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Nonlinear Time-Series Analysis

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Intelligent data analysis often requires one to extract meaningful conclusions about a complicated system using data from a single sensor. If the system is linear, a wealth of well-established, powerful time-series analysis techniques is available to the analyst. If it is not, the problem is much harder and one must resort to nonlinear dynamics theory in order to infer useful information from the data. IDA problems are often complicated by a simultaneous over-abundance and lack of data: reams of information about the voltage output of a power substation, for instance, but no data about other important quantities, such as the temperatures inside the transformers. Data-mining techniques[14] provide some useful ways to deal successfully with the sheer volume of information that constitutes one part of this problem. The second part of the problem is much harder. If the target system is highly complex — say, an electromechanical device whose dynamics is governed by three metal blocks, two springs, a pulley, several magnets, and a battery — but only one of its important properties (e.g., the position of one of the masses) is sensor-accessible, the data analysis procedure would appear to be fundamentally limited. Delay-coordinate embedding, a technique developed by the dynamics community, is a way to get around this problem; it lets one reconstruct the internal dynamics of a complicated nonlinear system from a single time series. That is, one can often use delay-coordinate embedding to infer useful information about internal (and unmeasurable) transformer temperatures using only their output voltages. The reconstruction produced by delay-coordinate embedding is not, of course, completely equivalent to the internal dynamics in all situations,
or embedding would amount to a general solution to control theory’s observer problem: how to identify all of the internal state variables of a system and infer their values from the signals that can be observed. However, a single-sensor reconstruction, if done right, can still be extremely useful because its results are guaranteed to be topologically (i.e., qualitatively) identical to the internal dynamics. This means that conclusions drawn about the reconstructed dynamics are also true of the internal dynamics of the system inside the black box.

If the system under examination is linear, the analysis is comparatively straightforward and the tools — the topic of section 1 of this chapter — are well developed. One can characterize the data using statistics (mean, standard deviation, etc.), fit curves to them (functional approximation), and plot various kinds of graphs to aid one’s understanding of the behavior. If a more-detailed analysis is required, one typically represents the system in an input + transfer function → output manner, using any of a wide variety of time- or frequency-domain models. This kind of formalism admits a large collection of powerful reasoning techniques, such as superposition and the notion of transforming back and forth between the time and frequency domains. The latter is particularly useful, as many signal processing operations are much easier in one domain than the other (e.g., manipulation of differential equations, which is trivial in the frequency domain and difficult in the time domain).

Nonlinear systems pose a special challenge to intelligent data analysis; their mathematics is vastly harder and many standard time-series analysis techniques simply do not apply. Chaotic systems, for instance, exhibit broad-band behavior, which makes many traditional signal processing operations useless. One cannot decompose problems in the standard “input + transfer function → output” manner, nor can one simply low-pass filter the data to remove noise, as the high-frequency components are essential elements of the signal. The concept of a discrete set of spectral components does not make sense in many nonlinear problems, so using transforms to move between time and frequency domains — a standard technique that lets one transform differential equations into algebraic ones and vice versa, making the former much easier to work with — does not work. For these and related reasons, nonlinear dynamicists eschew most forms of spectral analysis. Because they are soundly based in nonlinear dynamics theory and rest firmly on the formal definition of invariants, however, the analysis methods described in section 2 of this chapter do not suffer from the kinds of limitations that apply to traditional linear analysis methods.
Delay-coordinate embedding, where past values of a signal are used to reconstruct its dynamics, is particularly important for intelligent data analysis because fully observable systems are rare in engineering practice; as a rule, many — often, most — of a system’s state variables either are physically inaccessible or cannot be measured with available sensors. Worse yet, the true state variables may not be known to the user; temperature, for instance, can play an important and often unanticipated role in the behavior of an electronic circuit. Embedding, the topic of section 3 of this chapter, is a partial solution to this problem; it not only yields useful information about the behavior of the unmeasured variables, but also gives some indication of how many independent state variables actually exist inside the black box.

The objective of this chapter is to cover a representative sampling of the time-series analysis techniques that have been developed by the nonlinear dynamics community over the past decade, with a specific emphasis on how those techniques would be useful in IDA applications. Other communities, of course, have developed different approaches to nonlinear time-series analysis. One of the more famous is Tukey’s “exploratory data analysis,” a sleuthing approach that emphasizes (and supports) visual examination over blind, brute-force digestion of data into statistics and regression curves[47]. Some of the more-recent developments in this field attempt to aid — or even augment — the analyst’s abilities in unconventional ways, ranging from 3D virtual-reality displays to haptics (representing the data as a touch pattern, which has been proposed for reading mammograms[29]) or data sonification. By way of background, section 1 quickly reviews some of the traditional methods that apply to linear systems. Section 2 covers the bare essentials of dynamical systems theory and practice; these are the basis of the general theory of dynamics that applies to any system, linear or nonlinear. If all of the important properties of the target system can be identified and measured and the data are basically noise-free, these techniques, alone, can provide a very good solution to many nonlinear data-analysis problems. If there are fewer sensors than state variables, however, one must call upon the methods described in section 3 in order to reconstruct the dynamics before one can apply the section 2 methods. Noise is a much more difficult problem. There exist techniques that “filter” nonlinear time-series data, turning the nonlinearity to advantage and reducing the noise by an exponentially large factor[13], but the mathematics of this is well beyond the scope of this discussion. This chapter continues with two extended examples that demonstrate both the analysis methods of section 2 and the
Figure 1: The “input + transfer function → output” framework of traditional signal processing. Top: time domain. Bottom: frequency domain.

delay-coordinate reconstruction techniques of section 3, and concludes with some discussion of the utility of these methods in intelligent data analysis.

1 Linear Systems Analysis

The basic framework of traditional signal analysis[40] is schematized in figure 1; in it, an input signal is applied to a system to yield an output. One can describe this process in the time domain, using the impulse response $h(t)$ to model the system, or in the frequency domain, using the frequency response transfer function $H(s)$. The impulse response of a system is its transient response to a quick kick ($x(t_0) = 1$; $x(t) = 0 \forall t \neq t_0$); the frequency response $H(s)$ describes, for all $s$, what the system does to a sinusoidal input of frequency $s$. $H(s)$ is a complex function; it is most frequently written (and plotted) in magnitude ($|H(s)|$) and angle ($\angle H(s)$) form, but sometimes appears as $Re\{H(s)\}$ and $Im\{H(s)\}$.

Decomposing a problem in this “input + transfer function → output” manner is very useful; it allows one to apply powerful reasoning techniques like
superposition\(^1\). The problem with figure 1 is that systems can react very differently to different inputs at different times — that is, \(h(t)\) and \(H(s)\) may depend on the magnitude of \(x\), or they may have time-dependent coefficients. Either situation negates almost all of the advantages of both of the input/transfer function/output paradigms shown in the figure. Nonlinearity (the former case) and nonstationarity (the latter) are treated later in this chapter; in the remainder of this section, we assume linearity and time invariance.

The top paradigm in figure 1 is easier to think about, but the bottom is mathematically much easier to work with. In particular, deriving \(y(t)\) from \(x(t)\) and \(h(t)\) involves a convolution:

\[
y(t) = x(t) * h(t) = \int_{-\infty}^{+\infty} x(\tau)h(t - \tau)d\tau
\]

whereas the frequency-domain calculation only requires multiplication:

\[
Y(s) = X(s)H(s)
\]

The frequency domain has a variety of other powerful features. The spectrum is easy to interpret; the peaks of \(|H(s)|\) correspond to the natural frequencies ("modes") of the system and hence, loosely speaking, to the number of degrees of freedom. Differential equations become algebraic equations when transformed into the frequency domain, and signal separation is a trivial operation. Because of these advantages, engineers are trained to transform problems into the frequency domain, perform any required manipulations (e.g., filtering) in that domain, and then reverse-transform the results back into the time domain.

Traditional analysis methods characterize a linear system by describing \(h(t)\) or \(H(s)\). Depending on the demands of the application, this description — the "model" — can range from the highly abstract to the very detailed:

1. descriptive models: e.g., the sentence "as water flows out of a bathtub, the level in the tub decreases"

2. numerical models: a table of the water level in the tub versus time

\(^1\)If the inputs \(x_1\) and \(x_2\) produce the outputs \(y_1\) and \(y_2\), respectively, then the input \(x_1 + x_2\) will produce \(y_1 + y_2\).
3. graphical models: the same information, but in graphical form
4. statistical models: the mean and standard deviation of the water level
5. functional models: a least-squares fit of a line to the water level data
6. analytic models: an equation, algebraic or differential, that relates outflow and water level

The simplicity of the first item on the list is deceptive. Qualitative models like this are quite powerful — indeed, they are the basis for most human reasoning about the physical world. A circuit designer, for instance, reasons about the gain-bandwidth tradeoff of a circuit, and understands the system in terms of a balance between these two quantities ("if the gain goes up, the bandwidth (and hence the speed) goes down..."). Many traditional analysis methods are also based on qualitative models. One can, for instance, compute the location of the natural frequencies of a system from the ring frequency and decay time of its impulse response \( h(t) \) or the shape of its frequency response \( H(s) \); the latter also lets one compute the speed (rise time) and stability (gain or phase margin) of the system. Step and ramp response — how the system reacts to inputs of the form

\[
x(t) = 0 \quad t < 0 \\
x(t) = 1 \quad t \geq 0
\]

and

\[
x(t) = 0 \quad t < 0 \\
x(t) = t \quad t \geq 0
\]

respectively — also yield useful data analysis results; see [38] for details. Though qualitative models are very powerful, they are also very difficult to represent and work with explicitly; doing so effectively is the focus of the qualitative reasoning/qualitative physics community[49].

As noted and discussed by many authors (e.g., [46]), tables of numbers are much more useful to humans when they are presented in graphical form. For this reason, numerical models (item 2) are rarely used, and many IDA researchers, among others, have devoted much effort to finding and codifying systematic methods for portraying a data set and highlighting its important
features. Another way to make numbers more useful is to digest them into statistical values \[50\] like means, medians, and standard deviations, or to use the methods of functional approximation (e.g., chapter 10 of \[17\]) and regression to fit some kind of curve to the data.

Sometimes, none of these abstractions and approximations is adequate for the task at hand and one must use an analytic model. Again, these come in many flavors, ranging from algebraic expressions to partial differential equations. One of the simplest ways to use an algebraic equation to describe a system’s behavior is to model its output (or next state) as a weighted sum of its previous states. That is, if one has measured a series of values \( \{ x_i(t) \} \) of some system state variable \( x_i \), one predicts its output using the equation:

\[
y(t) = \sum_{l=0}^{L} b_l x_i(t - l) \tag{1}
\]

The technical task in fitting such an \( L^{th} \)-order moving average (MA) model to a data set involves choosing the window size \( L \) and finding appropriate values for the \( b_l \). A weighted average of the last \( L \) values is a simple smoothing operation, so this equation describes a low-pass filter. The impulse response of such a filter is described by the coefficients \( b_l \): as \( l \) goes from 0 to \( L \), the impulse first “hits” \( b_0 \), then \( b_1 \), and so on. Because this response dies out after \( L \) timesteps, equation (1) is a member of the class of finite impulse response (FIR) filters.

Autoregressive (AR) models are similar to MA models, but they are designed to account for feedback:

\[
y(t) = \sum_{m=0}^{M} a_m y(t - m) + x_i(t) \tag{2}
\]

Because of the feedback loop, the output can continue indefinitely even if the input \( x_i \) is zero, and so this type of model is called an infinite impulse response (IIR) filter. The dependence of \( y(t) \) on previous values of \( y \) also complicates the process of finding coefficients \( a_m \) that fit the model to a data set; see, e.g., \[5\] for more details.

The obvious next step is to combine MA and AR models:

\[
y(t) = \sum_{l=0}^{L} b_l x_i(t - l) + \sum_{m=0}^{M} a_m y(t - m) \tag{3}
\]

7
This “ARMA” model is both more general and more difficult to work with than its predecessors; one must choose \( L \) and \( M \) intelligently and use \( z \)-transform methods to find the coefficients (again, see [5] for this methodology). Despite these difficulties, ARMA models and their close relatives (e.g., ARIMA models) have “dominated all areas of time-series analysis and discrete-time signal processing for more than half a century”[48].

Models like those in the ARMA family capture the input/output behavior of a system in some set of test cases. For some tasks, such as controller design, an input/output model is inadequate and one really needs a model of that internal dynamics: a differential equation that accounts for the system’s dependence on present and previous states. Differential equations capture a system’s physics in a general way, but they are much more difficult to work with than the algebraic models described above. They are also much more difficult to construct; reverse-engineering the governing ODEs of a system from observations of its behavior — that is, solutions of those ODEs — is a non-trivial task (indeed, it amounts to inverting Runge-Kutta!). This procedure, which is known as system identification in the control-theory literature, is fairly straightforward if the system involved is linear; the textbook approach[27] is to choose a generic ODE system \( \ddot{x} = B\dot{x} \), fast-Fourier-transform the sensor data, and use the characteristics of the resulting impulse response to determine the coefficients of the matrix \( B \). The natural frequencies, which appear as spikes on the impulse response, yield the system’s eigenvalues; the off-diagonal elements can be determined via a residual analysis of the mode shapes between those spikes.

A linear, time-invariant system can be described quite nicely by these kinds of models, but nonstationarity or nonlinearity can throw a large wrench in the works. The standard textbook approach[8] to nonstationary data analysis involves special techniques that recognize the exact form of the nonstationarity (e.g., linear trend) and various machinations that transform the time series into stationary form, at which point one can use ARMA methods. Nonlinearity is not so easy to get around. It can be shown, for instance, that ARMA coefficients and the power spectrum (i.e., Fourier coefficients) contain the same information. Two very different nonlinear systems, however, may have almost indistinguishable spectra, so methods in the ARMA family break down in these cases\(^2\). Spectral similarity of dissimilar systems also has important implica-

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\(^2\)One can construct a patchwork of local-linear ARMA models[44] in situations like this, but such tactics contribute little to global system analysis and understanding.
tions for signal separation. In linear systems, it is often safe to assume (and easy to recognize) that the "important" parts of the signal are lower down on the frequency scale and easily separable from the noise (which is assumed to be high frequency), and it is easy to implement digital filters that remove components of a signal above a specified cutoff frequency[34]. In nonlinear systems, as described in more detail in the following section, the important parts of the signal often cover the entire spectrum, making signal separation a difficult proposition. Nonlinearity is even more of a hurdle in system identification: constructing dynamic models of linear systems is nontrivial, but human practitioners consider nonlinear system identification to be a "black art," and automating the process[6] is quite difficult.

2 Nonlinear Dynamics Basics

A dynamical system is something whose behavior evolves with time: binary stars, transistor radios, predator-prey populations, differential equations, the air stream past the cowl of a jet engine, and myriad other examples of interest to scientists and engineers in general and intelligent data analysts in particular. The bulk of an engineering or science education and the vast majority of the data analysis methods in current use, some of which are outlined in the previous section, are focused on linear systems, like a mass on a spring; systems whose governing equations do not include power terms, transcendental functions, etc. Most natural and man-made systems, however, are nonlinear. The behavior of nonlinear systems is far richer than that of linear systems, which makes them both much more difficult and much more interesting to analyze.

The state variables of a dynamical system are the fundamental quantities needed to describe it fully — angular position \( \theta \) and velocity \( \omega = \dot{\theta} \) for a pendulum, for instance, or capacitor voltages and inductor currents in an electronic circuit. The number \( n \) of state variables is known as the dimension of the system; a pendulum is a two-dimensional system, while a three-capacitor circuit has three dimensions. Simple systems like this that have a finite number of state variables can be described by ordinary differential equation (ODE) models like

\[
\ddot{x}(t) = -\frac{k}{m} x(t)
\]  
(4)
for a mass $m$ on a spring, where $x$ is the displacement from the equilibrium position and $k$ is the spring constant, or

$$\ddot{\theta}(t) = -mg \sin \theta(t)$$

(5)

for a pendulum of mass $m$ moving under the influence of gravity $g$. Equation (4) is linear and equation (5), because of the sine term, is not; in both systems, $n = 2$. If the number of state variables in the system is infinite — e.g., a moving fluid, whose physics is influenced by the pressure, temperature and velocity at every point — the system is called spatiotemporally extended, and one must use partial differential equation (PDE) models[12] to describe it properly. In this chapter, we will confine our attention to finite-dimensional dynamical systems that admit ODE models. Because the universe is by and large nonlinear, we will concentrate on methods that are general and powerful enough to handle all dynamical systems — not just linear ones. Finally, since most natural and man-made systems are not only nonlinear but also dissipative — that is, they lose some energy to processes like friction — we will not cover the methods of conservative or Hamiltonian dynamics[3, 33].

Much of traditional systems analysis, as described in the previous section, focuses on time-series or frequency-domain data. The nonlinear dynamics community, in contrast, relies primarily upon the state-space representation, plotting the behavior on the $n$-dimensional space ($\mathbb{R}^n$) whose axes are the state variables. In this representation, the damped oscillation of a mass bouncing on a spring (shown at the top of figure 2) manifests not as a pair of decaying sinusoidal time-domain signals, as shown in the middle plot on the figure, but rather as a spiral, as shown in the bottom plot. State space trajectories like this — system behavior (i.e., ODE solutions) for particular initial conditions — only implicitly contain time information; as a result, they make the geometry of the equilibrium behavior easy to recognize and analyze.

Dissipative dynamical systems have attractors: invariant state-space structures that remain after transients have died out. A useful way to think about this is to envision the "flow" of the dynamics causing the state to evolve towards a "low point" in the state-space landscape (cf., a raindrop running downhill into an ocean). There are four different kinds of attractors:

- fixed or equilibrium points (cf., the state-space plot at the bottom of Figure 1)
- periodic orbits (a.k.a. limit cycles)
Figure 2: A simple harmonic oscillator (a) undergoing a damped oscillation to a fixed point: (b) time-domain representation (c) state-space representation.
• quasiperiodic attractors

• chaotic or "strange" attractors

Fixed points — states from which the system does not move — can be stable or unstable. In the former case, perturbations will die out; in the latter, they will grow. A commonplace example of a stable fixed point is a marble at rest in the bottom of a bowl; the same marble balanced precariously on the rim of that bowl is at an unstable fixed point. Limit cycles are signals that are periodic in the time domain and closed curves in state space; an everyday example is the behavior of a healthy\(^3\) human heart. Quasiperiodic orbits and chaotic attractors are less familiar and harder to analyze, but no less common or interesting. The latter, in particular, are fascinating. They have a fixed, complicated, and highly characteristic geometry, much like an eddy in a stream, and yet nearby trajectories on a chaotic attractor move apart exponentially fast with time, much as two nearby wood chips will take very different paths through the same eddy. Trajectories cover chaotic attractors densely, visiting every point to within arbitrary \(\epsilon\), and yet they never quite repeat exactly. These properties translate to the very complex, almost-random, and yet highly structured behavior that has intrigued scientists and engineers for the last twenty years or so. Further discussion of chaotic systems, complete with a variety of examples, appears in section 4.

Attractor type is an important nonlinear data analysis feature, and there are a variety of ways for computer algorithms to recognize it automatically from state-space data. One standard geometric classification approach is cell dynamics\[^{23}\], wherein one divides the state space into uniform boxes. In Figure 3, for example, the limit cycle trajectory — a sequence of two-vectors of floating-point numbers measured by a finite-precision sensor — can be represented as the cell sequence

\[
[(1, 0), (2, 0), (3, 0), (4, 0), (4, 1), (5, 1), (5, 2), (4, 2), (3, 2), (3, 3), (4, 3), (4, 4), ...]
\]

Because multiple trajectory points are mapped into each cell, this discretized representation of the dynamics is significantly more compact than the original series of floating-point numbers and therefore much easier to work with. This is particularly important when complex systems are involved, as the number of

\[^{3}\text{One of the heart's pathological behaviors, termed ventricular fibrillation, is actually a chaotic behavior.}\]
cells in the grid grows exponentially with the number of dimensions\textsuperscript{4}. Though the approximate nature of this representation does abstract away much detailed information about the dynamics, it preserves many of its important invariant properties; see [20] or [31] for more details. This point is critical to the utility of the method; it means that conclusions drawn from the discretized trajectory are also true of the real trajectory — for example, a repeating sequence of cells in the former, as in Figure 3, implies that the $\mathbb{R}^n$ dynamics is on a limit cycle.

Much as a bowl can have several low spots or a mountain range can include many drainages, nonlinear systems can have multiple attractors of different types. Each attractor lies in a unique basin of attraction (all the points in the bowl or mountain range from which a marble or raindrop will end up at that attractor), and those basins partition\textsuperscript{5} the state space. A linear system, on the other hand, can have only one fixed point, and its basin — if it is stable — is all of $\mathbb{R}^n$. Dissipation, the notion of transient behavior that dies out, and the requirement that attractors are proper subsets of their basins are fundamentally linked. Dynamicists think about basin/attractor dynamics using the state-space contraction metaphor: initial conditions anywhere inside the boundary of a basin of attraction will converge to the associated attractor,

\textsuperscript{4}The example of Figure 3 is two-dimensional, but the cell dynamics formalism generalizes easily to arbitrary dimension.

\textsuperscript{5}This is a slight abuse of the technical term "partition;" nonattracting sets — which have no basins of attraction — can exist in dynamical systems, and basins technically do not include their boundaries.
so one envisions a volume of initial conditions spread out across the basin, all eventually converging to the attractor. (Conservative systems — those in which energy is conserved — preserve state-space volumes and do not have attractors.) Basins are very important for nonlinear data analysis. Attractors in neighboring basins can be quite different, and so small differences in initial conditions matter; a raindrop a millimeter away from a sharp mountain ridge will take a radically different path if a light breeze comes up. This can be a useful way to approach the analysis of a system that appears to have several behavior modes. Basin boundaries can be computed using the grid-based techniques described in the previous paragraph, as well as a variety of other approaches; see [18] or section 10.3.3 of [36] for more details.

The fixed nature of an attractor of a dynamical system is critically important to the approach to intelligent data analysis that is outlined in this chapter; it implies that the dynamical invariants of such attractors — their immutable mathematical properties — do not depend on how these attractors are viewed⁶, and therefore that analysis techniques that measure those invariants should yield the same results in the face of transformations like coordinate changes, for instance. Stability is such an invariant: a stable fixed point should not become unstable if one recalibrates a sensor. Topological dimension is another: a fixed point should not appear as a limit cycle when viewed from another angle. The nonlinear dynamics literature defines dozens of other dynamical invariants and proposes hundreds of algorithms for computing them; see [2] for a readable and comprehensive introduction. The two most common invariants in this list are the Lyapunov exponent λ, which measures how fast neighboring trajectories diverge, and the family of fractal dimensions, so named because they can take on non-integer values, which measure how much of \( R^n \) a trajectory actually occupies.

The Lyapunov exponent is defined as:

\[
\lambda = \lim_{t \to \infty} \frac{1}{t} \ln |m_i(t)|
\]

(6)

where the \( m_i(t) \) are the eigenvalues of the variational system (the matrix-valued linear differential equation that governs the growth of a small variation in the initial condition; see appendix B of [36] for details). A \( n \)-dimensional system has \( n \) \( \lambda \)s, each measuring the expansion rate, in one "direction," of the

⁶ within some limits, of course

14
distance between two neighboring trajectories. $\lambda$ is the nonlinear generalization of the real part of an eigenvalue; a positive $\lambda$ implies exponential growth of a perturbation along the unstable manifold, the nonlinear-systems generalization of the eigenvector associated with a positive-real-part eigenvalue. A negative $\lambda$ implies exponential shrinkage of the perturbation along the stable manifold that is the analog of the stable eigenvector. A system that has all negative $\lambda$s in some region is said to be “stable in the sense of Lyapunov,” and its trajectories relax to some proper subset of that region (the attractor). A system with all positive $\lambda$s is unstable in all directions. A zero $\lambda$ implies less-than-exponential growth, which generally takes place along the attractor. State-space contraction, part of the formal definition of dissipation, requires that $\Sigma \lambda_i < 0$ in any dissipative system.

The point of retooling the definition of dimension to allow for non-integer values is to be able to accurately characterize objects that are “between” two topological dimensions. A Cantor set, for example, contains an infinite number of zero-dimensional objects (points) but their topological dimension is still zero. Fractal dimensions capture this property; for a middle-third removed Cantor set, for example, the capacity dimension is 0.63. This invariant is used commonly by the nonlinear dynamics community because many (not all) chaotic attractors have fractal state-space structure — that is, their attractors have non-integer values of the fractal dimension. The most-common algorithm for computing any fractal dimension of a set $A$, loosely described, is to discretize state space into $\epsilon$-boxes, count the number of boxes\(^7\) occupied by $A$, and let $\epsilon \to 0$:

$$d_c = \lim_{\epsilon \to 0} \left\{ \frac{\log(N(A,\epsilon))}{\log(1/\epsilon)} \right\} \quad (7)$$

where $N(A, \epsilon)$ is the number of closed balls of radius $\epsilon > 0$ needed to cover $A$. (Strictly speaking, one doesn't just count the boxes, but rather accumulates the value of some measure on each boxes; see the discussion of equation (8) in section 3.2.) In reality, floating-point arithmetic and computational complexity place obvious limits on the $\epsilon \to 0$ part of equation (7); in practice, one repeats the dimension calculation for a range of $\epsilon$s and finds the power-law asymptote in the middle of the log-log plot of dimension versus $\epsilon$.

Dynamical invariants like $\lambda$ and $d_c$ can be used to classify attractors. In a $n$-dimensional system, there are $n$ Lyapunov exponents $\lambda_i$ and

\(^7\)Hence the term “box-counting dimension.”
• A stable fixed point has \( n \) negative \( \lambda \)s (since perturbations in any direction will die out) and a fractal dimension of zero

• An attracting limit cycle has one zero \( \lambda \) and \( n - 1 \) negative \( \lambda \)s (since perturbations off the attractor will die out, and a perturbation along the orbit remains constant) and a fractal dimension of one

• A chaotic attractor has one zero \( \lambda \) (along the attractor), at least one positive \( \lambda \) and — generally but not always — a non-integer fractal dimension. The positive \( \lambda \) reflects chaos's hallmark “sensitive dependence on initial conditions;” the system’s tendency to force neighboring trajectories apart.

Nonlinear analysis tools that target attractor type, basin geometry, dynamical invariants, etc. are harder to implement than the kinds of techniques that one can apply to a linear system, and their implications are generally less wide-ranging. If the system under consideration is linear, data analysis is relatively easy and one can make more (and more-powerful) inferences from the results. Where nonlinear systems are concerned, however, traditional methods often do not apply; in these problems, time-series analysis is much harder and the conclusions one can draw from the results are limited in range. This stems from the inherent mathematical difficulties of the domain, and is essentially unavoidable. If one is faced with a nonlinear problem, one must use the more difficult (and perhaps unfamiliar) methods covered in this chapter. The reader who is interested in delving deeper into this field should consult any of the dozens of good nonlinear dynamics books that are currently in print. An excellent overall starting point is [42], the basic mathematics is covered particularly well in [22], a comprehensive collection of algorithms appears in [36], and an entertaining popular overview may be found in [41].

3 Delay-Coordinate Embedding

Given a time series from a sensor on a single state variable \( x_t \) in a \( n \)-dimensional dynamical system, delay-coordinate embedding lets one reconstruct a useful version of the internal dynamics\(^8\) of that system. If the embedding is performed correctly, the theorems involved guarantee that the reconstructed dynamics is

\(^8\)That is, the state-space trajectory \( \{\tilde{x}(t)\} \), where \( \tilde{x} = \{x_1, x_2, \ldots, x_n\} \)
topologically (i.e., qualitatively) identical to the true dynamics of the system, and therefore that the dynamical invariants are also identical. This is an extremely powerful correspondence; it implies that conclusions drawn from the \textit{embedded} or \textit{reconstruction-space} dynamics are also true of the real — unmeasured — dynamics. This implies, for example, that one can reconstruct the dynamics of the earth’s weather simply by setting a thermometer on a windowsill.

There are, of course, some important caveats. Among other things, a correct embedding requires at least twice as many dimensions as the internal dynamics — a requirement that makes reconstruction of the weather thoroughly impractical, as it is a fluid dynamics system and thus infinite-dimensional. Moreover, even if the dynamics of the system under examination is simple, its precise dimension is often very hard to measure and rarely known \textit{a priori}. This is the main source of the hard problems of delay-coordinate embedding, which are discussed in more detail — together with some solutions — in the following sections.

\subsection{3.1 Embedding: the basic ideas}

Consider a data set comprised of samples $x_i(t)$ of a single state variable $x_i$ in a $n$-dimensional system, measured once every $\Delta t$ seconds, such as the example sensor time series shown in table 1. To embed such a data set, one constructs $m$-dimensional reconstruction-space vectors $\vec{R}(t)$ from $m$ time-delayed samples of the $x_i(t)$, such that

$$\vec{R}(t) = [x_i(t), x_i(t - \tau), x_i(t - 2\tau), \ldots, x_i(t - (m - 1)\tau)]$$

or

$$\vec{R}(t) = [x_i(t), x_i(t + \tau), x_i(t + 2\tau), \ldots, x_i(t + (m - 1)\tau)]$$

For example, if the time series in table 1 is embedded in two dimensions ($m = 2$) with a delay $\tau = 0.005$, the first few points in the reconstruction-space trajectory are:

\begin{align*}
(1.6352 & \quad 1.6260) \\
(1.6337 & \quad 1.6230) \\
(1.6322 & \quad 1.6214) \\
(1.6306 & \quad 1.6214)
\end{align*}
\[
\begin{array}{|c|c|c|c|}
\hline
x_i(t) & t & x_i(t) & t \\
\hline
1.6352 & 0.000 & 1.6214 & 0.008 \\
1.6337 & 0.001 & 1.6183 & 0.009 \\
1.6322 & 0.002 & 1.6183 & 0.010 \\
1.6306 & 0.003 & 1.6168 & 0.011 \\
1.6276 & 0.004 & 1.6137 & 0.012 \\
1.6260 & 0.005 & 1.6107 & 0.013 \\
1.6230 & 0.006 & 1.6076 & 0.014 \\
1.6214 & 0.007 & 1.6045 & 0.015 \\
\hline
\end{array}
\]

Table 1: An example data set: samples of one state variable $x_i$, measured every $\Delta t = 0.001$ seconds.

\[(1.6276\ 1.6183)\]
\[(1.6260\ 1.6183)\]
\[
\ldots
\]

If $m = 5$ and $\tau = 0.003$, the first few points of the trajectory are:

\[(1.6352\ 1.6306\ 1.6230\ 1.6183\ 1.6137)\]
\[(1.6337\ 1.6276\ 1.6214\ 1.6183\ 1.6107)\]
\[(1.6322\ 1.6260\ 1.6214\ 1.6168\ 1.6076)\]
\[(1.6306\ 1.6230\ 1.6183\ 1.6137\ 1.6045)\]
\[
\ldots
\]

The act of sampling a single system state variable $x_i(t)$ is equivalent to projecting an $n$-dimensional state-space dynamics down onto a single axis; the embedding process demonstrated above is akin to "unfolding" or "re-inflating" such a projection, albeit on different axes: the $m$ delay coordinates $x_i(t)$, $x_i(t-\tau)$, $x_i(t-2\tau)$, etc. instead of the $n$ true state variables $x_1(t)$, $x_2(t)$, \ldots, $x_n(t)$. The central theorem\cite{43} relating such embeddings to the true internal dynamics, which is generally attributed to Takens, was proved in \cite{35} and made practical in \cite{39}; informally, it states that given enough dimensions ($m$) and the right delay ($\tau$), the reconstruction-space dynamics and the true, unobserved state-space dynamics are topologically identical. More precisely, the reconstruction-space and state-space trajectories are guaranteed
to be diffeomorphic if $m = 2n+1$, where $n$ is the true dimension of the system\textsuperscript{9}.

Diffeomorphisms — transformations that are invertible, differentiable, and that possess differentiable inverses — preserve topology but not necessarily geometry; an attractor reconstructed using delay-coordinate embedding may look very different from the true attractor, but the former can be stretched and bent into the shape of the latter without “crossing over” itself. The $2n+1$ requirement of the theorem is really a brute-force worst-case limit for eliminating projection-induced crossings. The self-intersection point in figure 4(a), for example, makes the 2D projection of that curve not diffeomorphic to a circle; viewed from another angle, however, as in part (b), the curve is indeed smoothly deformable into a circle. $2n + 1$ is simply the minimum number of dimensions required to eliminate all such crossings, so lower-dimension embeddings may well be correct. This can, in fact, be exploited in deriving a tighter and easy-to-compute bound on $m$ that is valid in “almost every” situation\textsuperscript{39}.

The topological equivalence guaranteed by the Takens theorem is a powerful concept: it lets one draw sensible, justifiable conclusions about the full dynamics of an $n$-dimensional system using only the output of a single sensor. In particular, many properties of the dynamics are preserved by diffeomorphisms; if one computes them from a correct embedding, the answer will hold for the true internal dynamics as well. There are, of course, some important conditions on the theorem, and the difficulties that they pose are the source of most of the effort and subtlety in these types of methods. Specifically, in

\textsuperscript{9}$\tau$ is missing from these requirements because the theoretical conditions upon it are far less stringent and limiting, as described in the second paragraph of the next section.

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order to embed a data set, one needs \( m \) and \( \tau \), and neither of these parameters can be measured or derived from the data set, either directly or indirectly, so algorithms like those described in the following section rely on numeric and geometric heuristics to estimate them.

From a qualitative standpoint, embedding is not as outlandish as it might initially appear. The state variables in a nonlinear system are generally coupled to one another temporally by the dynamics, so using quantities that resemble forward differences as the axes of a reconstruction space makes some sense. (Techniques like divided differences can, in theory, be used to derive velocities from position data; in practice, however, these methods often fail because the associated arithmetic magnifies sensor error.) One can think of the \( x_i(t) \), \( x_i(t - \tau) \), etc., as independent coordinates that are nonlinearly related to the true state variables. The specifics of that relationship may not be obvious; the important point is that the form of that relationship ensures that the reconstructed dynamics \( \hat{\mathbf{x}}(t) \in \mathbb{R}^m \) is diffeomorphic to the true dynamics \( \mathbf{x}(t) \in \mathbb{R}^n \).

### 3.2 Finding appropriate embedding parameters

The time-series analysis literature contains scores of methods that use a variety of heuristics to solve the central problem of delay-coordinate reconstruction: given a one-dimensional time series from a dynamical system of unknown dimension, estimate values for the dimension \( m \) and delay \( \tau \) that will guarantee a correct embedding. Many of these algorithms are somewhat ad hoc; almost all are computationally expensive, highly sensitive to sensor and algorithm parameters, and some produce surprisingly different results, even on the same data set. See [2] for a recent summary and the FAQ for the newsgroup sci.nonlinear[1] for a list of public-domain software implementations of many of these algorithms. This chapter covers only a few of the most widely accepted and/or interesting representatives of each class of methods.

The delay \( \tau \) governs whether or not the coordinates \( x(t - j\tau) \) are indeed independent. If \( \tau \) is small, the reconstruction-space trajectory will lie very near the main diagonal. As long as the structure is not infinitely thin, this type of embedding is theoretically correct; in practice, however, finite-precision arithmetic on fixed-length (and possibly noisy) trajectories can easily generate apparent crossings in situations like this. If \( \tau \) is too large, on the other hand,
successive points $\tilde{R}(t)$ and $\tilde{R}(t + \Delta t)$, where $\Delta t$ is the sampling interval, will be uncorrelated and the larger spacing of the points in $\tilde{R}(t)$ again interferes numerically with topological equivalence. Ideally, then, one wants a time window for $\tau$ that is long enough for the system state to evolve to a visible (with respect to machine $\epsilon$) but not excessive extent.

One way to compute such an estimate is to perform some sort of averaged autocorrelation of successive points in the time series $x_i(t)$ or in the embedded trajectory $\tilde{R}(t)$ — e.g., autocorrelation or average mutual information[15] — as a function of $\tau$. For very small $\tau$, these statistics will be close to 1.0, since successive reconstruction-space trajectory points are very close to one another$^{10}$. For larger $\tau$, successive points become increasingly uncorrelated. The first minimum in the distribution is a sensible choice for $\tau$: qualitatively, it corresponds to the “smallest” $\tau$ for which the dynamics has caused nearby trajectory points to become somewhat uncorrelated (i.e., new information has been introduced between samples). This choice was originally proposed[15] by Fraser; other authors suggest using other features of the autocorrelation curve to choose good values for $\tau$ (e.g., the first maximum, with the rationale that these “close returns” correspond to natural periods of the system). Note that since one can compute average mutual information (AMI) from one- and two-embeddings (that is, $m = 1$ and $m = 2$), this kind of procedure does not require one to first find a correct value for $m$.

The Pineda-Sommerer (P-S) algorithm[37], which solves both halves of the embedding parameter problem at once, is more esoteric and complicated. Its input is a time series; its outputs are a delay $\tau$ and a variety of different estimates of the dimension $m$. The procedure has three major steps: it estimates $\tau$ using the mutual information function, uses that estimated value $\tau_0$ to compute a temporary estimate $E$ of the embedding dimension, and uses $E$ and $\tau_0$ to compute the generalized dimensions $D_q$, members of a parametrized family of fractal dimensions. Generalized dimensions are defined as

$$D_q = \frac{1}{q - 1} \limsup_{\epsilon \to 0} \frac{\log \sum_i p_i^q}{\log \epsilon}$$

(8)

where $p_i$ is some measure of the trajectory on box $i$. $D_0$, $D_1$, and $D_2$ are known, respectively, as the capacity, information, and correlation dimensions. The actual details of the P-S algorithm are quite involved; we will only give a qualitative description:

$^{10}$Note that $\tilde{R}(t) = x_i(t)$ if $m = 1$. 

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• Construct 1- and 2-embeddings of the data for a range of \( \tau \)s and compute the saturation dimension \( D^*_1 \) of each; the first minimum in this function is \( \tau_0 \). The \( D^*_1 \) computation entails:

• Computing the information dimension \( D_1 \) for a range of embedding dimensions \( E \) and identifying the saturation point of this curve, which occurs at embedding dimension \( D^*_1 \). The \( D_1 \) computation entails:

• Embedding the data in \( E \)-dimensional space, dividing that space into \( E \)-cubes that are \( \epsilon \) on a side, and computing \( D_1 \) using equation (8) with \( q = 1 \).

P-S incorporates an ingenious complexity-reduction technique in the fractal dimension calculation: the \( \epsilon \)s (see equation (7)) are chosen to be of the form \( 2^{-k} \) for integers \( k \) and the data are integerized; this allows most of the mathematical operations to proceed at the bit level and vastly accelerates the algorithm.

The false near neighbor (FNN) algorithm[28], which takes a \( \tau \) and a time series and produces a lower bound on \( m \), is far simpler than P-S. (As mentioned above, upper bounds for \( m \) are often chosen to be the smallest integer greater than twice the capacity dimension, \( D_0 \), of the data, in accordance with [39].) FNN is based on the observation that neighboring points may in reality be projections of points that are very far apart, as shown in Figure 5. The algorithm starts with \( m = 1 \), finds each point's nearest neighbor, and then embeds the data with \( m = 2 \). If the point separations change abruptly between the one- and two-embeddings, then the points were false neighbors (like A and C in the \( x \)-projection of Figure 5). The FNN algorithm continues adding dimensions and re-embedding until an acceptably small\(^{11}\) number of false near neighbors remains, and returns the last \( m \)-value as the estimated dimension. This algorithm is computationally quite complex; finding the nearest neighbors of \( N \) points requires \( O(N^2) \) distance calculations and comparisons. This can be reduced to \( O(N \log N) \) using a K-D tree implementation[16].

As should be obvious from the content and tone of this introduction, estimating \( \tau \) and \( m \) is algorithmically ad hoc, computationally complex, and numerically sensitive. For this reason, among others, nonlinear time-series analysis techniques that do not require embedding are extremely attractive. Recent evidence[26] suggests that the recurrence plot — a two-dimensional

\(^{11}\)An algorithm that removes all false near neighbors can be unduly sensitive to noise.
Figure 5: The geometric basis of the FNN algorithm. If this curve is projected onto the $x$ axis, the points A, B, and C appear to be near neighbors, even though C is quite distant in the 2D view. Differences between one- and two-embeddings of these data will expose false near neighbors like the [A,C] pair.

representation of a single trajectory wherein the time series spans both ordinate and abscissa and each point $(i, j)$ on the plane is shaded according to the distance between the two corresponding trajectory points $y_i$ and $y_j$ — may be such a technique. Among their other features, recurrence plots also work well on nonstationary data; see the following section for an example and more discussion.

4 Examples

In this section, we demonstrate some of the algorithms described in the previous section using two examples, one simulated and one real.

4.1 The Lorenz system

In the early 1960s[32], Edward Lorenz derived a simple model of the physics of a fluid that is being heated from below:

$$
\ddot{\bar{x}}(t) = \frac{d}{dt} \bar{\dot{x}}(t) = \begin{bmatrix} \dot{x}(t) \\ \dot{y}(t) \\ \dot{z}(t) \end{bmatrix} = \begin{bmatrix} a(y(t) - x(t)) \\ rx(t) - y(t) - x(t)z(t) \\ x(t)y(t) - bz(t) \end{bmatrix}
$$

(9)
Figure 6: State-space plots of Lorenz system behavior with $a = 10$ and $b = 8/3$: (a) a stable fixed point for $r = 15$ (b) a chaotic attractor for $r = 45$ (c) a periodic orbit for $r = 100$.

This $3^{rd}$-order ($n = 3$) ODE system is a rough approximation of a much more complex model: the Navier-Stokes PDEs for fluid flow. The state variables $x$, $y$, and $z$ are convective intensity, temperature variation, and the amount of deviation from linearity in the vertical convection profile, respectively; the coefficients $a$ and $r$ are physical parameters of the fluid — the Prandtl and Rayleigh numbers — and $b$ is the aspect ratio. This set of equations is one of the most common examples in the nonlinear dynamics literature. At low $r$ values, its solutions exhibit damped oscillations to simple fixed-point equilibria, the first category on the list of attractor types on page 10, as shown in figure 6(a). For higher $r$, which translates to a higher heat input, the convection rolls in the modeled fluid persist, in a complicated, highly structured, and nonperiodic way; see part (b) of figure 6 for an example. This behavior, reported in a 1963 paper entitled “Deterministic Nonperiodic Flow,” led Lorenz to recognize the classic “sensitive dependence on initial conditions” in the context of a fixed attractor geometry that is now a well-known hallmark of chaos. (The term “chaos” was coined twelve years later[30].) If $r$ is raised further, the convection rolls become periodic — the second category in the list on page 10. See part (c) of the figure for an example.

The trajectories plotted in figure 6 include complete information about all three of the state variables. In a real system, this may be an overly optimistic scenario; while temperature is not hard to measure, the other state variables are not so easy, so a full state-space picture of the dynamics — information that is amenable to the techniques of section 2 — may well be unavailable. Using the theory and techniques described in section 3, however, one can reconstruct the internal dynamics of this system from a time-series sampling of one of its
state variables — say, the \( x \) coordinate of the chaotic attractor in part (b) of figure 6, which is plotted in time-domain form in figure 7(a). After embedding those data in delay coordinates, one can then apply the nonlinear state-space analysis methods of section 2 to the results.

The first step in the embedding process is to decide upon a delay, \( \tau \). The first minimum in the AMI results shown in figure 7 falls at roughly \( \tau = 0.09 \) seconds\(^{12} \). Using this \( \tau \), the false-near neighbor results (part (b) of figure 7) suggest an embedding dimension of two or three, depending on one’s interpretation of the heuristic “acceptably small percentage” threshold in the algorithm. The box-counting dimension of this data set levels off at roughly 1.8 for \( m = 2 \) and above, as can be seen in part (c) of the figure. Following [39], this would imply an upper-bound embedding dimension of four.

It can be difficult to keep this menagerie of dimensions straight. In this example, the true dimension is known: \( n = 3 \). The time series \( x(t) \) in figure 7(a) is a one-dimensional projection of the \( R^3 \) trajectory in figure 6(b) onto the \( x \) axis. In the worst case, the Takens theorem tells us that an accurate reconstruction may require as many as \( m = 2n + 1 = 7 \) embedding dimensions in order to assure topological conjugacy to the true dynamics. Recall that this is a very pessimistic upper bound; in practice, slightly more opportunistic algorithms like the one proposed in [39] are able to make better bounds estimates — values for \( m \) that are lower than \( 2n + 1 \) and, at the same time, that avoid projection-induced topological inequivalencies between the true and reconstructed dynamics. In making such estimates, many of these algorithms make use of the fact that attractors do not occupy all of \( R^n \). The fractal dimension of the \( a = 10, r = 45, b = 8/3 \) Lorenz attractor, for instance, is somewhere between 1 and 2, depending upon which algorithm one uses; the calculated capacity dimension \( D_0 \) of the trajectory in figure 6(b), in particular, is 1.8, implying an upper bound of \( m = 4 \). Even this estimate is somewhat pessimistic; fractal dimension is a highly digested piece of information: a lumped parameter that compresses all the geometric information of an attractor into a single number. Because the FNN algorithm is based upon a more-detailed examination of the geometry, its results (\( m = 3 \), in this case) are a good lower bound.

Figure 8 shows embeddings of the Lorenz time series of figure 7 with \( m = 3 \) and various \( \tau \)s. Note how this reconstructed attractor starts out as a thin

\(^{12}\)The \( x \)-axis of the plot is measured in multiples of the sample interval of 0.002 second.
Figure 7: The $x$ coordinate of the chaotic Lorenz signal from part (b) of figure 5 and the corresponding embedding parameter analysis. (a) time series (b) average mutual information (AMI) as a function of the delay $\tau$ (c) false-near neighbor (FNN) percentage as a function of embedding dimension $m$ (d) box-counting dimension ($D_0$) as a function of $m$. AMI, FNN and $D_0$ results courtesy of Joe Iwanski.
Figure 8: Embeddings of the chaotic Lorenz signal from the previous three figures with $m = 3$ and various delays, plotted in 2D projection. The formal requirements of the embedding process — which these attractors meet — guarantees that they are topologically identical to the true attractor in figure 5(b).
band near the main diagonal and "inflates" with increasing $\tau$. The sample interval in this data set was not much smaller than the $\tau$ returned by the AMI algorithm, so the "thinnest" reconstruction is fairly wide. Note, too, the resemblance of these reconstructed attractors to the true state-space trajectory in figure 6(b) and how that resemblance changes with $\tau$. The whole point of doing an embedding is that the former can be deformed smoothly into the latter — even the $\tau = 0.5$ reconstruction, where the similarity (let alone the diffeomorphism!) are hard to visualize — and that the dynamical invariants of true (figure 6(b)) and reconstructed (figure 8) attractors are identical. That is, a fixed point in the reconstructed dynamics implies that there is a fixed point in the true dynamics, and so on. As noted before, this is the power of embedding: one can use nonlinear dynamics analysis techniques on its results and safely extend those conclusions to the hidden internal dynamics of the system under examination.

It would, of course, be ideal if one could avoid all of these embedding machinations and analyze the scalar time series directly. As mentioned at the end of section 3, recurrence plots (RPs) are relatively new and potentially quite powerful nonlinear time-series analysis tools whose results appear to be independent of embedding dimension in many cases[26]. An RP is a two-dimensional representation of a single trajectory; the time series is spread out along both $x$ and $y$ axes of the plot, and each pixel is colored according to the distance between the corresponding points — that is, if the 117th point on the trajectory is 14 distance units away from the 9435th point and the distance range 13–15 corresponds to the color red, the pixel lying at (117, 9435) on the RP will be shaded red. Figure 9 shows a recurrence plot (RP) of a short segment of the Lorenz signal in part (a) of figure 7. Different types of attractors leave clear and suggestive signatures in RPs; it is easy to recognize a periodic signal, for instance, and chaotic attractors exhibit the type of intricate patterns that are visible in the figure. Formalized classification of these signatures, however, is a difficult problem — and a current research topic. There are well-developed statistical approaches[26, 45], but structural/metric analysis (e.g., via pattern recognition) is still an open problem[25].

### 4.2 The driven pendulum

A time-series plot of a data set from an angle sensor on a parametrically forced pendulum — a solid aluminum arm that rotates freely on a standard bearing,
Figure 9: Recurrence plots (RPs) of a short segment of the Lorenz data from part (a) of figure 6. The pixel at $i,j$ is shaded to reflect the distance between the $i$th and $j$th point in the time series. On the unthresholded recurrence plot (UTRP) on the left, each pixel is coded according to the color bar shown to the right of the UTRP; in the thresholded RP to the right, pixels are black if the distance falls within some prescribed threshold corridor and white otherwise. Results courtesy of Joe Iwanski.
driven vertically by a motor through a simple linkage — is shown in part (a) of figure 10. An actuator controls the drive frequency and a sensor (an optical encoder) measures its angular position. The behavior of this apparently simple device is really quite complicated and interesting: for low drive frequencies, it has a single stable fixed point, but as the drive frequency is raised, the attractor undergoes a series of bifurcations. In the sensor data, this manifests as interleaved chaotic and periodic regimes[11]. The driven pendulum is also interesting from a modeling standpoint; at high resolutions, the backlash in the bearings invalidates the standard textbook model. Modeling these effects is critical, for instance, to the accurate deployment of the space shuttle’s manipulator arm[24].

The test run plotted in figure 10 was chosen for this example because the pendulum is oscillating in a chaotic manner, which rules out many traditional time-series analysis methods. The chaos manifests as seemingly structured, almost-periodic patterns in the time-series signal: oscillations that are quite similar but not identical and that almost (but not quite) repeat. Though these patterns are highly suggestive, they are very difficult to describe or classify in the time domain; in a state-space view, however, the characteristic structure of a chaotic attractor becomes patently obvious. Unfortunately, direct state-space analysis of this system is impossible. Only angle data are available; there is no angular velocity sensor and attempts to compute angular velocity via divided differences from the angle data yield numerically obscured results because the associated arithmetic magnifies the discretization error in angle (from the sensor resolution) and time (from timebase variation in the data channel).

Delay-coordinate embedding, however, produces a clean, easily analyzable picture of the dynamics that is guaranteed to be diffeomorphic to the system’s true dynamics. As in the Lorenz example, the embedding procedure begins with an estimation of $\tau$. AMI results on the chaotic pendulum data set, shown in part (b) of figure 10, suggest a delay of 0.022 seconds (roughly 11 clicks at a sample interval of 0.002 seconds). FNN results constructed using this $\tau$ (part (c) of the figure) suggest an embedding dimension of $m = 3$. The capacity dimension $D_0$ — part (d) — varies between 1.7 and 2.1, implying an upper bound of $m = 5$, following [39].

In the Lorenz example of the previous section, the true dimension $n$ was known. In the experimental pendulum setup, this is not the case. Presumably, three of the state variables are the bob angle $\theta$, the angular velocity $\omega$, and
Figure 10: A chaotic sensor data set from a parametrically forced pendulum: (a) time-domain plot of the bob angle, measured modulo $2\pi$ (b) AMI (c) FNN and (d) $D_0$ results, all courtesy of Joe Iwanski.
the time\textsuperscript{13} $t$; if, however, the device is shaking the lab bench or contracting and expanding with ambient temperature, other forces may come into play and other state variables may have important roles in the dynamics. The results described in the previous paragraph, which suggest that the dynamical behavior of the pendulum is low-dimensional ($m = 3 - 5$, specifically), imply that the system is probably not influenced by variables like lab bench position or temperature. Higher $m$ values from the estimation algorithms would suggest otherwise.

Figure 11 shows embeddings for various $\tau$s; note how a small $\tau$, as in the Lorenz example, creates a reconstruction that hugs the main diagonal, and how that reconstructed attractor unfolds as $\tau$ grows. The pendulum data were greatly oversampled, so it is possible to create a thinner embedding than in the Lorenz example, as shown in part (a) of this figure. This is the type of reconstruction whose topologically conjugacy to the true dynamics is effectively destroyed by noise and numerical problems; note the apparent overlap of trajectories and sprinkling of noisy points just outside the true attractor in the $\tau = 0.01$ and $\tau = 0.02$ embeddings.

As before, once one has a successful reconstruction of the dynamics, all of the analysis tools described in section 2 can be brought to bear upon it, and their conclusions can be assumed to hold for the system’s full underlying behavior.

5 Why Nonlinear Dynamics and Embedding are Useful for Intelligent Data Analysis

One of the more common — and more difficult — problems faced by an engineer or scientist is to analyze the dynamics of a complicated nonlinear system, given only measurements of one state variable. The techniques described in section 3 of this chapter, coupled with the theory covered in section 2, make significant inroads on this problem, allowing one to draw useful, justifiable, and sensible conclusions about a nonlinear system from the output of a single sensor. Specifically, a correct embedding of a data set from a single sensor on a black-box system is guaranteed to have the same dynamical invariants as the $n$-dimensional dynamics of the system inside the black box, and those

\textsuperscript{13}In a driven or nonautonomous system, time is an exogenous variable.
Figure 11: Embeddings of the pendulum data set from part (a) of figure 9.
invariants are useful tools for intelligent data analysis. Time-series analysis tools for linear systems are much easier to understand, implement, and use, but the universe is by and large nonlinear, so the application range of those kinds of tools is limited. Filtering out noise, for example, is fairly straightforward when one is working with data from a linear system: one simply transforms the data into the frequency domain uses a low-pass filter. In nonlinear systems, separating signal from noise is problematic, as the former is often broadband and thus the two are intermingled. (Noise, incidentally, is infinite-dimensional, so its implications for embedding dimension calculations are dire; recall the \(2n + 1\) requirement in the embedding theorems.) There has been some recent work on nonlinear "filtering" algorithms[19], including filtered delay-coordinate embedding[39] and an intriguing technique that exploits the stable and unstable manifold structure of a chaotic attractor to compress the noise ball. The latter method requires complete knowledge of the dynamics — the ODEs that govern the system. Since reverse-engineering ODEs from time-series samples of their solutions is an open problem for nonlinear systems, this filtering approach is hard to put into practice. One can, however, approximate the ODEs with local-linear models and get some reasonable results; see [13] for more details. In some cases, noise can actually be turned to advantage; its presence in a time series can allow the modeler to "explore" more of the state space[7].

One popular technique that may be conspicuous by its absence from this chapter is the neural net. Neural nets[21], which are essentially nonlinear regression networks, model the input/output behavior of a system. They are very good at learning the patterns in a data set, and hence are very effective at predicting what a system will do next. However, they do not model the underlying physics in a human-comprehensible form. It is very difficult to learn anything useful about a system by examining a neural net that has been "trained" on that system, so this technique has been omitted from this discussion. Their ability to predict, however, makes neural nets potentially useful to intelligent data analysis in a somewhat counterintuitive fashion: if one needs more data, one can train a neural net on the time series and then use it to augment that data set, generating new points that are consistent with the dynamics[9].

Nonlinear dynamics techniques like the ones described in this chapter may be more difficult to understand and use than the more-familiar linear ones, but they are more broadly applicable — indeed, the latter can be viewed as
a subset of the former. This family of theory and technique is valuable not only for time-series analysis, but also for many other tasks, such as modeling and prediction[10]. The kind of local-linear model mentioned in the previous paragraph, for instance, has been successfully used to predict the behavior of systems ranging from roulette wheels[4] to physiological disease patterns, currencies markets, and Bach fugues[48].

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References


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