we write it as \(*w\); e.g., \(*01011\). We represent a state \(\hat{S}\) as a set of suffixes. The function \(\hat{S}\) maps a finite history \(\hat{s}\) to that \(\hat{S}\) which contains a suffix of \(\hat{s}\). We shall arrange it so that the assignment of histories to states is never ambiguous.

One suffix \(*w\) is the child of another \(*v\), if \(w = av\), where \(a\) is a single symbol. That is, a child suffix is longer, going into the past, by one symbol than its parent. A suffix \(*w\) is a descendant of its ancestor \(*v\) if \(w = av\), where \(v\) is any (non-null) string.

In addition to a set of suffixes, each \(\hat{S}\) is associated with a distribution for the next observable \(S^{-1}\); i.e., \(P(S^{-1} = a|\hat{S} = \hat{S})\) is defined for each \(a \in A\) and each \(\hat{S}\). We call this conditional distribution the states’s morph.

The null hypothesis is that the process being modeled is Markovian on the basis of the states in \(\hat{S}\); that is,

\[
P(S^{-1} = aS^{-L-1}) = P(S^{-1} = \hat{S} = \hat{S}(S^{-L-1})) \quad \text{(4)}
\]

Naturally, one can apply a standard statistical test — e.g., the Kolmogorov-Smirnov (KS) test\(^6\) — to this hypothesis at a specified significance level, denoted \(\alpha\). If one uses the KS test, as we do here, one avoids directly estimating the morph conditional distribution and simply uses empirical frequency counts. Recall that the significance level is the probability of type-I error (rejecting the null when it is true). Generally, the KS test has higher power than other, similar tests, such as \(\chi^2\) (Rayner and Best, 1989). That is, the KS test has a lower probability of type-II error, of accepting the null hypothesis when it is false. We will see in Part II that the algorithm is not very sensitive to the precise significance test employed.

We modify the set \(\hat{S}\) only when the null hypothesis is rejected. When we reject the null hypothesis, we fall back on a restricted alternative hypothesis, which is that we have the right set of conditional distributions, but have assigned them to the wrong histories. We therefore try to assign child suffixes whose morphs differ from their parents to existing states. Only if this alternative is itself rejected do we create additional, new distributions to explain or capture the apparent non-Markovianess. This increases the cardinality of \(\hat{S}\).

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\(^5\) In general this calculation cannot be done directly for standard HMMs, which are nondeterministic (Blackwell and Koopmans, 1957).

Throughout, we shall write $\nu(A)$ for the number of times the event $A$ happens in the data $s^N$ — the count of $A$.

Thus, there are four CSSR parameters: the measurement alphabet size $k$, the length $N$ of the data stream $s^N$, the length $L_{\text{max}}$ of the longest history to be considered, and the significance level $\alpha$ of the null-hypothesis statistical test. There are three procedures in CSSR: Initialize, Homogenize, and Determinize.

I. Initialize. Set $L = 0$ and $\tilde{\mathcal{S}} = \{\tilde{\sigma}_0\}$, where $\tilde{\sigma}_0 = \{\ast \lambda\}$; i.e., $\tilde{\sigma}_0$ contains only the null sequence $\lambda$. We regard $\ast \lambda$ as a suffix of any history, so that initially $\tilde{e}$ maps all histories to $\tilde{\sigma}_0$. The morph of $\tilde{\sigma}_0$ is defined by

$$P(\tilde{S} = a|\tilde{\mathcal{S}} = \tilde{\sigma}_0) = P(\tilde{S}^{-1} = a),$$

so the initial model is that the process is a sequence of independent, identically-distributed random variables. As a consequence, the statistical complexity vanishes ($C(\tilde{\mathcal{S}}) \log_2 1 = 0$) and the entropy rate is maximal ($h(\tilde{\mathcal{S}}) = H[\tilde{S}^{-1} = a] \leq \log_2 k$).

II. Homogenize. We first generate states whose members are homogeneous for the next symbol — states whose histories all lead to the same morph. Said differently, we generate states whose member histories have no significant differences in their individual morphs. We do this as follows.

1. For each $\tilde{\sigma} \in \tilde{\mathcal{S}}$, calculate $\hat{P}_N(\tilde{S}^{-1}|\tilde{\mathcal{S}} = \tilde{\sigma})$ — the future distribution from that state, given the data sequence.

   (a) When $L = 0$ and the only suffix is $\ast \lambda$ and we could have seen any history; so we use Eq. (5) above.

   (b) For each sequence $\tilde{s}^L \in \tilde{\sigma}$, estimate $\hat{P}_N(\tilde{S}^{-1} = a|\tilde{S} = \tilde{s}^L)$. The naive maximum-likelihood estimate,

   $$\hat{P}_N(\tilde{S}^{-1} = a|\tilde{S} = \tilde{s}^L) = \frac{\nu(\tilde{S}^{-1} = a, \tilde{S} = \tilde{s}^L)}{\nu(\tilde{S} = \tilde{s}^L)},$$

   is simple and well adapted to the later parts of the procedure, but other estimators could be used. This distribution is the morph of the history $\tilde{s}^L$.

   (c) The morph of the state $\tilde{\sigma}$ is the weighted average of the morphs of its histories $\tilde{s}^L \in \tilde{\sigma}$, with weights proportional to $\nu(\tilde{S} = \tilde{s}^L)$:

   $$\hat{P}_N(\tilde{S}^{-1} = a|\tilde{\mathcal{S}} = \tilde{\sigma}) = \frac{1}{z} \sum_{\tilde{s}^L \in \tilde{\sigma}} \nu(\tilde{S} = \tilde{s}^L) \hat{P}_N(\tilde{S}^{-1} = a|\tilde{S} = \tilde{s}^L),$$

   where $z = \sum_{\tilde{s}^L \in \tilde{\sigma}} \nu(\tilde{S} = \tilde{s}^L)$ is the number of occurrences in $s^N$ of suffixes in $\tilde{\sigma}$.

2. For each $\tilde{\sigma} \in \tilde{\mathcal{S}}$, test the null (Markovian) hypothesis. For each length-$L$ history $\tilde{s}^L \in \tilde{\sigma}$ and each $a \in A$, generate the suffix $as^L$ of length $L + 1$ — a child suffix of $\tilde{s}^L$.

   (a) Estimate the morph of $as^L$ by the same method as used above, Eq. (6).

   (b) If the morphs of $as^L$ and $\tilde{\sigma}$ do not differ according to the significance test, add $as^L$ to $\tilde{\sigma}$.
(c) If they do differ, then test whether there are any states in $\tilde{S}$ whose morphs do not differ significantly from that of $a^{s_i}_{-L}$. If so, add $a^{s_i}_{-L}$ to the state whose morph its morph matches most closely, as measured by the score of the significance test. (This is the “restricted alternative hypothesis” mentioned above.)

(d) However, if the morph of $a^{s_i}_{-L}$ is significantly different from the morphs of all existing states, then create a new state and add $a^{s_i}_{-L}$ to it, along with its morph.

(e) Recalculate the morphs of states from which sequences have been added or deleted.

3. Increment $L$ by one.

4. Repeat steps 1–3 until reaching the maximum history length $L_{\text{max}}$.

At the end of state homogenization, no history is in a state whose morph is significantly different from its own. Moreover, every state’s morph is significantly different from every other state’s morph. The causal states have this property, but their transitions are also deterministic and so we need another procedure to “determinize” $\tilde{S}$ (see Proposition 7).

III. Determinize.

1. Eliminate transient states from the current state-transition structure, leaving only recurrent states.

2. For each state $\tilde{\sigma} \in \tilde{S}$:

   (a) For each $a \in A$:

      i. Calculate $\tilde{\sigma}(s^\sigma a)$ for all $s^\sigma \in \tilde{\sigma}$ — these are the successor states on symbol $a$ of the histories $s^\sigma$ — by finding $\tilde{\sigma}' \in \tilde{S}$ such that $(s^\sigma a)^L_{-1} \in \tilde{\sigma}'$.

      ii. If there are $n > 1$ successor states on $a$, create $n - 1$ new states, each with $\tilde{\sigma}$’s ($L = 1$) morph. Partition histories in $\tilde{\sigma}$ between $\tilde{\sigma}$ and the new states so that all histories in $\tilde{\sigma}$ and the new states have the same successor on $a$. Go to i.

   (b) If every history $s^\sigma \in \tilde{\sigma}$ has the same successor on $a$, for every $a$, go on to the next state.

3. From the new, deterministic states eliminate those which are transient.

Since this procedure only produces smaller (fewer-suffix) states, and a state with one suffix string in it cannot be split, the procedure terminates, if only by assigning each history its own state. When it terminates, $\tilde{S}$ will be a set of states with deterministic transitions. Moreover, since we create deterministic states by splitting homogeneous states, the deterministic states remain homogeneous.

Now, as we noted, the causal states are the minimal states that have a homogeneous distribution for the next symbol and are deterministic. If we had access to the exact conditional distributions from the underlying process, therefore, and did not have to estimate the morphs, this procedure would return the causal states. Instead, it returns a set of states that, in a sense set by the chosen significance test, cannot be statistically distinguished from them.

3.2 Example: The Even Process

To illustrate the workings of CSSR, let’s see how it reconstructs an easy-to-describe process with two states — the even process of Weiss (1973) — that, despite its simplicity, has several nontrivial properties. We use data from a simulation of the process and typical parameter settings for the algorithm.

7. Actually, to which of these states $a^{s_i}_{-L}$ is assigned is irrelevant in the limit where $N \to \infty$; but the choice we use here is convenient, plausible, and can be implemented consistently.
Figure 1: The even process: a strictly sofic, non-Markovian stochastic process.

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</tr>
<tr>
<td>101</td>
<td>837</td>
<td>0101</td>
<td>0</td>
<td>1101</td>
<td>836</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>110</td>
<td>1654</td>
<td>0110</td>
<td>814</td>
<td>1110</td>
<td>841</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>111</td>
<td>3378</td>
<td>0111</td>
<td>841</td>
<td>1111</td>
<td>2537</td>
<td></td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

Table 1: Count statistics for words of length 4 and shorter from $10^4$ samples of the even process of Fig. 1. The total count at each length is 9,996; 9996 words.

We have a two-symbol alphabet, $A = \{0, 1\}$. There are two recurrent states, labeled $A$ and $B$ in Fig. 1. State $A$ can either emit a 0 and return to itself, or emit a 1 and go to $B$; we take the version where these options are equally likely. State $B$ always emits a 1 and goes to $A$. The labeled transition matrices $T$ are thus

$$
T^{(0)} = \begin{bmatrix} 0.5 & 0 \\ 0 & 0 \end{bmatrix} \quad \text{and} \quad T^{(1)} = \begin{bmatrix} 0 & 0.5 \\ 1.0 & 0 \end{bmatrix}.
$$

(8)

We take $L_{\max} = 3$ and $\alpha = 0.01$. We simulated the even process for $10^4$ time steps and accumulated the sequence statistics for words of length 4 and shorter, given in Table 1. Since one slides a window of size $L_{\max} + 1 = 4$ through the data stream, there are 9,996 length-4 words. Observe in Table 1 that sequences containing odd-length blocks of 1s, bounded by 0s, do not appear. That is, words of the form $\{012k+1\}0$, $k = 1, 2, \ldots$ are forbidden.

Overall, the data contains 3309 0s and 6687 1s, since for simplicity we fix the total number of words at each length to be 9,996. The initial state $A_{L=0}$ formed at $L = 0$, containing the null suffix $*\lambda$, therefore produces 1s with probability $\approx 0.669$.

The null suffix $*\lambda$ has two children, $*0$ and $*1$. At $L = 1$ the probability of producing a 1, conditional on the suffix $*0$, is $P(1|*0) \approx 1655/3309 \approx 0.500$, which is significantly different from the distribution for the null suffix. Similarly, the probability of producing a 1, conditional on the suffix $*1$, is $P(1|*1) \approx 5032/6690 \approx 0.752$, which is also significantly different from that of the parent state. We thus produce two new states $B_{L=1}$ and $C_{L=1}$, one containing the suffix $*1$ and one the suffix $*0$, respectively. There is now a total of three states: $A_{L=0} = \{*\lambda\}$, $B_{L=1} = \{*1\}$, and $C_{L=1} = \{*0\}$.

Examining the second generation of children at $L = 2$, one finds the conditional probabilities of producing 1s given in Table 2. The morphs of the first two suffixes, $*00$ and $*10$, do not differ
Table 2: Conditional counts having read two symbols.

| Suffix *w | P(1 | * w) |
|-----------|---------|
| *10       | 837/1655 ≈ 0.506 |
| *01       | 1655/1655 = 1.000 |
| *11       | 3378/5032 ≈ 0.671 |

Table 3: Conditional counts for child suffixes of state $C_{L=1}$’s $L = 2$ suffixes.

| Suffix *w | P(1 | * w) |
|-----------|---------|
| *000      | 422/836 ≈ 0.505 |
| *100      | 396/818 ≈ 0.484 |
| *010      | 0/837 = 0.0  |
| *110      | 837/1655 ≈ 0.506 |

Table 4: Conditional counts for child suffixes of state $D_{L=2}$.

| Suffix *w | P(1 | * w) |
|-----------|---------|
| *001      | 818/818 = 1.000 |
| *101      | 837/837 = 1.000 |
Pattern Discovery in Time Series I

| Suffix *w | P(1 | * w) |
|-----------|----------|
| *011      | 0.508    |
| *111      | 0.751    |

Table 5: Conditional counts for child sufixes of state $A_{L=0}$'s $L = 2$ sufix *11.

Figure 2: The full $\epsilon$-machine reconstructed from $10^4$ samples of the even process of Fig. 1. State $A$ (inscribed circle) is the unique start state.

3.3 Time Complexity

Procedure I (Initialize) takes a time proportional to the number $N$ of symbols in the data stream.

Procedure II (Homogenize) is a little subtler. For each history, checking whether it belongs to the same state as its parent, repartitioning, and the creation of a new state (if needed) can all be done in constant time. However, removing ancestors after repartitioning or state-creation takes a time proportional to (at most) $L_{\text{max}}$, since there can be no more than $L_{\text{max}}$ ancestors. Since there are at most $k^L_{\text{max}}$ sufixes, the total time for Procedure II is proportional to $L_{\text{max}}k^L_{\text{max}}$.

The time needed for Procedure III (Determinize) can be divided into three parts: collecting the transition structure, removing transient states, and determinizing. (We remove transient states twice, but this only introduces a constant factor.) The time to collect the transition structure is at most $kk^L_{\text{max}}$. The time to remove transients is $s^2k$, where $s$ is the number of states (before removal).
The time needed to make one determinizing pass is \( k \cdot L \), and the maximum number of passes needed is \( k \cdot L_{\text{max}} \), since this is the worst case for the number of states which could be made. So the worst-case time for determinization is \( k \cdot 2 \cdot L_{\text{max}} + 1 \). Now, \( s \) is \( k \cdot L_{\text{max}} \), at most, so the overall time for Procedure III is \( O(k \cdot 2 \cdot L_{\text{max}} + 1) + O(N) \), which asymptotically is \( O(k \cdot 2 \cdot L_{\text{max}} + 1) + O(N) \). Observe that this is linear in the data size \( N \). It is exponential in the alphabet size \( k \), but the exponent of \( 2 \cdot L_{\text{max}} + 1 \) is very much a loose worst-case result. It applies only in extreme cases; e.g., when every string spawns its own state, almost all of which are transient. In practice, the run time is much shorter than this bound would lead one to fear. Average-case results replace the number of strings, here bounded by \( k \cdot L_{\text{max}} \), with the typical number of distinct sequences of length \( L_{\text{max}} \), the process generates with positive probability, which is \( 2^{L_{\text{max}}} \) (Cover and Thomas, 1991). Similarly, bounding \( s \), the number of states, by \( k \cdot L_{\text{max}} \) is generally excessive and better approximated by \( 2^C \).

4. Reliability and Rates of Convergence

We wish to show that the estimated \( \epsilon \)-machines produced by the CSSR algorithm converge to the original process’s true \( \epsilon \)-machine. Specifically, we will show that the set \( \hat{S} \) of states it estimates converges in probability to the correct set \( S \) of causal states: i.e., the probability that \( \hat{S} \neq S \) goes to zero as \( N \to \infty \). To do this, we first look at the convergence of the empirical conditional frequencies to the true morphs and then use this to show that the set \( \hat{S} \) of states converges. Finally, we examine the convergence of predictions once the causal states are established and say a few words about how, from a dynamical-systems viewpoint, the \( \epsilon \)-machine acts as a kind of attractor for the algorithm.

Throughout, we make the following assumptions:

1. The process is conditionally stationary. Hence the ergodic theorem for stationary processes applies (Grimmett and Stirzaker, 1992, Sec. 9.5) and time averages converge almost surely to the state-space average for whatever ergodic component the process happens to be in.

2. The original process has only a finite number of causal states.

3. Every state contains at least one suffix of finite length. That is, there is some finite \( L \) such that every state contains a suffix of length no more than \( L \). This does not mean that \( L \) symbols of history always suffice to determine the state, just that it is possible to synchronize — in the sense of Crutchfield and Feldman (2001a) — to every state after seeing no more than \( L \) symbols.

We also estimate conditional probabilities (morphs) by simple maximum likelihood. One can substitute other estimators — e.g., maximum entropy — and only the details of the convergence arguments would change.

4.1 Convergence of the Empirical Conditional Probabilities to the Morphs

The past \( \overrightarrow{S} \) and the future \( \overleftarrow{S} \) of the process are independent given the causal state \( \sigma \in S \) (Prop. 3). More particularly, the past \( \overrightarrow{S} \) and the next future symbol \( \overleftarrow{S}^{-1} \) are independent given the causal state. Hence, \( \overrightarrow{S} \) and \( \overleftarrow{S} \) are independent, given a history suffix sufficient to fix us in a cell of the causal-state partition. Now, the morph for that suffix is a multinomial distribution over \( A \), which gets sampled independently each time the suffix occurs. Our estimate of the morph is the empirical distribution obtained by IID samples from that multinomial. Write this estimated distribution as \( \hat{P}_N(\overrightarrow{S} = a | \overleftarrow{S} = s^{-1}) \), recalling that \( P(\overrightarrow{S} = a | \overleftarrow{S} = s^{-1}) \) denotes the true morph. Does \( \hat{P}_N \to P \) as \( N \) grows?
Since we use the KS test, it is reasonable to employ the variational distance. Given two distributions $P$ and $Q$ over $A$, the variational distance is

$$d(P, Q) = \sum_{a \in A} |P(X = a) - Q(X = a)|.$$  \hfill (9)

Scheffé showed that

$$d(P, Q) = 2 \max_{A \subseteq 2^A} |P(X \in A) - Q(X \in A)|,$$  \hfill (10)

where $2^A$ is the power-set of $A$, of cardinality $2^k$ (Devroye and Lugosi, 2001).

Chernoff’s inequality (Vidyasagar, 1997) tells us that, if $X_1, X_2, \ldots, X_n$ are IID Bernoulli random variables and $A_n$ is the mean of the first $n$ of the $X_i$, with probability $\mu$ of success, then

$$P(|A_n - \mu| \geq t) \leq 2e^{-2nt^2}.$$  \hfill (11)

Applying this to our estimation problem and letting $n = \nu(s^L)$ be the number of times we have seen the suffix $s^L$, we have:

$$P(d(\hat{P}_n(s^L | S = s^L), P(s^L | S = s^L)) \geq t)$$

$$= P \left( t < \sum_{a \in A} \left| \hat{P}_n(s^L = a | S = s^L) - P(s^L = a | S = s^L) \right| \right)$$

$$= P \left( 2t < \max_{A \subseteq 2^A} \left| \hat{P}_n(s^L \in A | S = s^L) - P(s^L \in A | S = s^L) \right| \right)$$

$$\leq \sum_{A \subseteq 2^A} P \left( \left| \hat{P}_n(s^L \in A | S = s^L) - P(s^L \in A | S = s^L) \right| > 2t \right)$$

$$\leq \sum_{A \subseteq 2^A} 2e^{-2n(2t)^2}$$

$$= 2^{k+1} e^{-8nt^2}.$$  \hfill (17)

Thus, the empirical conditional distribution (morph) for each suffix converges exponentially fast to its true value, as the suffix count $n$ grows.

Suppose there are $s$ suffixes across all the states. We have seen the $i^{th}$ suffix $n_i$ times; abbreviate the $s$-tuple $(n_1, \ldots, n_s)$ by $n$. We want the empirical distribution associated with each suffix to be close to the empirical distribution of all the other suffixes in its state. We can ensure this by making all the empirical distributions close to the state’s morph. In particular, if all distributions are within $t$ of the morph, then (by the triangle inequality) every distribution is within $2t$ of every other distribution. Call the probability that this is not true $q(t, n)$. This is the probability that at least one of the empirical distributions is not within $t$ of the state’s true morph. $q(t, n)$ is at most the sum of the probabilities of each suffix being an outlier. Hence, if there are $s$ suffixes in total, across all the states, then the probability that one or more suffixes differs from its true morph by $t$ or more is

$$q(t, n) \leq \sum_{i=1}^{s} 2ke^{-8n_i t^2}$$

$$\leq 2^{k+1} 8e^{-8nt^2},$$  \hfill (19)

where $m$ is the least of the $n_i$. Now, this formula is valid whether we interpret $s$ as the number of histories actually seen or as the number of histories needed to infer the true states. In the first

8. This metric is compatible with most other standard tests, too.
case, Eq. (19) tells us how accurate we are on the histories seen; in the latter, over all the ones we need to see. This last is clearly what we need to prove overall convergence, but knowing \( s \), in that sense, implies more knowledge of the process’s structure than we start with. However, we can upper bound \( s \) by the maximum number of morphs possible -- \( s \leq (k^{L+1} - 1)/(k-1) \) -- so we only need to know (or, in practice, pick) \( L \).

Which string is least-often seen — which \( n_i \) is \( m \) — generally changes as more data accumulates. However, we know that the frequencies of strings converge almost surely to probabilities as \( N \to \infty \). Since there are only a finite number of strings of length \( L \) or less, it follows that the empirical frequencies of all strings also converge almost surely (den Hollander, 2000). (The rate of convergence depends on the entropy \( D(P|P) \) of the empirical distribution relative to the true.) Our observed process is a random function of the process’s causal states. Now, for any pair of distributions \( P(X) \) and \( Q(X) \) and any measurable function \( f : X \to Y \),

\[
d(P(X), Q(X)) \geq d(P(Y), Q(Y)) ,
\]

whether \( f \) is either a deterministic or a random function. Hence, the empirical sequence distribution, at any finite length, converges to the true distribution at least as quickly as the state distributions. That is, they converge with at least as large an exponent. Therefore, under our assumptions, for each \( i \), \( n_i/N \to p_i \) exponentially fast — there is an increasing function \( r(\varepsilon) \), \( r(0) = 0 \), and a constant \( C \) such that, for each \( \varepsilon \geq 0 \),

\[
P \left( \left| \frac{n_i}{N} - p_i \right| > \varepsilon \right) \leq Ce^{-N\varepsilon} .
\]

\( r(\varepsilon) \) is known as the large-deviation rate function.) The probability that \( m \leq N(p^* - \varepsilon) \) is clearly no more than the probability that, for one or more \( i \), \( n_i \leq N(p^* - \varepsilon) \). Since the lowest-probability strings need to make the smallest deviations from their expectations to cause this, the probability that at least one \( n_i \) is below \( N(p^* - \varepsilon) \) is no more than \( k\^L \) times the probability that the count of the least probable string is below \( N(p^* - \varepsilon) \). The probability that an empirical count \( n_i \) is below its expectation by \( \varepsilon \), in turn, is less than the probability that it deviates from its expectation by \( \varepsilon \) in either direction. That is,

\[
P(m \leq N(p^* - \varepsilon)) \leq P \left( \bigcup \n_i \leq N(p^* - \varepsilon) \right) \leq \sum \n_i \leq N(p^* - \varepsilon) \leq k^L P(n^* \leq N(p^* - \varepsilon)) \leq k^L P \left( \left| \frac{n^*}{N} - p^* \right| > \varepsilon \right) \leq Ck^L e^{-N\varepsilon} .
\]

9. Write \( F_i \) for the event that the frequencies of string \( i \) fail to converge to the probability. From the ergodic theorem, \( P(F_i) = 0 \) for each \( i \). If there are \( s \) strings, the event that represents one or more failures of convergence is \( \bigcup_{i=1}^s F_i \). Using Bonferroni’s inequality, \( P(\bigcup_{i=1}^s F_i) \leq \sum_{i=1}^s P(F_i) = s \cdot 0 = 0 \). Hence, all \( s \) strings converge together with probability 1.
With a bound on \( m \), we can fix an overall exponential bound on the error probability \( q \):

\[
q(t, n) \leq \inf_{r \geq 0} C 2^{k+1} k^L \left( \frac{KL}{k - 1} - 1 \right) e^{-8N(p^* - \epsilon)^2} e^{-Nr(\epsilon)}.
\]

(28)

Solving out the infimum would require knowledge of \( r(\epsilon) \), which is difficult to come by. Whatever \( r(\epsilon) \) may be, however, the bound is still exponential in \( N \).

The above bound is crude for small \( N \) and especially for small \( t \). In the limit \( t \to 0 \), it tells us that a probability is less than some positive integer, which, while true, is not at all sharp. It becomes less crude, however, as \( N \) and \( t \) grow. In any case, it suffices for the present purposes.

We can estimate \( p^* \) from the reconstructed \( \epsilon \)-machine by calculating its fluctuation spectrum (Young and Crutchfield, 1993). Young and Crutchfield demonstrated that the estimates of \( p^* \) obtained in this way become quite accurate with remarkably little data, just as, we shall see in Part II, estimates of the entropy rate \( h_\mu \) do. At the least, calculating \( p^* \) provides a self-consistency check on the reconstructed \( \epsilon \)-machine.

4.2 Analysis of Error Probabilities

Let us consider the kinds of statistical error each of the algorithm’s three procedures can produce.

Since it merely sets up parameters and data structures, nothing goes wrong in Procedure I (Initialize).

Procedure II (Homogenize) can make two kinds of errors. First, it can group together histories with different distributions for the next symbol. Second, it can fail to group together histories that have the same distribution. We will analyze both cases together.

It is convenient here to introduce an additional term. By analogy with the causal states, the precausal states are defined by the following equivalence relation: Two histories are precausally equivalent when they have the same length-1 morph. The precausal states are then the coarsest states (partition) that are homogeneous for \( S \). The causal states are either the same as the precausal states or a refinement of them. Procedure II ought to deliver the correct partition of histories into precausal states.

Suppose \( \overline{s}_i \) and \( \overline{s}_j \) are suffixes in the same state, with counts \( n_i \) and \( n_j \). No matter how large their counts, there is always some variational distance \( t \) such that the significance test will not separate estimated distributions differing by \( t \) or less. If we make \( n_i \) large enough, then, with probability arbitrarily close to one, the estimated distribution for \( \overline{s}_i \) is within \( t/2 \) of the true morph, and similarly for \( \overline{s}_j \). Thus, the estimated morphs for the two suffixes are within \( t \) of each other and will be merged. Indeed, if a state contains any finite number of suffixes, by obtaining a sufficiently large sample of each, we can ensure (with arbitrarily high probability) that they are all within \( t/2 \) of the true morph and so within \( t \) of each other and thus merged. In this way, the probability of inappropriate splitting can be made arbitrarily small.

If each suffix’s conditional distribution is sufficiently close to its true morph, then any well behaved test will eventually separate suffixes that belong to different morphs. More concretely, let the variational distance between the morphs for the pair of states \( \sigma_i \) and \( \sigma_j \) be \( d_{ij} \). A well behaved test will distinguish two samples whose distance is some constant fraction of this or more — say, \( 3d_{ij}/4 \) or more — if \( n_i \) and \( n_j \) are large enough. By taking \( n_i \) large enough, we can make sure, with probability arbitrarily close to 1, that the estimated distribution for \( \sigma_i \) is within some small distance of its true value — say, \( d_{ij}/8 \). We can do the same for \( \sigma_j \) by taking \( n_j \) large enough. Therefore, with probability arbitrarily close to one, the distance between the estimated morphs for \( \sigma_i \) and \( \sigma_j \) is at least \( 3d_{ij}/4 \), and \( i \) and \( j \) are, appropriately, separated. Hence, the probability of erroneous non-separations can be made as small as desired.

Therefore, by taking \( N \) large enough, we can make the probability that the correct precausal states are inferred arbitrarily close to 1.
If every history is correctly assigned to a precausal state, then nothing can go wrong in Procedure III (Determinize). Take any pair of histories \( \tilde{s}_1 \) and \( \tilde{s}_2 \) in the same precausal state: either they belong to the same causal state or they do not. If they do belong to the same causal state, then by determinism, for every string \( w \), \( \tilde{s}_1 w \) and \( \tilde{s}_2 w \) belong to the same causal state. Since the causal states are refinements of the precausal states, this means that \( \tilde{s}_1 w \) and \( \tilde{s}_2 w \) also belong to the same precausal state. Contrarily, if \( \tilde{s}_1 \) and \( \tilde{s}_2 \) belong to different causal states, they must give different probabilities for some strings. Pick the shortest such string \( w \) (or any one of them, if there is more than one) and write it as \( w = va \), where \( a \) is a single symbol. Then the probability of \( a \) depends on whether we saw \( \tilde{s}_1 v \) or \( \tilde{s}_2 v \). So \( \tilde{s}_1 v \) and \( \tilde{s}_2 v \) have distinct length-1 morphs and belong to different precausal states. Hence, determinization will separate \( \tilde{s}_1 \) and \( \tilde{s}_2 \), since, by hypothesis, the precausal states are correct. Thus, histories will always be separated, if they should be, and never, if they should not.

Since determinization always refines the partition with which it starts and the causal states are a refinement of the precausal states, there is no chance of merging histories that do not belong together. Hence, Procedure III will always deliver the causal states, if it starts with the precausal states. We will not examine the question of whether Procedure III can rectify a mistake made in Procedure II. Experientially, this depends on the precise way determinization is carried out and, most typically, if the estimate of the precausal states is seriously wrong, determinization only compounds the mistakes. Procedure III does not, however, enhance the probability of error.

To conclude: If the number of causal states is finite and \( L \) is sufficiently large, the probability that the states estimated are not the causal states becomes arbitrarily small, for sufficiently large \( N \). Hence the CSSR algorithm, considered as an estimator, is (i) consistent (Cramér, 1945), (ii) probably approximately correct (Kearns and Vazirani, 1994, Vapnik, 2000), and (iii) reliable (Spirtes et al., 2001, Kelly, 1996), depending on which field one comes from.

### 4.3 Dynamics of the Learning Algorithm

We may consider the CSSR algorithm itself to be a stochastic dynamical system, moving through a state space of possible history-partitions or \( \varepsilon \)-machines. What we have seen is that the probability that CSSR does not infer the correct states (partition) — that it does not have the causal architecture right — drops exponentially as time goes on. By the Borel-Cantelli lemma, therefore, CSSR outputs \( \varepsilon \)-machines that have the wrong architecture only finitely many times before fixing on the correct architecture forever. Thus, the proper architecture is a kind of absorbing region in \( \varepsilon \)-machine space. In fact, almost all of the algorithm’s trajectories end up there and stay. (There may be other absorbing regions, but only a measure-0 set of inference trajectories reach them.)

We are not satisfied with getting the correct causal architecture, however. We also want the morphs to converge. Here we can exploit the fact that the estimated morph for each causal state is an average over the histories it contains. If there are \( s_c \) causal states and the least frequently sampled one has been seen \( m_o \) times, then reasoning parallel to the above tells us that the probability any of our estimated morphs differs from its true morph by \( t \) or more is at most \( 2^{k+1}s_c\exp(-8m_0t^2) \). Moreover, since the causal states form an irreducible Markov chain, \( m_o \) will converge exponentially quickly to \( Np^*_c \).

As a general note, while the probability of errors of a given size goes down exponentially with \( N \), this does not imply that the expectation of the error is exponentially decreasing. Rather, the expected variation difference between the true and empirical distributions for a multinomial goes as \( N^{-1/2} \) (Devroye and Lugosi, 2001). Since \( N \) appears in the form for the error probability only in the combination \( Np^*_c \), we expect a variational error scaling as \( 1/\sqrt{Np^*_c} \). Now, the asymptotic equipartition theorem (Cover and Thomas, 1991) tells us that, for large \( L \), every string that appears

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10. Otherwise, \( \tilde{s}_1 \) and \( \tilde{s}_2 \) must assign different probabilities to \( v \), and so \( w = va \) is not the shortest string on which they differ.
with positive probability has a probability of approximately $Lh_\mu$. Hence $p^* \to Lh_\mu$, too. This suggests that, for large $N$ and large $L$, the variational error should scale as $1/\sqrt{N L h_\mu}$. Readers may judge the quality of the data-collapse of the rescaled error for themselves in the tests presented in Part II.

4.4 State-Merging $\epsilon$-Machine Inference Algorithms

Existing $\epsilon$-machine reconstruction procedures use what one might call state compression or merging. The default assumption is that each distinct history encountered in the data is a distinct causal state. Histories are then merged into causal states when their morphs are close. Kindred merging procedures can learn hidden Markov models (Stolcke and Omohundro, 1993) and finite automata (Trakhtenbrot and Barzdin, 1973, Murphy, 1996).

The standard $\epsilon$-machine inference algorithm is the subtree-merging algorithm introduced by Crutchfield and Young (Crutchfield and Young, 1989, Crutchfield, 1992). The algorithm begins by building a $k$-ary tree of some pre-set depth $D$, where paths through the tree correspond to sequences of observations of length $D$, obtained by sliding a length-$D$ window through the data stream (or streams, if there are several). If $D = 4$, say, and the sequence abba is encountered, the path in the tree will start at the root node, take the edge labeled $a$ to a new node, then take the outgoing edge labeled $b$ to a third node, then the edge labeled $b$ from that, and finally the edge labeled $a$ to a fifth node, which is a leaf. An edge of the tree is labeled, not just with a symbol, but also with the number of times that edge has been traversed in scanning through the data stream. Denote by $\nu(a_i|n)$ the number on the $a_i$ edge going out of node $n$ and $N$ the total number of sequences entered into the tree. The tree is a $D$-block Markov model of the process. Each level $l$ gives an estimate of the distribution of length-$l$ words in the data stream.

The traversal-counts are converted into empirical conditional probabilities by normalization:

$$\hat{P}_N(a_i|n) = \frac{\nu(a_i|n)}{\sum_{a_j} \nu(a_j|n)}.$$ 

Thus, attached to each non-leaf node is an empirical conditional distribution for the next symbol. If node $n$ has descendants to depth $K$, then it has (by implication) a conditional distribution for futures of length $K$.

The merging procedure is now as follows. Consider all nodes with subtrees of depth $L = D/2$ (Crutchfield, 1992). Take any two of them. If all the empirical probabilities attached to the length-$L$ path in their subtrees are within some constant $\delta$ of one another, then the two nodes are equivalent. They should be merged with one another. The new node for the root will have incoming links from both the parents of the old nodes. This procedure is to be repeated until no further merging is possible.\(^{11}\) It is clear that, given enough data, a long enough $L$, and a small enough $\delta$, the subtree algorithm will converge on the causal states.

All other methods for $\epsilon$-machine reconstruction currently in use are also based on merging. Take, for instance, the “topological” or “modal” merging procedure of Perry and Binder (1999). They consider the relationship between histories and futures, both (in the implementation) of length $L$. Two histories are assigned to the same state if the sets of futures that can succeed them are identical.\(^{12}\) The distribution over those futures is then estimated for each state, not for each history.

As we said, the default assumption of current state-merging methods is that each history is its own state. The implicit null model of the process is thus the most complex possible, given the length of histories available. This null model is whittled down by merging, and the estimated statistical complexity falls as $N$ grows. State-merging algorithms, so to speak, converge “from above” on the true model. In all this, they are the opposite of CSSR. State splitting starts by putting every

11. Since the criterion for merging is not a true equivalence relation (it lacks transitivity), the order in which states are examined for merging matters, and various tricks exist for dealing with this. See, e.g., Hanson (1993).

12. This is an equivalence relation, but it is not causal equivalence.
history in one state — a zero-complexity null model that is elaborated by splitting. It converges from below, both in the number of states and in the complexity. Conceivably, one could develop unbiased estimates of complexity by combining merging and splitting algorithms properly.

Unfortunately, state-merging has inherent difficulties. For instance: what is a reasonable value of morph similarity $\delta$? Clearly, as the amount of data increases and the law of large numbers makes empirical probabilities converge to true probabilities, $\delta$ should grow smaller. But it is grossly impractical to calculate what $\delta$ should be, since the null model itself is so complicated. (Current best practice is to pick $\delta$ as though the process were an IID multinomial, which is the opposite of the algorithm’s implicit null model.) In fact, using the same $\delta$ for every pair of tree nodes is unreliable. The nodes will not have been visited equally often, being associated with different tree depths, so the conditional probabilities in their subtrees vary in accuracy. An analysis of the convergence of empirical distributions, of the kind we made in Section 4.1 above, could give us a handle on $\delta$, but reveals another difficulty. CSSR must estimate $2^k$ probabilities for each history — one for each member of the power-set of the alphabet. The subtree-merging algorithm, however, must estimate the probability of each member of the power set of future sequences, i.e., $2^{k+L}$ probabilities. This is an exponentially larger number, and the corresponding error bounds would be worse by this factor.

The theorems in Shalizi and Crutchfield (2001) say a great deal about the causal states: they are deterministic, they are Markovian, and so on. No previous reconstruction algorithm made use of this information to guide its search. Subtree-merging algorithms can return nondeterministic states, for instance, which cannot possibly be the true causal states. While the subtree algorithm converges, and other merging algorithms probably do too, CSSR should do better, both in terms of the kind of result it delivers and the rate at which it approaches the correct result.

5. Conclusion

5.1 Future Directions

A number of directions for future work present themselves. Elsewhere, we have developed extensions of computational mechanics to transducers and interacting time series and to spatio-temporal dynamical systems (Crutchfield, 1994, Feldman and Crutchfield, 1998, Hanson and Crutchfield, 1997, Shalizi and Crutchfield, 2002). It is clear that the present algorithm can be applied to transducers, and we feel that it can be applied to spatiotemporal systems. Work on both extensions is currently underway.

One can easily extend the formalism of computational mechanics to processes that take on continuous values at discrete times, but this has never been implemented. Much of the machinery we employed here carries over straightforwardly to the continuous setting, e.g., empirical process theory for IID samples (Devroye and Lugosi, 2001) or the Kolmogorov-Smirnov test. The main obstacle to simply using CSSR in its present form is the need for continuous interpolation between the (necessarily finite) measurements. However, all methods of predicting continuous-valued processes must likewise impose some interpolation scheme, and perhaps one along the lines of Bosq (1998) or Fraser and Dimitriadis (1993) would work. Continuous-valued, continuous-time processes raise more difficult questions, which we shall not even attempt to sketch here.

Potentially, $\epsilon$-machines and our algorithm can be applied in any domain where HMMs have proved their value (e.g., bioinformatics (Baldi and Brunak, 1998)) or where there are poorly-understood processes generating sequential data, such as speech, in which one wishes to find nonobvious or very complex patterns.

13. It is sometimes claimed (Palmer et al., 2002) that the nondeterminism is due to nonstationarity in the data stream. While a nonstationary source can cause the subtree-merging algorithm to return nondeterministic states, the algorithm is quite capable of doing this when the source is IID.
5.2 Summary

We have presented a new algorithm for pattern discovery in time series. Given samples of a conditionally stationary process, the algorithm reliably infers the process’s causal architecture. Under reasonable conditions on the process, the algorithm almost surely returns an incorrect architecture only finitely many times. The time complexity of the algorithm is linear in the data size. We show in Part II that it works reliably on stochastic processes defined over subshifts of finite type, strictly sofic systems, and HMMs, and reasonably on infinite-state processes. In addition to the arguments made here that CSSR will consistently outperform prior causal-state-merging algorithms, in Part II we demonstrate that it performs significantly better than conventional HMM algorithms and context-tree methods.

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Appendix A. Information Theory

Our notation and terminology follows that of Cover and Thomas (1991).

Given a random variable $X$ taking values in a discrete set $A$, the entropy $H[X]$ of $X$ is

$$H[X] = - \sum_{a \in A} P(X = a) \log_2 P(X = a).$$

$H[X]$ is the expectation value of $- \log_2 P(X)$. It represents the uncertainty in $X$, interpreted as the mean number of binary distinctions (bits) needed to identify the value of $X$. Alternately, it is the minimum number of bits needed to encode or describe $X$. Note that $H[X] = 0$ if and only if $X$ is (almost surely) constant.

The joint entropy $H[X;Y]$ of two variables $X$ and $Y$ is the entropy of their joint distribution:

$$H[X;Y] = - \sum_{a \in A, b \in B} P(X = a, Y = b) \log_2 P(X = a, Y = b).$$

The conditional entropy of $X$ given $Y$ is


$H[X|Y]$ is the average uncertainty remaining in $X$, given a knowledge of $Y$.

The mutual information $I[X;Y]$ between $X$ and $Y$ is

$$I[X;Y] = H[X] - H[X|Y].$$

It gives the reduction in $X$’s uncertainty due to knowledge of $Y$ and is symmetric in $X$ and $Y$.

The entropy rate $h_\mu$ of a stochastic process $\ldots, S_{-2}, S_{-1}, S_0, S_1, S_2, \ldots$ is

$$h_\mu = \lim_{L \to \infty} \frac{1}{L} H[S_1 \ldots S_L | S_{-L} \ldots S_1] = H[S_1 | S_0].$$
(The limit always exists for conditionally stationary processes.) \( h \) measures the process’s unpredictability, in the sense that it is the uncertainty which remains in the next measurement even given complete knowledge of the past.

**References**


