Abstract Before we apply nonlinear techniques, for example those inspired by chaos theory, to dynamical phenomena occurring in nature, it is necessary to first ask if the use of such advanced techniques is justified by the data. While many processes in nature seem very unlikely a priori to be linear, the possible nonlinear nature might not be evident in specific aspects of their dynamics. The method of surrogate data has become a very popular tool to address such a question. However, while it was meant to provide a statistically rigorous, foolproof framework, some limitations and caveats have shown up in its practical use. In this paper, recent efforts to understand the caveats, avoid the pitfalls, and to overcome some of the limitations, are reviewed and augmented by new material. In particular, we will discuss specific as well as more general approaches to constrained randomisation, providing a full range of examples. New algorithms will be introduced for unevenly sampled and multivariate data and for surrogate spike trains. The main limitation, which lies in the interpretability of the test results, will be illustrated through instructive case studies. We will also discuss some implementational aspects of the realisation of these methods in the TISEAN software package.

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1 Introduction

A nonlinear approach to analysing time series data can be motivated by two distinct reasons. One is intrinsic to the signal itself while the other is due to additional knowledge we may have about the nature of the observed phenomenon. As for the first motivation, it might be that the arsenal of linear methods has been exploited thoroughly but all the efforts left certain structures in the time series unaccounted for. As for the second, a system may be known to include nonlinear components and therefore a linear description seems unsatisfactory in the first place. Such an argument is often heard for example in brain research — nobody expects for example the brain to be a linear device. In fact, there is ample evidence for nonlinearity in particular in small assemblies of neurons. Nevertheless, the latter reasoning is rather dangerous. The fact that a system contains nonlinear components does not prove that this nonlinearity is also reflected in a specific signal we measure from that system. In particular, we do not know if it is of any practical use to go beyond the linear approximation when analysing the signal. After all, we do not want our data analysis to reflect our prejudice about the underlying system but to represent a fair account of the structures that are present in the data. Consequently, the application of nonlinear time series methods has to be justified by establishing nonlinearity in the time series.

Suppose we had measured the signal shown in Fig. in some biological setting. Visual inspection immediately reveals nontrivial structure in the serial correlations. The data fails a test for Gaussianity, thus ruling out a Gaussian linear stochastic process as its source. Depending on the assumptions we are willing to make on the underlying process, we might suggest different origins for the observed strong “spikyness” of the dynamics. Superficially, low dimensional chaos seems unlikely due to the strong fluctuations, but maybe high dimensional dynamics? A large collection of neurons could intermittently synchronise to give rise to the burst episodes. In fact, certain artificial neural network models show qualitatively similar dynamics. The least interesting explanation, however, would be that all the spikyness comes from a distortion by the measurement procedure and all the serial correlations are due to linear stochastic dynamics. Occam’s razor tells us that we should be able to rule out such a simple explanation before we venture to construct more complicated models.

Surrogate data testing attempts to find the least interesting explanation that cannot be ruled out based on the data. In the above example, the data shown in Fig. this would be the hypothesis that the data has been generated by a stationary Gaussian linear stochastic process (equivalently, an autoregressive moving average or ARMA process) that is observed through an invertible, static, but possible nonlinear observation function:

\[ s_n = s(x_n), \quad \{x_n\} : \text{ARMA}(M, N). \] (1)

Neither the order \(M, N\), the ARMA coefficients, nor the function \(s(\cdot)\) are assumed to be known. Without explicitly modeling these parameters, we still know that such a process would show characteristic linear correlations (reflecting the ARMA structure) and a characteristic single time probability distribution (reflecting the action of \(s(\cdot)\) on the original Gaussian distribution). Figure shows a surrogate time series that is designed to have exactly these properties in common with the data but to be as random as possible otherwise. By a proper statistical test we can now look for additional structure that is present in the data but not in the surrogates.

In the case of the time series in Fig. there is no additional structure since it has been generated by the rule

\[ s_n = \alpha x_n^3, \quad x_n = 0.9 x_{n-1} + \eta_n \] (2)

where \(\{\eta_n\}\) are Gaussian independent increments and
α is chosen so that the data have unit variance\(^1\) This means that the strong nonlinearity that generates the bursts is due to the distorted measurement that enhances ordinary fluctuations, generated by linear stochastic dynamics.

In order to systematically exclude simple explanations for time series observations, this paper will discuss formal statistical tests for nonlinearity. We will formulate suitable null hypotheses for the underlying process or for the observed structures themselves. In the former case, null hypotheses will be extensions of the statement that the data were generated by a Gaussian linear stochastic processes. The latter situation may occur when it is difficult to properly define a class of possibly underlying processes but we want to check if a particular set of observables gives a complete account of the statistics of the data. We will attempt to reject a null hypothesis by comparing the value of a nonlinear parameter taken on by the data with its probability distribution. Since only exceptional cases allow for the exact or asymptotic derivation of this distribution unless strong additional assumptions are made, we have to estimate it by a Monte Carlo re-sampling technique. This procedure is known in the nonlinear time series literature as the method of surrogate data, see Refs. \([6–8]\). Most of the body of this paper will be concerned with the problem of generating an appropriate Monte Carlo sample for a given null hypothesis.

We will also dwell on the proper interpretation of the outcome of such a test. Formally speaking, this is totally straightforward: A rejection at a given significance level means that if the null hypothesis is true, there is certain small probability to still see the structure we detected. Non-rejection means even less: either the null hypothesis is true, or the discriminating statistics we are using fails to have power against the alternative realised in the data. However, one is often tempted to go beyond this simple reasoning and speculate either on the nature of the nonlinearity or non-stationarity that lead to the rejection, or on the reason for the failure to reject.

Since the actual quantification of nonlinearity turns out to be the easiest — or in any case the least dangerous — part of the problem, we will discuss it first. In principle, any nonlinear parameter can be employed for this purpose. They may however differ dramatically in their ability to detect different kinds of structures. Unfortunately, selecting the most suitable parameter has to be done without making use of the data since that would render the test incorrect: If the measure of nonlinearity has been optimised formally or informally with respect to the data, a fair comparison with surrogates is no longer possible. Only information that is shared by data and surrogates, that is, for example, linear correlations, may be considered for guidance. If multiple data sets are available, one could use some sequences for the selection of the nonlinearity parameter and others for the actual test. Otherwise, it is advantageous to use one of the parameter free methods that can be set up with very little detailed knowledge of the data.

Since we want to advocate to routinely use a nonlinearity test whenever nonlinear methods are planned to be applied, we feel that it is important to make a practical implementation of such a test easily accessible. Therefore, one branch of the TISEAN free software package \([9]\) is devoted to surrogate data testing. Appendix \([A]\) will discuss the implementational aspects necessary to understand what the programs in the package do.

## 2 Detecting weak nonlinearity

Many quantities have been discussed in the literature that can be used to characterise nonlinear time series. For the purpose of nonlinearity testing we need such quantities that are particular powerful in discriminating linear dynamics and weakly nonlinear signatures — strong nonlinearity is usually more easily detectable. An important objective criterion that can be used to guide the preferred choice is the discrimination power of the resulting test. It is defined as the probability that the null hypothesis is rejected when it is indeed false. It will obviously depend on how and how strongly the data actually deviates from the null hypothesis.

### 2.1 Higher order statistics

Traditional measures of nonlinearity are derived from generalisations of the two-point auto-covariance function or the power spectrum. The use of higher order cumulants as well as bi- and multi-spectra is discussed for example in Ref. \([10]\). One particularly useful third order quantity\(^2\) is

\[
\phi^{\text{rev}}(\tau) = \frac{1}{N - \tau} \sum_{n=\tau+1}^{N} (s_n - s_{n-\tau})^3, \quad (3)
\]

\(^1\) In order to simplify the notation in mathematical derivations, we will assume throughout this paper that the mean of each time series has been subtracted and it has been rescaled to unit variance. Nevertheless, we will often transform back to the original experimental units when displaying results graphically.

\(^2\) We have omitted the commonly used normalisation to second moments since throughout this paper, time series and their surrogates will have the same second order properties and identical pre-factors do not enter the tests.
since it measures the asymmetry of a series under time reversal. (Remember that the statistics of linear stochastic processes is always symmetric under time reversal. This can be most easily seen when the statistical properties are given by the power spectrum which contains no information about the direction of time.) Time reversibility as a criterion for discriminating time series is discussed in detail in Ref. \cite{1}, where, however, a different statistic is used to quantify it. The concept itself is quite folklore and has been used for example in Refs. \cite{3,2}.

Time irreversibility can be a strong signature of nonlinearity. Let us point out, however, that it does not imply a dynamical origin of the nonlinearity. We will later (Sec. \ref{sec:7.1}) give an example of time asymmetry generated by a measurement function involving a nonlinear time average.

### 2.2 Phase space observables

When a nonlinearity test is performed with the question in mind if nonlinear deterministic modeling of the signal may be useful, it seems most appropriate to use a test statistic that is related to a nonlinear deterministic approach. We have to keep in mind, however, that a positive test result only indicates nonlinearity, not necessarily determinism. Since nonlinearities tests are usually performed on data sets which do not show unambiguous signatures of low-dimensional determinism (like clear scaling over several orders of magnitude), one cannot simply estimate one of the quantitative indicators of chaos, like the fractal dimension or the Lyapunov exponent. The formal answer would almost always be that both are probably infinite. Still, some useful test statistics are at least inspired by these quantities. Usually, some effective value at a finite length scale has to be computed without establishing scaling region or attempting to approximate the proper limits.

In order to define an observable in $m$–dimensional phase space, we first have to reconstruct that space from a scalar time series, for example by the method of delays:

$$s_n = (s_{n-(m-1)\tau}, s_{n-(m-2)\tau}, \ldots, s_n) \quad \text{(4)}$$

One of the more robust choices of phase space observable is a nonlinear prediction error with respect to a locally constant predictor $F$ that can be defined by

$$\gamma(m, \tau, \epsilon) = \left( \frac{1}{N} \sum [s_{n+1} - F(s_n)]^2 \right)^{1/2} \quad \text{(5)}$$

The prediction over one time step is performed by averaging over the future values of all neighbouring delay vectors closer than $\epsilon$ in $m$ dimensions.

We have to consider the limiting case that the deterministic signature to be detected is weak. In that case, the major limiting factor for the performance of a statistical indicator is its variance since possible differences between two samples may be hidden among the statistical fluctuations. In Ref. \cite{13}, a number of popular measures of nonlinearity are compared quantitatively. The results can be summarised by stating that in the presence of time-reversal asymmetry, the particular quantity Eq.(3) that derives from the three-point autocorrelation function gives very reliable results. However, many nonlinear evolution equations produce little or no time-reversal asymmetry in the statistical properties of the signal. In these cases, simple measures like a prediction error of a locally constant phase space predictor, Eq.(\ref{equation:5}), performed best. It was found to be advantageous to choose embedding and other parameters in order to obtain a quantity that has a small spread of values for different realisations of the same process, even if at these parameters no valid embedding could be expected.

Of course, prediction errors are not the only class of nonlinearity measures that has been optimised for robustness. Notable other examples are coarse-grained redundancies \cite{10,16} and, at an even higher level of coarse-graining, symbolic methods \cite{17}. The very popular method of false nearest neighbours \cite{18} can be easily modified to yield a scalar quantity suitable for nonlinearity testing. The same is true for the concept of unstable periodic orbits (UPOs) \cite{13,20}.

### 3 Surrogate data testing

All of the measures of nonlinearity mentioned above share a common property. Their probability distribution on finite data sets is not known analytically – except maybe when strong additional assumptions about the data are made. Some authors have tried to give error bars for measures like predictabilities (e.g. Barahona and Poon \cite{21}) or averages of pointwise dimensions (e.g. Skinner et al. \cite{24}) based on the observation that these quantities are averages (mean values or medians) of many individual terms, in which case the variance (or quartile points) of the individual values yield an error estimate. This reasoning is however only valid if the individual terms are independent, which is usually not the case for time series data. In fact, it is found empirically that nonlinearity measures often do not even follow a Gaussian distribution. Also the standard error given by Roulston \cite{22} for the mutual information is fully correct only for uniformly distributed data. His derivation assumes a smooth rescaling to uniformity. In practice, however, we have to
rescale either to exact uniformity or by rank-ordering uniform variates. Both transformations are in general non-smooth and introduce a bias in the joint probabilities. In view of the serious difficulties encountered when deriving confidence limits or probability distributions of nonlinear statistics with analytical methods, it is highly preferable to use a Monte Carlo resampling technique for this purpose.

3.1 Typical vs. constrained realisations

Traditional bootstrap methods use explicit model equations that have to be extracted from the data and are then run to produce Monte Carlo samples. This typical realisations approach can be very powerful for the computation of confidence intervals, provided the model equations can be extracted successfully. The latter requirement is very delicate. Ambiguities in selecting the proper model class and order, as well as the parameter estimation problem have to be addressed. Whenever the null hypothesis involves an unknown function (rather than just a few parameters) these problems become profound. A recent example of a typical realisations approach to creating surrogates in the dynamical systems context is given by Ref. [24]. There, a Markov model is fitted to a coarse-grained dynamics obtained by binning the two dimensional delay vector distribution of a time series. Then, essentially the transfer matrix is iterated to yield surrogate sequences. We will offer some discussion of that work later in Sec. [i].

As discussed by Theiler and Prichard [24], the alternative approach of constrained realisations is more suitable for the purpose of hypothesis testing we are interested in here. It avoids the fitting of model equations by directly imposing the desired structures onto the randomised time series. However, the choice of possible null hypothesis is limited by the difficulty of imposing arbitrary structures on otherwise random sequences. In the following, we will discuss a number of null hypotheses and algorithms to provide the adequately constrained realisations. The most general method to generate constrained realisations of time series [26] is described in Sec. [i].

Consider as a toy example the null hypothesis that the data consists of independent draws from a fixed probability distribution. Surrogate time series can be simply obtained by randomly shuffling the measured data. If we find significantly different serial correlations in the data and the shuffles, we can reject the hypothesis of independence. Constrained realisations are obtained by creating permutations without replacement. The surrogates are constrained to take on exactly the same values as the data, just in random temporal order. We could also have used the data to infer the probability distribution and drawn new time series from it. These permutations with replacement would then be what we called typical realisations.

Obviously, independence is not an interesting null hypothesis for most time series problems. It becomes relevant when the residual errors of a time series model are evaluated. For example in the BDS test for non-linearity [27], an ARMA model is fitted to the data. If the data are linear, then the residuals are expected to be independent. It has been pointed out, however, that the resulting test is not particularly powerful for chaotic data [28].

3.2 The null hypothesis: model class vs. properties

From the bootstrap literature we are used to defining null hypothesis for time series in terms of a class of processes that is assumed to contain the specific process that generated the data. For most of the literature on surrogate data, this situation hasn’t changed. One very common null hypothesis goes back to Theiler and coworkers [2] and states that the data have been generated by a Gaussian linear stochastic process with constant coefficients. Constrained realisations are created by requiring that the surrogate time series have the same Fourier amplitudes as the data. We can clearly see in this example that what is needed for the constrained realisations approach is a set of observable properties that is known to fully specify the process. The process itself is not reconstructed. But this example is also exceptional. We know that the class of processes defined by the null hypothesis is fully parametrised by the set of ARMA(\(M, N\)) models (autoregressive moving average, see Eq. [1] below). If we allow for arbitrary orders \(M\) and \(N\), there is a one-to-one correspondence between the ARMA coefficients and the power spectrum. The power spectrum is here estimated by the Fourier amplitudes. The Wiener–Khinchin theorem relates it to the autocorrelation function by a simple Fourier transformation. Consequently, specifying either the class of processes or the set of constraints are two ways to achieve the same goal. The only generalisation of this favourable situation that has been found so far is the null hypothesis that the ARMA output may have been observed by a static, invertible measurement function. In that case, constraining the single time probability distribution and the Fourier amplitudes is sufficient.

If we want to go beyond this hypothesis, all we can do in general is to specify the set of constraints we will impose. We cannot usually say which class of processes this choice corresponds to. We will have to be content with statements that a given set of statistical
parameters exhaustively describes the statistical properties of a signal. Hypotheses in terms of a model class are usually more informative but specifying sets of observables gives us much more flexibility.

3.3 Test design

Before we go into detail about the generation of surrogate samples, let us outline how an actual test can be carried out. Many examples are known of nonlinearity measures that aren’t even approximately normally distributed. It has therefore been advocated since the early days of “sigmas” the data lies outside these bounds. Such a reasoning implicitly assumes a Gaussian distribution. Of the nonlinearity measure by an error bar and deriving the significance from the number of “sigmas” the data lies outside these bounds. Such a reasoning implicitly assumes a Gaussian distribution.

Instead, we follow Theiler et al. by using a rank–order test. First, we select a residual probability $\alpha$ of a false rejection, corresponding to a level of significance $(1 - \alpha) \times 100\%$. Then, for a one–sided test (e.g. looking for small prediction errors only), we generate $M = 1/\alpha - 1$ surrogate sequences. Thus, including the data itself, we have $1/\alpha$ sets. Therefore, the probability that the data by coincidence has the smallest, say, prediction error is exactly $\alpha$, as desired. For a two–sided test (e.g. for time asymmetry which can go both ways), we would generate $M = 2/\alpha - 1$ surrogates, resulting in a probability $\alpha$ that the data gives either the smallest or the largest value.

For a minimal significance requirement of 95%, we thus need at least 19 or 39 surrogate time series for one– and two–sided tests, respectively. The conditions for rank based tests with more samples can be easily worked out. Using more surrogates can increase the discrimination power.

4 Fourier based surrogates

In this section, we will discuss a hierarchy of null hypotheses and the issues that arise when creating the corresponding surrogate data. The simpler cases are discussed first in order to illustrate the reasoning. If we have found serial correlations in a time series, that is, rejected the null hypothesis of independence, we may ask of what nature these correlations are. The simplest possibility is to explain the observed structures by linear two-point autocorrelations. A corresponding null hypothesis is that the data have been generated by some linear stochastic process with Gaussian increments. The most general univariate linear process is given by

$$s_n = \sum_{i=1}^{M} a_i s_{n-i} + \sum_{i=0}^{N} b_i \eta_{n-i},$$

(6)

where $\{\eta_n\}$ are Gaussian uncorrelated random increments. The statistical test is complicated by the fact that we do not want to test against one particular linear process only (one specific choice of the $a_i$ and $b_i$), but against a whole class of processes. This is called a composite null hypothesis. The unknown values $a_i$ and $b_i$ are sometimes referred to as nuisance parameters. There are basically three directions we can take in this situation. First, we could try to make the discriminating statistic independent of the nuisance parameters. This approach has not been demonstrated to be viable for any but some very simple statistics. Second, we could determine which linear model is most likely realised in the data by a fit for the coefficients $a_i$ and $b_i$, and then test against the hypothesis that the data has been generated by this particular model. Surrogates are simply created by running the fitted model. This typical realisations approach is the common choice in the bootstrap literature, see e.g. the classical book by Efron. The main drawback is that we cannot recover the true underlying process by any fit procedure. Apart from problems associated with the choice of the correct model orders $M$ and $N$, the data is by construction a very likely realisation of the fitted process. Other realisations will fluctuate around the data which induces a bias against the rejection of the null hypothesis. This issue is discussed thoroughly in Ref. where also a calibration scheme is proposed.

The most attractive approach to testing for a composite null hypothesis seems to be to create constrained realisations. Here it is useful to think of the measurable properties of the time series rather than its underlying model equations. The null hypothesis of an underlying Gaussian linear stochastic process can also be formulated by stating that all structure to be found in a time series is exhausted by computing first and second order quantities, the mean, the variance and the auto-covariance function. This means that a randomised sample can be obtained by creating sequences with the same second order properties as the measured data, but which are otherwise random. When the linear properties are specified by the squared amplitudes of the (discrete) Fourier transform

$$|S_k|^2 = \left| \frac{1}{\sqrt{N}} \sum_{n=0}^{N-1} s_n e^{2\pi kn/N} \right|^2,$$

(7)

that is, the periodogram estimator of the power spectrum, surrogate time series $\{\pi_n\}$ are readily created by
multiplying the Fourier transform of the data by random phases and then transforming back to the time domain:

\[ \tilde{s}_n = \frac{1}{\sqrt{N}} \sum_{k=0}^{N-1} e^{i\alpha_k} |S_k| e^{-i2\pi kn/N}, \]  

where \( 0 \leq \alpha_k < 2\pi \) are independent uniform random numbers.

### 4.1 Rescaled Gaussian linear process

The two null hypotheses discussed so far (independent random numbers and Gaussian linear processes) are not what we want to test against in most realistic situations. In particular, the most obvious deviation from the Gaussian linear process is usually that the data do not follow a Gaussian single time probability distribution. This is quite obvious for data obtained by measuring intervals between events, e.g. heart beats since intervals are strictly positive. There is however a simple generalisation of the null hypothesis that explains deviations from the normal distribution by the action of an invertible, static measurement function:

\[ s_n = s(x_n), \quad x_n = \sum_{i=1}^{M} a_i x_{n-i} + \sum_{i=0}^{N} b_i \eta_{n-i}. \]

We want to regard a time series from such a process as essentially linear since the only nonlinearity is contained in the — in principle invertible — measurement function \( s(\cdot) \).

Let us mention right away that the restriction that \( s(\cdot) \) must be invertible is quite severe and often undesired. The reason why we have to impose it is that otherwise we couldn’t give a complete specification of the process in terms of observables and constraints. The problem is further illustrated in Sec. 7.1 below.

The most common method to create surrogate data sets for this null hypothesis essentially attempts to invert \( s(\cdot) \) by rescaling the time series \( \{s_n\} \) to conform with a Gaussian distribution. The rescaled version is then phase randomised (conserving Gaussianity on average) and the result is rescaled to the empirical distribution of \( \{s_n\} \). The rescaling is done by simple rank ordering. Suppose we want to rescale the sequence \( \{s_n\} \) so that the rescaled sequence \( \{r_n\} \) takes on the same values as some reference sequence \( \{g_n\} \) (e.g. draws from a Gaussian distribution). Let \( \{g_n\} \) be sorted in ascending order and rank \( \{s_n\} \) denote the ascending rank of \( s_n \), e.g. rank \( s_n \) = 3 if \( s_n \) is the 3rd smallest element of \( \{s_n\} \). Then the rescaled sequence is given by

\[ r_n = g_{\text{rank}(s_n)}, \quad n = 1, \ldots, N. \]  

Figure 3: Discrepancy of the power spectra of human breath rate data (solid line) and 19 AAFT surrogates (dashed lines). Here the power spectra have been computed with a square window of length 64.

The amplitude adjusted Fourier transform (AAFT) method has been originally proposed by Theiler et al. It results in a correct test when \( N \) is large, the correlation in the data is not too strong and \( s(\cdot) \) is close to the identity. Otherwise, there is a certain bias towards a too flat spectrum, to be discussed in the following section.

### 4.2 Flatness bias of AAFT surrogates

It is argued in Ref. that for short and strongly correlated sequences, the AAFT algorithm can yield an incorrect test since it introduces a bias towards a slightly flatter spectrum. In Fig. 3 we see power spectral estimates of a clinical data set and of 19 AAFT surrogates. The data is taken from data set B of the Santa Fe Institute time series contest. It consists of 4096 samples of the breath rate of a patient with sleep apnoea. The sampling interval is 0.5 seconds. The discrepancy of the spectra is significant. A bias towards a white spectrum is noted: power is taken away from the main peak to enhance the low and high frequencies.

Heuristically, the flatness bias can be understood as follows. Amplitude adjustment attempts to invert the unknown measurement function \( s(\cdot) \) empirically. The estimate \( \hat{s}^{-1}(\cdot) \) of the inverse obtained by the rescaling of a finite sample to values drawn from a Gaussian distribution is expected to be consistent but it is not exact for finite \( N \). The sampling fluctuations of \( \delta_{\alpha} = \hat{s}^{-1}(s_n) - s^{-1}(s_n) \) will be essentially independent of \( n \) and thus spectrally white. Consequently, Gaussian scaling amounts to adding a white component to the spectrum, which therefore tends to become flatter under the procedure. Since such a bias can lead to spurious results, surrogates have to be refined before...
a test can be performed.

4.3 Iteratively refined surrogates

In Ref. [30], we propose a method which iteratively corrects deviations in spectrum and distribution from the goal set by the measured data. In an alternating fashion, the surrogate is filtered towards the correct Fourier amplitudes and rank-ordered to the correct distribution.

Let \( \{|S_k|^2\} \) be the Fourier amplitudes, Eq. (3), of the data and \( \{c_k\} \) a copy of the data sorted by magnitude in ascending order. At each iteration stage \( (i) \), we have a sequence \( \{\tau_n^{(i)}\} \) that has the correct distribution (coincides with \( \{c_k\} \) when sorted), and a sequence \( \{r_n^{(i)}\} \) that has the correct Fourier amplitudes given by \( \{|S_k|^2\} \). One can start with \( \{r_n^{(0)}\} \) being either an AAFT surrogate, or simply a random shuffle of the data.

The step \( \tau_n^{(i)} \to r_n^{(i)} \) is a very crude “filter” in the Fourier domain: The Fourier amplitudes are simply replaced by the desired ones, first, taking the (discrete) Fourier transform of \( \{\tau_n^{(i)}\} \):

\[
R_k^{(i)} = \frac{1}{\sqrt{N}} \sum_{n=0}^{N-1} \tau_n^{(i)} e^{i2\pi kn/N} .
\]

(11)

Then transform back, replacing the actual amplitudes by the desired ones, but keeping the phases \( e^{i\psi_k^{(n)}} = R_k^{(i)} / |R_k^{(i)}| \):

\[
r_n^{(i)} = \frac{1}{\sqrt{N}} \sum_{k=0}^{N-1} e^{i\psi_k^{(n)}} |S_k| e^{-i2\pi kn/N} .
\]

(12)

The step \( r_n^{(i)} \to r_n^{(i+1)} \) proceeds by rank ordering:

\[
\tau_n^{(i+1)} = c_{\text{rank}(r_n^{(i)})} .
\]

(13)

Intuitively, it can be understood that the iteration scheme is attracted to a fixed point \( \tau_n^{(\infty)} = \tau_n^{(i+1)} \) for large \( i \). Since the minimal possible change equals to the smallest nonzero difference \( c_n - c_{n-1} \) and is therefore finite for finite \( N \), the fixed point is reached after a finite number of iterations. The remaining discrepancy between \( \tau_n^{(\infty)} \) and \( \tau_n^{(\infty)} \) can be taken as a measure of the accuracy of the method. Whether the residual bias in \( \tau_n^{(\infty)} \) or \( \tau_n^{(\infty)} \) is more tolerable depends on the data and the nonlinearity measure to be used. For coarsely digitised data\(^3\), deviations from the discrete distribution can lead to spurious results whence \( \tau_n^{(\infty)} \) is the safer choice. If linear correlations are dominant, \( \tau_n^{(\infty)} \) can be more suitable.

The final accuracy that can be reached depends on the size and structure of the data and is generally sufficient for hypothesis testing. In all the cases we have studied so far, we have observed a substantial improvement over the standard AAFT approach. Convergence properties are also discussed in [30]. In Sec. 5.3 below, we will say more about the remaining inaccuracies.

4.4 Example: Southern oscillation index

As an illustration let us perform a statistical test for nonlinearity on a monthly time series of the Southern Oscillation Index (SOI) from 1866 to 1994 (1560 samples). For a reference on analysis of Southern Oscillation data see Graham et al. [32]. Since a discussion of this climatic phenomenon is not relevant to the issue at hand, let us just consider the time series as an isolated data item. Our null hypothesis is that the data is adequately described by its single time probability distribution and its power spectrum. This corresponds to the assumption that an autoregressive moving average (ARMA) process is generating a sequence that is measured through a static monotonic, possibly nonlinear observation function.

For a test at the 99% level of significance (\( \alpha = 0.01 \)), we generate a collection of \( 1/\alpha - 1 = 99 \) surrogate time series which share the single time sample probability distribution and the periodogram estimator with the data. This is carried out using the iterative method described in Sec. 4.3 above (see also Ref. [30]). Figure 4 shows the data together with one of the 99 surrogates.

As a discriminating statistics we use a locally constant predictor in embedding space, using three dimensional delay coordinates at a delay time of one month. Neighbourhoods were selected at 0.2 times the rms amplitude of the data. The test is set up in such a way that the null hypothesis may be rejected when the prediction error is smaller for the data than for all of the 99 surrogates. But, as we can see in Fig. 4, this is not the case. Predictability is not significantly reduced by destroying possible nonlinear structure. This negative result can mean several things. The prediction error statistics may just not have any power to detect the kind of nonlinearity present. Alternatively, the underlying process may be linear and the null hypothesis true. It could also be, and this seems the most likely option after all we know about the equations governing

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\(^3\) Formally, digitisation is a non-invertible, nonlinear measurement and thus not included in the null hypothesis. Constraining the surrogates to take exactly the same (discrete) values as the data seems to be reasonably safe, though. Since for that case we haven’t seen any dubious rejections due to discretisation, we didn’t discuss this issue as a serious caveat. This decision may of course prove premature.
Figure 4: Monthly values of the Southern Oscillation Index (SOI) from 1866 to 1994 (upper trace) and a surrogate time series exhibiting the same auto-covariance function (lower trace). All linear properties of the fluctuations and oscillations are the same between both tracings. However, any possible nonlinear structure except for a static rescaling of the data is destroyed in the lower tracing by the randomisation procedure.

Climate phenomena, that the process is nonlinear but the single time series at this sampling covers such a poor fraction of the rich dynamics that it must appear linear stochastic to the analysis.

Of course, our test has been carried out disregarding any knowledge of the SOI situation. It is very likely that more informed measures of nonlinearity may be more successful in detecting structure. We would like to point out, however, that if such information is derived from the same data, or literature published on it, a bias is likely to occur. Similarly to the situation of multiple tests on the same sample, the level of significance has to be adjusted properly. Otherwise, if many people try, someone will eventually, and maybe accidentally, find a measure that indicates nonlinear structure.

4.5 Periodicity artefacts

The randomisation schemes discussed so far all base the quantification of linear correlations on the Fourier amplitudes of the data. Unfortunately, this is not exactly what we want. Remember that the autocorrelation structure given by

\[ C(\tau) = \frac{1}{N - \tau} \sum_{n=\tau+1}^{N} s_n s_{n-\tau} \]  

(14)

corresponds to the Fourier amplitudes only if the time series is one period of a sequence that repeats itself every N time steps. This is, however, not what we believe to be the case. Neither is it compatible with the null hypothesis. Conserving the Fourier amplitudes of the data means that the periodic auto-covariance function

\[ C_p(\tau) = \frac{1}{N} \sum_{n=1}^{N} s_n s_{\text{mod}(n-\tau-1,N)+1} \]  

(15)

is reproduced, rather than \( C(\tau) \). This seemingly harmless difference can lead to serious artefacts in the surrogates, and, consequently, spurious rejections in a test. In particular, any mismatch between the beginning and the end of a time series poses problems, as discussed e.g. in Ref. [7]. In spectral estimation, problems caused by edge effects are dealt with by windowing and zero padding. None of these techniques have been successfully implemented for the phase randomisation of surrogates since they destroy the invertibility of the transform.

Let us illustrate the artefact generated by an end point mismatch with an example. In order to generate an effect that is large enough to be detected visually, consider 1500 iterates of the almost unstable AR(2) process, 

\[ s_n = 1.9 s_{n-1} - 0.9001 s_{n-2} + \eta_n \]  

(upper trace of Fig. 6). The sequence is highly correlated and there is a rather big difference between the first and the last points. Upon periodic continuation, we see a jump between \( s_{1500} \) and \( s_1 \). Such a jump has spectral power at all frequencies but with delicately tuned phases. In surrogate time series conserving the
Fourier amplitudes, the phases are randomised and the spectral content of the jump is spread in time. In the surrogate sequence shown as the lower trace in Fig. 6, the additional spectral power is mainly visible as a high frequency component. It is quite clear that the difference between the data and such surrogates will be easily been picked up by, say, a nonlinear predictor, and can lead to spurious rejections of the null hypothesis.

The problem of non-matching ends can often be overcome by choosing a sub-interval of the recording such that the end points do match as closely as possible \[\gamma_{\text{jump}} = \left( \frac{s_1 - s_N}{\sum_{n=1}^{N}(s_n - \langle s \rangle)^2} \right)^2 \]

and the mismatch in the first derivative by

\[\gamma_{\text{slip}} = \left( \frac{(s_2 - s_1) - (s_N - s_{N-1})}{\sum_{n=1}^{N}(s_n - \langle s \rangle)^2} \right)^2 \]

The fractions \(\gamma_{\text{jump}}\) and \(\gamma_{\text{slip}}\) give the contributions to the total power of the series of the mismatch of the end points and the first derivatives, respectively. For the series shown in Fig. 6, \(\gamma_{\text{jump}} = 0.45\%\) and the end effect dominates the high frequency end of the spectrum. By systematically going through shorter sub-sequences of the data, we find that a segment of 1350 points starting at sample 102 yields \(\gamma_{\text{jump}} = 10^{-5}\%\) or an almost perfect match. That sequence is shown as the upper trace of Fig. 7 together with a surrogate (lower trace). The spurious “crinkliness” is removed.

In practical situations, the matching of end points is a simple and mostly sufficient precaution that should not be neglected. Let us mention that the SOI data discussed before is rather well behaved with little end-to-end mismatch \((\gamma_{\text{jump}} < 0.004\%\)) Therefore we didn’t have to worry about the periodicity artefact.

The only method that has been proposed so far that strictly implements \(C(\tau)\) rather than \(C_p(\tau)\) is given in Ref. \[26\] and will be discussed in detail in Sec. 5 below. The method is very accurate but also rather costly in terms of computer time. It should be used in cases of doubt and whenever a suitable sub-sequence cannot be found.

### 4.6 Iterative multivariate surrogates

A natural generalisation of the null hypothesis of a Gaussian linear stochastic process is that of a multivariate process of the same kind. In this case, the process is determined by giving the cross-spectrum in addition to the power spectrum of each of the channels. In Ref. \[34\], it has been pointed out that phase randomised surrogates are readily produced by multiplying the Fourier phases of each of the channels by the same set of random phases since the cross-spectrum reflects relative phases only. The authors of Ref. \[34\]
did not discuss the possibility to combine multivariate phase randomisation with an amplitude adjustment step. The extension of the iterative refinement scheme introduced in Sec. 4.3 to the multivariate case is relatively straightforward. Since deviations from a Gaussian distribution are very common and may occur due to a simple invertible rescaling due to the measurement process, we want to give the algorithm here.

Recall that the iterative scheme consists of two procedures which are applied in an alternating fashion until convergence to a fixed point is achieved. The amplitude adjustment procedure by rank ordering is readily applied to each channel individually. However, the spectral adjustment in the Fourier domain has to be modified. Let us introduce a second index in order to denote the $M$ different channels of a multivariate time series $\{s_{n,m}, \ n = 1, \ldots, N, \ m = 1, \ldots, M\}$. The change that has to be applied to the “filter” step, Eq. (12), is that the phases $\psi_{k,m}$ have to be replaced by phases $\phi_{k,m}$ with the following properties. (We have dropped the superscript $(i)$ for convenience.) The replacement should be minimal in the least squares sense, that is, it should minimise

$$h_k = \sum_{m=1}^{M} |e^{i\phi_{k,m}} - e^{i\psi_{k,m}}|^2. \quad (18)$$

Also, the new phases must implement the same phase differences exhibited by the corresponding phases $e^{i\phi_{k,m}} = S_{k,m}/|S_{k,m}|$ of the data:

$$e^{i(\phi_{k,m_2} - \phi_{k,m_1})} = e^{i(\rho_{k,m_2} - \rho_{k,m_1})}. \quad (19)$$

The last equation can be fulfilled by setting $\phi_{k,m} = \rho_{k,m} + \alpha_k$. With this, we have $h_k = \sum_{m=1}^{M} 2 - 2 \cos(\alpha_k - \psi_{k,m} + \rho_{k,m})$ which is extremal when

$$\tan \alpha_k = \frac{\sum_{m=1}^{M} \sin(\psi_{k,m} - \rho_{k,m})}{\sum_{m=1}^{M} \cos(\psi_{k,m} - \rho_{k,m})}. \quad (20)$$

The minimum is selected by taking $\alpha_k$ in the correct quadrant.

As an example, let us generate a surrogate sequence for a simultaneous recording of the breath rate and the instantaneous heart rate of a human during sleep. The data is again taken from data set B of the Santa Fe Institute time series contest [32]. The 1944 data
points are an end-point matched sub-sequence of the data used as a multivariate example in Ref. [26]. In the latter study, which will be commented on in Sec. 6.2 below, the breath rate signal had been considered to be an input and therefore not been randomised. Here, we will randomise both channels under the condition that their individual spectra as well as their cross-correlation function are preserved as well as possible while matching the individual distributions exactly. The iterative scheme introduced above took 188 iterations to converge to a fixed point. The data and a bi-variate surrogate is shown in Fig. 8. In Fig. 8 the cross-correlation functions of the data and one surrogate are plotted. Also, for comparison, the same for two individual surrogates of the two channels. The most striking difference between data and surrogates is that the coherence of the breath rate is lost. Thus, it is indeed reasonable to exclude the nonlinear structure in the heart rate, by taking the breath rate as a given input signal. Such an analysis is however beyond the scope of the method discussed in this section. First of all, specifying the full cross-correlation function to a fixed signal plus the autocorrelation function over-specifies the problem and there is no room for randomisation. In Sec. 6.2 below, we will therefore revisit this problem. With the general constrained randomisation scheme to be introduced below, it will be possible to specify a limited number of lags of the auto- and cross-correlation functions.

5 General constrained randomisation

Randomisation schemes based on the Fourier amplitudes of the data are appropriate in many cases. However, there remain some flaws, the strongest being the severely restricted class of testable null hypotheses. The periodogram estimator of the power spectrum is about the only interesting observable that allows for the solution of the inverse problem of generating random sequences under the condition of its given value.

In the general approach of Ref. [26], constraints (e.g. autocorrelations) on the surrogate data are implemented by a cost function which has a global minimum when the constraints are fulfilled. This general framework is much more flexible than the Fourier based methods. We will therefor discuss it in some detail.

5.1 Null hypotheses, constraints, and cost functions

As we have discussed previously, we will often have to specify a null hypothesis in terms of a complete set of observable properties of the data. Only in specific cases (e.g. the two point autocorrelation function), there is a one-to-one correspondence to a class of models (here the ARMA process). In any case, if \( \{ s_n \} \) denotes a surrogate time series, the constraints will most often be of (or can be brought into) the form

\[
F_i(\{ s_n \}) = 0, \quad i = 1, \ldots, I.
\]  

Such constraints can always be turned into a cost function

\[
E(\{ s_n \}) = \left( \sum_{i=1}^{I} |w_i F_i(\{ s_n \})|^q \right)^{1/q}.
\]  

The fact that \( E(\{ s_n \}) \) has a global minimum when the constraints are fulfilled is unaffected by the choice of the weights \( w_i \neq 0 \) and the order \( q \) of the average. The least squares or \( L^2 \) average is obtained at \( q = 2 \), \( L^1 \) at \( q = 1 \) and the maximum distance when \( q \to \infty \). Geometric averaging is also possible (and can be formally obtained by taking the limit \( q \to 0 \) in a proper way). We have experimented with different choices of \( q \) but we haven’t found a choice that is uniformly superior to others. It seems plausible to give either uniform weights or to enhance those constraints which are particularly difficult to fulfill. Again, conclusive empirical results are still lacking.

Consider as an example the constraint that the sample autocorrelation function of the surrogate \( \bar{C}(\tau) = \langle \bar{s}_n \bar{s}_{n-\tau} \rangle \) (data rescaled to zero mean and unit variance) are the same as those of the data, \( C(\tau) = \langle s_n s_{n-\tau} \rangle \). This is done by specifying zero discrepancy as a constraint \( F_{\tau}(\{ s_n \}) = \bar{C}(\tau) - C(\tau), \quad \tau = 1, \ldots, \tau_{\text{max}} \). If the correlations decay fast, \( \tau_{\text{max}} \) can be restricted, otherwise \( \tau_{\text{max}} = N - 1 \) (the largest available lag). Thus, a possible cost function could read

\[
E = \max_{\tau=0}^{\tau_{\text{max}}} \left| \bar{C}(\tau) - C(\tau) \right|.
\]  

Other choices of \( q \) and the weights are of course also possible.

In all the cases considered in this paper, one constraint will be that the surrogates take on the same values as the data but in different time order. This ensures that data and surrogates can equally likely be drawn from the same (unknown) single time probability distribution. This particular constraint is not included in the cost function but identically fulfilled by considering only permutations without replacement of the data for minimisation.

By introducing a cost function, we have turned a difficult nonlinear, high dimensional root finding problem into a minimisation problem. This leads to extremely many false minima whence such a strategy is discouraged for general root finding problems [42].
Here, the situation is somewhat different since we need to solve Eq. (23) only over the set of all permutations of $\{s_n\}$. Although this set is big, it is still discrete and powerful combinatorial minimisation algorithms are available that can deal with false minima very well. We choose to minimise $E(\{s_n\})$ among all permutations $\{s_n\}$ of the original time series $\{s_n\}$ using the method of simulated annealing. Configurations are updated by exchanging pairs in $\{s_n\}$. The annealing scheme will decide which changes to accept and which to reject. With an appropriate cooling scheme, the annealing procedure can reach any desired accuracy. Apart from simulated annealing, genetic algorithms have become very popular for this kind of problems and there is no reason why they couldn’t be used for the present purpose as well.

5.2 Computational issues of simulated annealing

Simulated annealing is a very powerful method of combinatorial minimisation in the presence of many false minima. Simulated annealing has a rich literature, classical references are Metropolis et al. \cite{36} and Kirkpatrick \cite{37}, more recent material can be found for example in Vidal \cite{38}. Despite its many successful applications, using simulated annealing efficiently is still a bit of an art. We will here discuss some issues we have found worth dealing with in our particular minimisation problem. Since the detailed behaviour will be different for each cost function, we can only give some general guidelines.

The main idea behind simulated annealing is to interpret the cost function $E$ as an energy in a thermodynamic system. Minimising the cost function is then equivalent to finding the ground state of a system. A glassy solid can be brought close to the energetically optimal state by first heating it and subsequently cooling it. This procedure is called “annealing”, hence the name of the method. If we want to simulate the thermodynamics of this tempering procedure on a computer, we notice that in thermodynamic equilibrium at some finite temperature $T$, system configurations should be visited with a probability according to the Boltzmann distribution $e^{-E/T}$ of the canonical ensemble. In Monte Carlo simulations, this is achieved by accepting changes of the configuration with a probability $p = 1$ if the energy is decreased ($\Delta E < 0$) and $p = e^{-\Delta E/T}$ if the energy is increased, ($\Delta E \geq 0$). This selection rule is often referred to as the Metropolis step. In a minimisation problem, the temperature is the parameter in the Boltzmann distribution that sets its width. In particular, it determines the probability to go “up hill”, which is important if we need to get out of false minima.

Figure 10: Building up correlations by pairwise permutation. Suppose we want to generate the strong anti-correlation present in the data (upper trace) by minimising $E = |C(1) - C(1)|$. The annealing started with a random permutation (middle trace, $E = 1.129$). At a given intermediate state (lower trace, $E = 0.256$), exchanging the points $a$ and $b$ increases the cost to $E = 0.2744$ while exchanging $c$ and $d$ creates negative correlation and reduces the cost to $E = 0.002$.

In order to anneal the system to the ground state of minimal “energy”, that is, the minimum of the cost function, we want to first “melt” the system at a high temperature $T$, and then decrease $T$ slowly, allowing the system to be close to thermodynamic equilibrium at each stage. If the changes to the configuration we allow to be made connect all possible states of the system, the updating algorithm is called ergodic. Although some general rigorous convergence results are available, in practical applications of simulated annealing some problem-specific choices have to be made. In particular, apart from the constraints and the cost function, one has to specify a method of updating the configurations and a schedule for lowering the temperature. In the following, we will discuss each of these issues.

Concerning the choice of cost function, we have already mentioned that there is a large degeneracy in that many cost functions have an absolute minimum whenever a given set of constraints if fulfilled. The convergence properties can depend dramatically on the choice of cost function. Unfortunately, this dependence seems to be so complicated that it is impossible even to discuss the main behaviour in some gen-
erality. In particular, the weights \( w_i \) in Eq. (22) are sometimes difficult to choose. Heuristically, we would like to reflect changes in the \( I \) different constraints about equally, provided the constraints are independent. Since their scale is not at all set by Eq. (21), we can use the \( w_i \) for this purpose. Whenever we have some information about which kind of deviation would be particularly problematic with a given test statistic, we can give it a stronger weight. Often, the shortest lags of the autocorrelation function are of crucial importance, whence we tend to weight autocorrelations by 1/\( \tau \) when they occur in sums. Also, the \( C(\tau) \) with larger \( \tau \) are increasingly ill-determined due to the fewer data points entering the sums. As an extreme example, \( C(N - 1) = s_{14}N - 1 \) shows huge fluctuations due to the lack of self-averaging. Finally, there are many more \( C(\tau) \) with larger \( \tau \) than at the crucial short lags.

A way to efficiently reach all permutations by small individual changes is by exchanging randomly chosen (not necessarily close-by) pairs. How the interchange of two points can affect the current cost is illustrated schematically in Fig. 11. Optimising the code that computes and updates the cost function is essential since we need its current value at each annealing step — which are expected to be many. Very often, an exchange of two points is reflected in a rather simple update of the cost function. For example, computing \( C(\tau) \) for a single lag \( \tau \) involves \( O(N) \) multiplications. Updating \( C(\tau) \) upon the exchange of two points \( i < j \) only requires the replacement of the terms \( s_i s_{i - \tau}, s_{i + \tau} s_i, s_j s_{j - \tau}, \) and \( s_{j + \tau} s_j \) in the sum. Note that cheap updates are a source of potential mistakes (e.g. avoid subtracting terms twice in the case that \( i = j - \tau \) but also of roundoff errors. To ensure that the assumed cost is always equal to the actual cost, code carefully and monitor roundoff by computing a fresh cost function occasionally.

Further speed-up can be achieved in two ways. Often, not all the terms in a cost function have to be added up until it is clear that the resulting change goes up hill by an amount that will lead to a rejection of the exchange. Also, pairs to be exchanged can be selected closer in magnitude at low temperatures because large changes are very likely to increase the cost.

Many cooling schemes have been discussed in the literature [18]. We use an exponential scheme in our work. We will give details on the — admittedly largely ad hoc — choices that have been made in the TISEAN implementation in Appendix A. We found it convenient to have a scheme available that automatically adjusts parameters until a given accuracy is reached. This can be done by cooling at a certain rate until we are stuck (no more accepted changes). If the cost is not low enough yet, we melt the system again and cool at a slower rate.

5.3 Example: avoiding periodicity artefacts

Let us illustrate the use of the annealing method in the case of the standard null hypothesis of a rescaled linear process. We will show how the periodicity artefact discussed in Sec. 4.3 can be avoided by using a more suitable cost function. We prepare a surrogate for the data shown in Fig. 8 (almost unstable AR(2) process) without truncating its length. We minimise the cost function given by Eq. (23), involving all lags up to \( \tau_{\text{max}} = 100 \). Also, we excluded the first and last points from permutations as a cheap way of imposing the long range correlation. In Fig. 11 we show progressive stages of the annealing procedure, starting from a
random scramble. The temperature $T$ is decreased by 0.1% after either $10^6$ permutations have been tried or $10^4$ have been successful. The final surrogate neither has spuriously matching ends nor the additional high frequency components we saw in Fig. 6. The price we had to pay was that the generation of one single surrogate took 6 h of CPU time on a Pentium II PC at 350 MHz. If we had taken care of the long range correlation by leaving the end points loose but taking $\tau_{\text{max}} = N - 1$, convergence would have been prohibitively slow. Note that for a proper test, we would need at least 19 surrogates. We should stress that this example with its very slow decay of correlations is not the case, for example, if we include all possible lags of the autocorrelation function, which gives as many minimisation problems, we are not really interested in the solutions that put sacrificing 10% of the points to get rid of the end point mismatch is preferable here to spending several days of CPU time on the annealing scheme. In other cases, however, we may not have such a choice.

5.4 Combinatorial minimisation and accuracy

In principle, simulated annealing is able to reach arbitrary accuracy at the expense of computer time. We should, however, remark on a few points. Unlike other minimisation problems, we are not really interested in the solutions that put the probability distribution over the $N!$ permutations, unless $N$ is really small or the constraints grossly over-specify the problem. This can be the case, for example, if we include all possible lags of the autocorrelation function, which gives as many (nonlinear) equations as unknowns, $I = N$. These may close for small $N$ in the space of permutations. In such extreme situations, it is possible to include extra cost terms penalising closeness to one of the trivial transformations of the data. Let us note that if the surrogates are “too similar” to the data, this does not in itself affect the validity of the test. Only the discrimination power may be severely reduced.

Now, if we don’t want to reach $E = 0$, how can we be sure that there are enough independent realisations with $E \approx 0$? The theoretical answer depends on the form of the constraints in a complicated way and cannot be given in general. We can, however, offer a heuristic argument that the number of configurations with $E$ smaller than some $\Delta E$ grows fast for large $N$. Suppose that for large $N$ the probability distribution of $E$ converges to an asymptotic form $p(E)$. Assume further that $\tilde{p}(\Delta E) = \text{Prob}(E < \Delta E) = \int_0^{\Delta E} p(E) dE$ is nonzero but maybe very small. This is evidently true for autocorrelations, for example. While thus the probability to find $E < \Delta E$ in a random draw from the distribution of the data may be extremely small, say $\tilde{p}(\Delta E) = 10^{-45}$ at 10 sigmas from the mean energy, the total number of permutations, figuring as the number of draws, grows as $N! \approx (N/e)^N \sqrt{2\pi N}$, that is, much faster than exponentially. Thus, we expect the number of permutations with $E < \Delta E$ to be $\propto \tilde{p}(\Delta E)N!$. For example, $10^{-45} \times 1000! \approx 10^{2522}$.

In any case, we can always monitor the convergence of the cost function to avoid spurious results due to residual inaccuracy in the surrogates. As we will discuss below, it can also be a good idea to test the surrogates with a linear test statistic before performing the actual nonlinearity test.

5.5 The curse of accuracy

Strictly speaking, the concept of constrained realisations requires the constraints to be fulfilled exactly, a practical impossibility. Most of the research efforts reported in this article have their origin in the attempt to increase the accuracy with which the constraints are implemented, that is, to minimise the bias resulting from any remaining discrepancy. Since most measures of nonlinearity are also sensitive to linear correlations, a side effect of the reduced bias is a reduced variance of such estimators. Paradoxically, thus the enhanced accuracy may result in false rejections of the null hypothesis on the ground of tiny differences in some nonlinear characteristics. This important point has been recently put forth by Kugiumutzis [20].

Consider the highly correlated autoregressive process $x_n = 0.99x_{n-1} - 0.8x_{n-2} + 0.65x_{n-3} + \eta_n$, measured by the function $s_n = s(x_n) = x_n|x_n$ and then normalised to zero mean and unit variance. The strong correlation together with the rather strong static nonlinearity makes this a very difficult data set for the generation of surrogates. Figure 12 shows the bias and variance for a linear statistic, the unit lag autocorrelation $C_p(1)$, Eq. (15), as compared to its goal value given by the data. The left part of Fig. 13 shows $C_p(1)$ versus the iteration count $i$ for 200 iterative surrogates, $i = 1$ roughly corresponding to AAFT surrogates. Although the mean accuracy increases dramatically compared to the first iteration stages, the data consistently remains outside a $2\sigma$ error bound. Since nonlinear parameters will also pick up linear correlations, we have to expect spurious results in such a case. In the right part, annealed surrogates are generated with a cost function $E = \max_{\tau=1}^{200} |C_p(\tau) - C_p(\tau)|/\tau$. The bias and variance of $C_p(1)$ are plotted versus the cost $E$. Since the cost function involves $C_p(1)$, it is not surprising that we see good convergence of the bias. It is also noteworthy that the variance is in any event large enough to
Figure 12: Bias and variance of unit lag autocorrelation $C_p(1)$ for ensembles of surrogates. Left part: $C_p(1)$ plotted versus the iteration count $i$ for 200 iterative surrogates. The AAFT method gives accuracies comparable to the value obtained for $i = 1$. Right part: $C_p(1)$ plotted versus the goal value of the cost function for 20 annealed surrogates. The horizontal line indicates the sample value for the data sequence. See text for discussion.

exclude spurious results due to remaining discrepancy in the linear correlations.

Kugiumtzis suggests to test the validity of the surrogate sample by performing a test using a linear statistic for normalisation. For the data shown in Fig. 12, this would have detected the lack of convergence of the iterative surrogates. Currently, this seems to be the only way around the problem and we thus recommend to follow his suggestion. With the much more accurate annealed surrogates, we haven’t so far seen examples of dangerous remaining inaccuracy, but we cannot exclude their possibility. If such a case occurs, it may be possible to generate unbiased ensembles of surrogates by specifying a cost function that explicitly minimises the bias. This would involve the whole collection of $M$ surrogates at the same time, including extra terms like

$$E_{\text{ensemble}} = \sum_{\tau=0}^{\tau_{\text{max}}} \left( \sum_{m=1}^{M} \overline{C}_m(\tau) - C(\tau) \right)^2.$$ (24)

Here, $\overline{C}_m(\tau)$ denotes the autocorrelation function of the $m$-th surrogate. In any event, this will be a very cumbersome procedure, in terms of implementation and in terms of execution speed and it is questionable if it is worth the effort.

6 Various Examples

In this section, we want to give a number of applications of the constrained randomisation approach. If the constraints consist only of the Fourier amplitudes and the single time probability distribution, the iteratively refined, amplitude adjusted surrogates discussed in Sec. 4.3 are usually sufficient if the end point artefact can be controlled and convergence is satisfactory. Even the slightest extension of these constraints makes it impossible to solve the inverse problem directly and we have to follow the more general combinatorial approach discussed in the previous section. The following examples are meant to illustrate how this can be carried out in practice.

6.1 Including non-stationarity

Constrained randomisation using combinatorial minimisation is a very flexible method since in principle arbitrary constraints can be realised. Although it is seldom possible to specify a formal null hypothesis for more general constraints, it can be quite useful to be able to incorporate into the surrogates any feature of the data that is understood already or that is uninteresting. Non-stationarity has been excluded so far by requiring the equations defining the null hypothesis to remain constant in time. This has a two-fold consequence. First, and most importantly, we must keep in mind that the test will have discrimination power against non-stationary signals as a valid alternative to the null hypothesis. Thus a rejection can be due to nonlinearity or non-stationarity equally well.

Second, if we do want to include non-stationarity in
the null hypothesis we have to do so explicitly. Let us illustrate how this can be done with an example from finance. The time series consists of 1500 daily returns (until the end of 1996) of the BUND Future, a derived German financial instrument. The data were kindly provided by Thomas Schürmann, WGZ-Bank Düsseldorf. As can be seen in the upper panel of Fig. 13, the sequence is non-stationary in the sense that the local variance and to a lesser extent also the local mean undergo changes on a time scale that is long compared to the fluctuations of the series itself. This property is known in the statistical literature as heteroscedasticity and modelled by the so-called GARCH [14] and related models. Here, we want to avoid the construction of an explicit model from the data but rather ask the question if the data is compatible with the null hypothesis of a correlated linear stochastic process with time dependent local mean and variance. We can answer this question in a statistical sense by creating surrogate time series that show the same linear correlations and the same time dependence of the running mean and running variance as the data and comparing a nonlinear statistic between data and surrogates. The lower panel in Fig. 13 shows a surrogate time series generated using the annealing method. The cost function was set up to match the autocorrelation function up to five days and the moving mean and variance in sliding windows of 100 days duration. In Fig. 13, the running mean and variance are shown as points and error bars, respectively, in the middle trace. The deviation of these between data and surrogate has been minimised to such a degree that it can no longer be resolved. A comparison of the time-asymmetry statistic Eq. (3) for the data and 19 surrogates did not reveal any discrepancy, and the null hypothesis could not be rejected.

6.2 Multivariate data

In Ref. [26], the flexibility of the approach was illustrated by a simultaneous recording of the breath rate and the instantaneous heart rate of a human subject during sleep. The interesting question was, how much of the structure in the heart rate data can be explained by linear dependence on the breath rate. In order to answer this question, surrogates were made that had the same autocorrelation structure but also the same cross-correlation with respect to the fixed input signal, the breath rate. While the linear cross-correlation with the breath rate explained the coherent structure of the heart rate, other features, in particular its asymmetry under time reversal, remained unexplained. Possible explanations include artefacts due to the peculiar way of deriving heart rate from inter-beat intervals, non-linear coupling to the breath activity, nonlinearity in the cardiac system, and others.

Within the general framework, multivariate data can be treated very much the same way as scalar time series. In the above example, we chose to use one of the channels as a reference signal which was not randomised. The rationale behind this was that we were not looking for nonlinear structure in the breath rate itself and thus we didn’t want to destroy any such structure in the surrogates. In other cases, we can decide either to keep or to destroy cross-correlations between channels. The former can be achieved by applying the same permutations to all channels. Due to the limited experience we have so far and the multitude of possible cases, multivariate problems have not been included in the TISEAN implementation yet.

6.3 Uneven sampling

Let us show how the constrained randomisation method can be used to test for nonlinearity in time series taken at time intervals of different length. Unevenly sampled data are quite common, examples include drill core data, astronomical observations or stock price notations. Most observables and algorithms cannot easily be generalised to this case which is particularly true for nonlinear time series methods. (See [41] for material on irregularly sampled time series.) Interpolating the data to equally spaced sampling times is not recommendable for a test for nonlinearity since one could not a posteriori distinguish between genuine structure and nonlinearity introduced spuriously by the interpolation process. Note that also zero padding is a nonlinear operation in the sense that stretches of zeroes are unlikely to be produced by any linear stochastic process.

For data that is evenly sampled except for a moderate number of gaps, surrogate sequences can be produced relatively straightforwardly by assuming the value zero during the gaps and minimising a standard cost function like Eq. (23) while excluding the gaps from the permutations tried. The error made in estimating correlations would then be identical for the data and surrogates and could not affect the validity of the test. Of course, one would have to modify the nonlinearity measure to avoid the gaps. For data sampled at incommensurate times, such a strategy can no longer be adopted. We then need different means to specify the linear correlation structure.

Two different approaches are viable, one residing in the spectral domain and one in the time domain. Consider a time series sampled at times \{t_n\} that need not be equally spaced. The power spectrum can then be estimated by the Lomb periodogram, as discussed for
example in Ref. [2]. For time series sampled at constant time intervals, the Lomb periodogram yields the standard squared Fourier transformation. Except for this particular case, it does not have any inverse transformation, which makes it impossible to use the standard surrogate data algorithms mentioned in Sec. 13. In Ref. 14, we used the Lomb periodogram of the data as a constraint for the creation of surrogates. Unfortunately, imposing a given Lomb periodogram is very time consuming because at each annealing step, the $O(N)$ spectral estimator has to be computed at $O(N_f)$ frequencies with $N_f \propto N$. Press et al. [12] give an approximation algorithm that uses the fast Fourier transform to compute the Lomb periodogram in $O(N \log N)$ time rather than $O(N^2)$. The resulting code is still quite slow.

As a more efficient alternative to the commonly used but computationally costly Lomb periodogram, let us suggest to use binned autocorrelations. They are defined as follows. For a continuous signal $s(t)$ (take $\langle s \rangle = 0$, $\langle s^2 \rangle = 1$ for simplicity of notation here), the autocorrelation function is $C(\tau) = \langle s(t)s(t-\tau) \rangle = (1/T) \int_0^T dt' s(t')s(t'-\tau)$. It can be binned to a bin size $\Delta$, giving $C_{\Delta}(\tau) = (1/\Delta) \int_{\tau-\Delta}^{\tau+\Delta} dt' C(\tau')$. We now have to approximate all integrals using the available values of $s(t_n)$. In general, we estimate

$$\int_a^b dt \ f(t) \approx (b-a) \sum \frac{B_n(a,b) \ f(t_n)}{|B_n(a,b)|} . \quad (25)$$

Here, $B_n(a,b) = \{ n : a < t_n \leq b \}$ denotes the bin ranging from $a$ to $b$ and $|B(a,b)|$ the number of its elements. We could improve this estimate by some interpolation of $f(\cdot)$, as it is customary with numerical integration but the accuracy of the estimate is not the central issue here. For the binned autocorrelation, this approximation simply gives

$$C_{\Delta}(\tau) \approx \frac{\sum B_n(\tau-\Delta, \tau) \ s(t_i)s(t_j)}{|B_n(\tau-\Delta, \tau)|} . \quad (26)$$

Here, $B_n(i,j) = \{ (i,j) : a < t_i - t_j \leq b \}$. Of course, empty bins lead to undefined autocorrelations. If we have evenly sampled data and unit bins, $t_i - t_{i-1} = \Delta$, $i = 2, \ldots, N$, then the binned autocorrelations coincide with ordinary autocorrelations at $\tau = i\Delta$, $i = 0, \ldots, N-1$.

Once we are able to specify the linear properties of a time series, we can also define a cost function as usual and generate surrogates that realise the binned autocorrelations of the data. A delicate point however is the choice of bin size. If we take it too small, we get bins that are almost empty. Within the space of permutations, there may be only a few ways then to generate precisely that value of $C_{\Delta}(\tau)$, in other words, we over-specify the problem. If we take the bin size too large, we might not capture important structure in the autocorrelation function.

As an application, let us construct randomised versions of part of an ice core data set, taken from the Greenland Ice Sheet Project Two (GISP2) [14]. An extensive data base resulting from the analysis of physical and chemical properties of Greenland ice up to a depth of 3028.8 m has been published by the National Snow and Ice Data Center together with the World Data Center-A for Paleoclimatology, National Geophysical Data Center, Boulder, Colorado [15]. A long ice core is usually cut into equidistant slices and initially, all measurements are made versus depth. Considerable expertise then goes into the dating of each slice [16]. Since the density of the ice, as well as the annual total deposition, changes with time, the final time series data are necessarily unevenly sampled. Furthermore, often a few values are missing from the record. We will study a subset of the data ranging back 10000 years in time, corresponding to a depth of 1564 m, and continuing until 2000 years before present. Figure 14 shows the sampling rate versus time for the particular ice core considered.

We use the $\delta^{18}O$ time series which indicates the deviation of the $\alpha = 18O/16O$ ratio from its standard value $\alpha_0$: $\delta^{18}O = 0.103(\alpha - \alpha_0)/\alpha_0$. Since the ratio of the condensation rates of the two isotopes depends on temperature, the isotope ratio can be used to derive a temperature time series. The upper trace in Fig. 14 shows the recording from 10000 years to 2000 years before present, comprising 538 data points.

In order to generate surrogates with the same linear properties, we estimate autocorrelations up to a lag of $\tau = 1000$ years by binning to a resolution of 5 y. A typical surrogate is shown as the lower trace in Fig. 14. We have not been able to detect any nonlinear struc-
ture by comparing this recording with 19 surrogates, neither using time asymmetry nor prediction errors. It should be admitted, however, that we haven’t attempted to provide nonlinearity measures optimised for the unevenly sampled case. For that purpose, also some interpolation is permissible since it is then part of the nonlinear statistic. Of course, in terms of geophysics, we are asking a very simplistic question here. We wouldn’t really expect strong nonlinear signatures or even chaotic dynamics in such a single probe of the global climate. All the interesting information — and expected nonlinearity — lies in the interrelation between various measurements and the assessment of long term trends we have deliberately excluded by selecting a subset of the data.

### 6.4 Spike trains

A spike train is a sequence of \(N\) events (for example neuronal spikes, or heart beats) occurring at times \(\{t_n\}\). Variations in the events beyond their timing are ignored. Let us first note that this very common kind of data is fundamentally different from the case of unevenly sampled time series studied in the last section in that the sampling instances \(\{t_n\}\) are not independent of the measured process. In fact, between these instances, the value of \(s(t)\) is undefined and the \(\{t_n\}\) contain all the information there is.

Very often, the discrete sequence if inter-event intervals \(x_n = t_n - t_{n-1}\) is treated as if it were an ordinary time series. We must keep in mind, however, that the index \(n\) is not proportional to time any more.

\[
s(t) = \sum_{n=1}^{N} \delta(t - t_n) .
\]

(27)

With \(\int dt \ s(t) e^{i \omega t} = \sum_{n=1}^{N} e^{-i \omega t_n}\), the periodogram spectral estimator is then simply obtained by squaring the (continuous) Fourier transform of \(s(t)\):

\[
P(\omega) = \left| \frac{1}{2\pi} \sum_{n=1}^{N} e^{-i \omega t_n} \right|^2 .
\]

(28)

Other spectral estimators can be derived by smoothing \(P(\omega)\) or by data windowing. It is possible to generate surrogate spike trains that realise the spectral estimator Eq.(28), although this is computationally very cumbersome. Again, we can take advantage of the relative computational ease of binned autocorrelations here.\footnote{Thanks to Bruce Gluckman for pointing this out to us.} Introducing a normalisation constant \(\alpha\), we can write \(C(\tau) = \alpha \int dt s(t)s(t - \tau) = \)
Thus, all we have to do is to count all possible intervals $t_i - t_j$ in a bin. The upper panel in Fig. 17 shows the binned autocorrelation function with bin size $\Delta = 0.02$ sec up to a lag of 6 sec for the heart beat data shown in Fig. 16. Superimposed is the corresponding curve for a surrogate that has been generated with the deviation from the binned autocorrelations of the data as a cost function. The two curves are practically indistinguishable. If we choose $\alpha$ such that $C(0) = 1$, we obtain:

$$C_\Delta(\tau) = \frac{|B_{ij}(\tau - \Delta, \tau)|}{N\Delta}. \quad (29)$$

Thus, all we have to do is to count all possible intervals $t_i - t_j$ in a bin. The upper panel in Fig. 17 shows the binned autocorrelation function with bin size $\Delta = 0.02$ sec up to a lag of 6 sec for the heart beat data shown in Fig. 16. Superimposed is the corresponding curve for a surrogate that has been generated with the deviation from the binned autocorrelations of the data as a cost function. The two curves are practically indistinguishable. If we choose $\alpha$ such that $C(0) = 1$, we obtain:

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7 Questions of interpretation

Having set up all the ingredients for a statistical hypothesis test of nonlinearity, we may ask what we can learn from the outcome of such a test. The formal answer is of course that we have, or have not, rejected a specific hypothesis at a given level of significance. How interesting this information is, however, depends on the null hypothesis we have chosen. The test is most meaningful if the null hypothesis is plausible enough so that we are prepared to believe it in the lack of evidence against it. If this is not the case, we may be tempted to go beyond what is justified by the test in our interpretation. Take as a simple example a recording of hormone concentration in a human. We can test for the null hypothesis of a stationary Gaussian linear random process by comparing the data to phase randomised Fourier surrogates. Without any test, we know that the hypothesis cannot be true since hormone concentration, unlike Gaussian variates, is strictly non-negative. If we failed to reject the null hypothesis by a statistical argument, we will therefore go ahead and reject it anyway by common sense, and the test was pointless. If we did reject the null hypothesis by finding a coarse-grained “dimension” which is significantly lower in the data than in the surrogates, the result formally does not give any new information but we might be tempted to speculate on the possible interpretation of the “nonlinearity” detected.

This example is maybe too obvious, it was meant only to illustrate that the hypothesis we test against is not often what we would actually accept to be true. Other, less obvious and more common, examples include signals which are known (or found by inspection) to be non-stationary (which is not covered by most null hypotheses), or signals which are likely to be measured in a static nonlinear, but non-invertible way. Before we discuss these two specific caveats in some more detail, let us illustrate the delicacy of these questions with a real data example.

Figure 18 shows as an intra-cranial recording of the neuronal electric field during an epileptic seizure, together with one iteratively generated surrogate data set [30] that has the same amplitude distribution and the same linear correlations or frequency content as the data. We have eliminated the end-point mismatch by truncating the series to 1875 samples. A test was scheduled at the 99% level of significance, using nonlinear prediction errors (see Eq. (3); $m = 3, \tau = 5, \epsilon = 0.2$) as a discriminating statistics. The nonlinear correlations we are looking for should enhance predictability and we can thus perform a one-sided test for a significantly smaller error. In a test with one data set and 99 surrogates, the likelihood that the data would
yield the smallest error by mere coincidence is exactly 1 in 100. Indeed, as can be seen in Fig. 19, the test just rejects the null hypothesis.

Unfortunately, the test itself does not give any guidance as to what kind of nonlinearity is present and we have to face notoriously ill-defined questions like what is the most natural interpretation. Similar spike-and-wave dynamics as in the present example has been previously reported as chaotic, but these findings have been questioned. Hernández and coworkers have suggested a stochastic limit cycle as a simple way of generating spike-and-wave-like dynamics.

If we represent the data in time delay coordinates — which is what we would usually do with chaotic systems — the nonlinearity is reflected by the “hole” in the centre (left panel in Fig. 20). A linear stochastic process could equally well show oscillations, but its amplitude would fluctuate in a different way, as we can see in the right panel of the same figure for an iso-spectral surrogate. It is difficult to answer the question if the nonlinearity could have been generated by a static mechanism like the measurement process (beyond the invertible rescaling allowed by the null hypothesis). Deterministic chaos in a narrower sense seems rather unlikely if we regard the prediction errors shown in Fig. 19. Although significantly lower than that of the surrogates, the absolute value of the nonlinear prediction error is still more than 50% of the rms amplitude of the data (which had been rescaled to unit variance). Not surprisingly, the correlation integral (not shown here) does not show any proper scaling region either. Thus, all we can hand back to the clinical researchers is a solid statistical result but the insight into what process is generating the oscillations is limited.

A recent suggestion for surrogates for the validation of unstable periodic orbits (UPOs) may serve as an example for the difficulty in interpreting results for more fancy null hypothesis. Dolan and coworkers coarse-grain amplitude adjusted data in order to extract a transfer matrix that can be iterated to yield typical realisations of a Markov chain. The rationale there is to test if the finding of a certain number of UPOs could be coincidental, that is, not generated

5 Contrary to what is said in Ref. 24, binning a two dimensional distribution yields a first order (rather than a second order) Markov process, for which a three dimensional binning would be needed to include the image distribution as well.
by dynamical structure. Testing against an order $D$ Markov model removes dynamical structure beyond the “attractor shape” (AS) in $D + 1$ dimensions. It is not clear to us what the interpretation of such a test would be. In the case of a rejection, they would infer a dynamical nature of the UPOs found. But that would most probably mean that in some higher dimensional space, the dynamics could be successfully approximated by a Markov chain acting on a sufficiently fine mesh. This is at least true for finite dimensional dynamical systems. In other words, we cannot see what sort of dynamical structure would generate UPOs but not show its signature in some higher order Markov approximation.

7.1 Non-dynamic nonlinearity

A non-invertible measurement function is with current methods indistinguishable from dynamic nonlinearity. The most common case is that the data are squared moduli of some underlying dynamical variable. This is supposed to be true for the celebrated sunspot numbers. Sunspot activity is generally connected with magnetic fields and is to first approximation proportional to the squared field strength. Obviously, sunspot numbers are non-negative, but also the null hypothesis of a monotonically rescaled Gaussian linear random process is true for the underlying signal, it is usually not true for filtered copies of it, in particular sequences of first differences, see Prichard [50] for a discussion of this problem.

The catch is that nonlinear deterministic dynamical systems may produce irregular time evolution, or chaos, and the signals generated by such processes will be easily found to be nonlinear by statistical methods. But many authors have confused cause and effect in this logic: deterministic chaos does imply nonlinearity, but not vice versa. The confusion is partly due to the heavy use of methods inspired by chaos theory, leading to arguments like “If the fractal dimension algorithm has power to detect nonlinearity, the data must have a fractal attractor!” Let us give a very simple and commonplace example where such a reasoning would lead the wrong way.

One of the most powerful [6,11,13] indicators of nonlinearity in a time series is the change of statistical properties introduced by a reversal of the time direction: Linear stochastic processes are fully characterised by their power spectrum which does not contain any information on the direction of time. One of the simplest ways to measure time asymmetry is by taking the first differences of the series to some power, see Eq.(3). Despite its high discrimination power, also for many but not all dynamical nonlinearities, this statistic has not been very popular in recent studies, probably since it is rather unspecific about the nature of the nonlinearity. Let us illustrate this apparent flaw by an example where time reversal asymmetry is generated by the measurement process.

Consider a signal generated by a second order autoregressive (AR(2)) process $x_n = 1.6x_{n-1} - 0.61x_{n-2} + \eta_n$. The sequence $\{\eta_n\}$ consists of independent Gaussian random numbers with a variance chosen such that the data have unit variance. A typical output of 2000 samples is shown as the upper panel in Fig. 21. Let the measurement be such that the data is rescaled by the strictly monotonic function $s_n = e^{x_n/2}$. The resulting sequence (see the lower panel in Fig. 21) still satisfies the null hypothesis formulated above. This is no longer the case if we take differences of this signal, a linear operation that superficially seems harmless for a “linear” signal. Taking differences turns the up-down-asymmetry of the data into a forward-backward asymmetry. As it has been pointed out by Prichard, [50] the static nonlinearity and linear filtering are not interchangeable with respect to the null hypothesis and the sequence $\{z_n = s_n - s_{n-5} = e^{x_n/2} - e^{x_{n-5}/2}\}$ must be considered nonlinear in the sense that it violates the null hypoth-

![Figure 21](image-url)
Figure 22: Moving differences $s_n - s_{n-5}$ of the sequence shown in Fig. 21 (upper), and a surrogate time series (lower). A formal test shows that the nonlinearity is significant at the 99% level.

Figure 23: A single spike is artificially introduced in an otherwise linear stochastic time sequence (upper). In the surrogate time series (lower), this leads to multiple short spikes. Although the surrogate data has the same frequency content and takes on the same set of values as the data, the remnants of the spike will lead to the detection of nonlinearity.

This impulse leads to the formation of a rather large spike. Such a sequence is shown in Fig. 23. Note that due to the correlations in the process, the spike covers more than a single measurement.

When we generate surrogate data, the first observation we make is that it takes the algorithm more than 400 iterations in order to converge to a reasonable tradeoff between the correct spectrum and the required distribution of points. Nevertheless, the accuracy is quite good — the spectrum is correct within 0.1% of the rms amplitude. Visual inspection of the lower panel of Fig. 23 shows that the spectral content — and the assumed values — during the single spike are represented in the surrogates by a large number of shorter spikes. The surrogates cannot know of an external kick. The visual result can be confirmed by a statistical test with several surrogates, equally well (99% significance) by a time asymmetry statistic or a nonlinear prediction error.

If non-stationarity is known to be present, it is necessary to include it in the null hypothesis explicitly. This is in general very difficult but can be undertaken in some well behaved cases. In Sec. 6.1 we discussed the simplest situation of a slow drift in the calibration of the data. It has been shown empirically that a slow drift in system parameters is not as harmful as expected. It is possible to generate surrogates for sliding windows and restrict the discriminating statistics to exclude the points at the window boundaries.

7.2 Non-stationarity

It is quite common in bio-medical time series (and elsewhere) that otherwise harmless looking data once in a while are interrupted by a singular event, for example a spike. It is now debatable whether such spikes can be generated by a linear process by nonlinear rescaling. We do not want to enter such a discussion here but merely state that a time series that covers only one or a few such events is not suitable for the statistical study of the spike generation process. The best working assumption is that the spike comes in by some external process, thus rendering the time series non-stationary. In any case, the null hypotheses we are usually testing against are not likely to generate such singular events autonomously. Thus, typically, a series with a single spike will be found to violate the null hypothesis, but, arguably, the cause is non-stationarity rather than non-linearity. Let us discuss as a simple example the same AR(2) process considered previously, this time without any rescaling. Only at a single instant, $n = 1900$, the system is kicked by a large impulse instead of the Gaussian variate $\eta_{1900}$. 

Noticing such a sequence (see the upper panel in Fig. 22) is found to be nonlinear at the 99% level of significance using the statistics given in Eq. (3), but also using nonlinear prediction errors. (Note that the nature of the statistic Eq. (3) requires a two-sided test.) A single surrogate series is shown in the lower panel of Fig. 22. The tendency of the data to raise slowly but to fall fast is removed in the linear surrogate, as it should.
It is quite obvious that special care has to be taken in such an analysis.

8 Conclusions: Testing a Hypothesis vs. Testing Against Surrogates

Most of what we have to say about the interpretation of surrogate data tests, and spurious claims in the literature, can be summarised by stating that there is no such thing in statistics as testing a result against surrogates. All we can do is to test a null hypothesis. This is more than a difference in words. In the former case, we assume a result to be true unless it is rendered obsolete by finding the same with trivial data. In the latter case, the only one that is statistically meaningful, we assume a more or less trivial null hypothesis to be true, unless we can reject it by finding significant structure in the data.

As everywhere in science, we are applying Occam’s razor: We seek the simplest — or least interesting — model that is consistent with the data. Of course, as always when such categories are invoked, we can debate what is “interesting”. Is a linear model with several coefficients more or less parsimonious than a nonlinear dynamical system written down as a one line formula? People unfamiliar with spectral time series methods often find their use and interpretation at least as demanding as the computation of correlation dimensions. From such a point of view it is quite natural to take the nonlinearity of the world for granted, while linearity needs to be established by a test against surrogates.

The reluctance to take surrogate data as what they are, a means to test a null hypothesis, is partly explainable by the choice of null hypotheses which are currently available for proper statistical testing. As we have tried to illustrate in this paper, recent efforts on the generalisation of randomisation schemes broaden the repertoire of null hypotheses. The hope is that we can eventually choose one that is general enough to be acceptable if we fail to reject it with the methods we have. Still, we cannot prove that there is no dynamics in the process beyond what is covered by the null hypothesis. From a practical point of view, however, there is not much of a difference between structure that is not there and structure that is undetectable with our observational means.

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A The TISEAN implementation

Starting with the publication of source code for a few nonlinear time series algorithms by Kantz and Schreiber, a growing number of programs has been put together to provide researchers with a library of common tools. The TISEAN software package is freely available in source code form and an introduction to the contained methods has been published in Ref. More recent versions of the package contain a comprehensive range of routines for the generation and testing of surrogate data. The general constrained randomisation scheme described in Sec. is implemented as an extendable framework that allows for the addition of further cost functions with relatively little effort. With few exceptions, all the code used in the examples in this paper is publicly available as part of TISEAN 2.0.

A.1 Measures of nonlinearity

A few programs in the package directly issue scalar quantities that can be used in nonlinearity testing. These are the zeroth order nonlinear predictors (predict and zeroth) which implement Eq. and the time reversibility statistic (timeres) implementing Eq. For a couple of other quantities, we have deliberately omitted a black box algorithm to turn the raw results into a single number. A typical example are the programs for dimension estimation (d2, c2, c2naive, and c1) which compute correlation sums for ranges of length scales \( \epsilon \) and embedding dimensions \( m \). For dimension estimation, these curves have to be interpreted with due care to establish scaling behaviour and convergence with increasing \( m \). Single numbers issued by black box routines have led to too many spurious results in the literature. Researchers often
forget that such numbers are not interpretable as fractal dimensions at all but only useful for comparison and classification. Without genuine scaling at small length scales, a data set that gives $D_2 = 4.2$ by some ad hoc method to estimate $D_2$ cannot be said to have more degrees of freedom, or be more “complex” than one that yields $D_2 = 3.5$.

This said, users are welcome to write their own code to turn correlation integrals, local slopes ($c2d$), Takens’ estimator ($c2t$), or Gaussian Kernel correlation integrals ($c2g$) into nonlinearity measures. The same situation is found for Lyapunov exponents ($lyap$), $k$, $1lyap_r$), entropies ($boxcount$) and other quantities. Since all of these have already been described in Ref. 8, we refer the reader there for further details.

A.2 Iterative FFT surrogates

The workhorse for the generation of surrogate data within the TISEAN package is the program surrogates. It implements the iterative Fourier based scheme introduced in Ref. 30 and discussed in Sec. 5. It has been extended to be able to handle multivariate data as discussed in Sec. 6. An FFT routine is used that can handle data sets of $N$ points if $N$ can be factorised using prime factors 2, 3, and 5 only. Routines that take arbitrary $N$ will end up doing a slow Fourier transform if $N$ is not factorisable with small factors. Occasionally, the length restriction results in the loss of a few points.

The routine starts with a random scramble as $\left\{ \tau^{(0)}_n \right\}$, performs as many iterates as necessary to reach a fixed point and then prints out $\tau^{(\infty)}_n$ or $s^{(\infty)}_n$, as desired. Further, the number of iterations is shown and the residual root mean squared discrepancy between $\tau^{(\infty)}_n$ and $s^{(\infty)}_n$. The number of iterations can be limited by an option. In particular, $i = 0$ gives the initial scramble as $\left\{ \tau^{(0)}_n \right\}$ or a non-rescaled FFT surrogate as $\left\{ \tau^{(0)}_n \right\}$. The first iterate, $\left\{ \tau^{(1)}_n \right\}$, is approximately (but not quite) equivalent to an AAFT surrogate. It is advisable to evaluate the residual discrepancy whenever the algorithm took more than a few iterations. In cases of doubt if the accuracy is sufficient, it may be useful to plot the autocorrelation function (corr or autocor) of the data and $\tau^{(\infty)}_n$, and, in the multivariate case, the cross-correlation function (xcor) between the channels. The routine can generate up to 999 surrogates in one call.

Since the periodicity artefact discussed in Sec. 6.3 can lead to spurious test results, we need to select a suitable sub-sequence of the data before making surrogates. For this purpose, TISEAN contains the program endtoend. Let $\left\{ s^{(n_0)}_n \right\} = s_{n+n_0}$ be a subsequence of length $\bar{N}$ and offset $n_0$. The program then computes the contribution of the end-to-end mismatch $(s^{(n_0)}_1 - s^{(n_0)}_{\bar{N}})^2$ to the total power in the sub-sequence:

$$\gamma^{(\text{jump})} = \frac{(s^{(n_0)}_1 - s^{(n_0)}_{\bar{N}})^2}{\sum_{n=1}^{\bar{N}} (s^{(n_0)}_n - \langle s^{(n_0)}_n \rangle)^2}$$

as well as the contribution of the mismatch in the first derivative

$$\gamma^{(\text{slip})} = \frac{\left[ (s^{(n_0)}_2 - s^{(n_0)}_1) - (s^{(n_0)}_\bar{N} - s^{(n_0)}_{\bar{N}-1}) \right]^2}{\sum_{n=1}^{\bar{N}} (s^{(n_0)}_n - \langle s^{(n_0)}_n \rangle)^2}$$

and the weighted average

$$\gamma^{(\bar{N},n_0)} = w \gamma^{(\text{jump})} + (1 - w) \gamma^{(\text{slip})}.$$  

The weight $w$ can be selected by the user and is set to 0.5 by default. For multivariate data with $M$ channels, $(1/M) \sum_{m=1}^{M} \gamma^{(\bar{N},n_0)}_m$ is used.

Now the program goes through a sequence of decreasing $\bar{N} = 2^i 3^j 5^k$, $i,j,k \in \mathbb{N}$, and for each $\bar{N}$ determines $n_0^\ast$ such that $\gamma^{(\bar{N},n_0^\ast)}$ is minimal. The values of $\bar{N}$, $n_0^\ast$, and $\gamma^{(\bar{N},n_0^\ast)}$ are printed whenever $\gamma$ has decreased. One can thus easily find a sub-sequence that achieves negligible end point mismatch with the minimal loss of data.

A.3 Annealed surrogates

For cases where the iterative scheme does not reach the necessary accuracy, or whenever a more general null hypothesis is considered, the TISEAN package offers an implementation of the constrained randomisation algorithm using a cost function minimised by simulated annealing, as introduced in Ref. 26 and described in Sec. 5. Since one of the main advantages of the approach is its flexibility, the implementation more resembles a toolbox than a single program. The main driving routine randomize takes care of the data input and output and operates the simulated annealing procedure. It must be linked together with modules that implement a cooling schedule, a cost function, and a permutation scheme. Within TISEAN, several choices for each of these are already implemented but it is relatively easy to add individual variants or completely different cost functions, cooling or permutation schemes. With the development structure provided, the final executables will then have names reflecting the components linked together, in the form randomize\_A\_B\_C, where $A$ is a cost function module, $B$ a cooling scheme, and $C$ a permutation scheme.

Currently, two permutation schemes are implemented. In general, one will use a scheme random
that selects a pair at random. It is, however, possible to specify a list of points to be excluded from the permutations. This is useful when the time series contains artifacts or some data points are missing and have been replaced by dummy values. It is planned to add a temperature-sensitive scheme that selects pairs close in magnitude at low temperatures. For certain cost functions (e.g. the spike train spectrum), an update can only be carried out efficiently if two consecutive points are exchanged. This is implemented in an alternative permutation scheme event.

The only cooling scheme supported in the present version of TISEAN (2.0) is exponential cooling \((\exp)\). This means that whenever a certain condition is reached, the temperature is multiplied by a factor \(\alpha < 1\). Apart from \(\alpha\) and the initial temperature \(T_0\), two important parameters control the cooling schedule. Cooling is performed either if a maximal total number of trials \(S_{\text{total}}\) is exceeded, or if a maximal number \(S_{\text{succ}}\) of trials has been successful since the last cooling. Finally, a minimal number of successes \(S_{\text{min}}\) can be specified below which the procedure is considered to be “stuck”. All these parameters can be specified explicitly. However, it is sometimes very difficult to derive reasonable values except by trial and error. Slow cooling is necessary if the desired accuracy of the constraint is high. It seems reasonable to increase \(S_{\text{succ}}\) and \(S_{\text{total}}\) with the system size, but also with the number of constraints incorporated in the cost function. It can be convenient to use an automatic scheme that starts with fast parameter settings and re-starts the procedure with slower settings whenever it gets stuck, until a desired accuracy is reached. The initial temperature can be selected automatically using the following algorithm. Start with an arbitrary small initial temperature. Let the system evolve for \(S_{\text{total}}\) steps (or \(S_{\text{succ}}\) successes). If less than \(2/3\) of the trials were successes, increase the initial temperature by a factor of ten to “melt” the system. This procedure is repeated until more than \(2/3\) successes are reached. This ensures that we start with a temperature that is high enough to leave all false minima. If the automatic scheme gets stuck (the low temperature allows too few changes to take place), it re-starts at the determined melting temperature. At the same time, the cooling rate is decreased by \(\alpha \to \sqrt{\alpha}\) and \(S_{\text{total}} \to \sqrt{2} S_{\text{total}}\). We suggest to create one surrogate with the automatic scheme and then use the final values of \(T_0\), \(\alpha\) and \(S_{\text{total}}\) for subsequent runs. Of course, other more sophisticated cooling schemes may be suitable depending on the specific situation. The reader is referred to the standard literature \[35\].

Several cost functions are currently implemented in TISEAN. Each of them is of the general form \(\|\mathbf{y}\|_2\) and the constraints can be matched in either the \(L^1\), \(L^2\), or the \(L^\infty\) (or maximum) norms. In the \(L^1\) and \(L^2\) norms, autocorrelations are weighted by \(w_\tau = 1/\tau\) and frequencies by \(w_\omega = 1/\omega\).

Autocorrelations (auto, or a periodic version autop) are the most common constraints available. Apart from the type of average, one has to specify the maximal lag \(t_{\text{max}}\) (see e.g. Eq.\[23\]). This can save a substantial fraction of the computation time if only short range correlations are present. For each update, only \(O(t_{\text{max}})\) terms have to be updated.

For unevenly sampled data (see Sec. \[6\]), the cost function uneven implements binned autocorrelations as defined by Eq.\[20\]. The update of the histogram at each annealing step takes a number of steps proportional to the number of bins. The user has to specify the bin size \(\Delta\) and the total lag time covered contiguous by the bins.

For surrogate spike trains, either the spike train periodogram Eq.\[23\] or binned correlations Eq.\[29\] can be used. In the former case, the cost function is coded in spikespec. The user has to give the total number of frequencies and the frequency resolution. Internally, the event times \(t_n\) are used. A computationally feasible update is only possible if two consecutive intervals \(t_n - t_{n-1}\) and \(t_{n+1} - t_n\) are exchanged by \(t_n \to t_{n-1} + t_{n+1} - t_n\) (done by the permutation scheme event). As a consequence, coverage of permutation space is quite inefficient. With binned autocorrelations spikeauto, intervals are kept internally and any two intervals may be swapped, using the standard permutation scheme random.

The documentation distributed with the TISEAN package describes how to add further cost functions. Essentially, one needs to provide cost function specific option parsing and input/output functions, a module that computes the full cost function and one that performs an update upon permutation. The latter should be coded very carefully. First it is the single spot that uses most of the computation time and second, it must keep the cost function consistent for all possible permutations. It is advisable to make extensive tests against freshly computed cost functions before entering production.

In future releases of TISEAN, it is planned to include routines for cross correlations in multivariate data, multivariate spike trains, and mixed signals. We hope that users take the present modular implementation as a starting point for the implementation of other null hypotheses.

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