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Nonlinear modal analysis using pattern recognition

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Abstract
The main objective of nonlinear modal analysis is to formulate a mathematical model of a nonlinear dynamical structure based on observations of input/output data from the dynamical system. Most theories regarding structural modal analysis are centred on the linear modal analysis which has proved to now to be the method of choice for the analysis of linear dynamic structures. However, for the majority of other structures, where the effect of nonlinearity becomes significant, then nonlinear modal analysis is a necessity. The objective of the current paper is to demonstrate a machine learning approach to output-only nonlinear modal decomposition using kernel independent component analysis and locally linear embedding analysis. The key element is to demonstrate a pattern recognition approach which exploits the idea of independence of principal components by learning the nonlinear manifold between the variables.

1 Introduction

The machine learning methods that are introduced in this paper aim to address the problem of validity that surrounds the modal analysis of nonlinear structures. Modal analysis is an important tool in structural dynamics as it is used to understand the dynamical characteristics of the structure. Many methods have been proposed in recent years regarding nonlinear analysis, such as nonlinear normal modes or the method of normal forms [1, 2, 3, 4, 5, 6, 7, 8, 9].

In this work a different approach is investigated through the usage of unsupervised pattern recognition techniques such as kernel independent component analysis (KICA) and locally linear embedding manifold learning (LLE). These methods serve two purposes, a reduction in the dimensionality by mapping the data from high-dimensional spaces to lower-dimensional spaces and a revealing of the hidden features of the data by learning the structure of the nonlinear manifold between the variables of interest. Of course this dimensionality reduction is accompanied by loss of some information; therefore, the goal in dimensionality reduction should be to preserve as much relevant information as possible.

The goal of these methods is one: to create uncorrelated variables but retaining the maximum possible variance of the original observations. The effect of structural nonlinearity on linear modal analysis is critical. Specifically, decoupling of the system into SDOF systems is lost and in turn superposition is lost. It is of critical importance to mention that these clever and advanced unsupervised algorithms can work with output-only data and can play a significant role in the model updating of nonlinear systems by giving crucial insight into the dynamical behaviour of the system.

The layout of the paper is as follows. Section 2 covers the main features of linear modal analysis using linear decoupling methods such as principal component analysis, while section 3 discusses an alternative approach of independent component analysis (ICA). Section 4 gives an example of nonlinear modal analysis based on the unsupervised learning techniques that are mentioned in sections 2 and 3. Section 5 discusses how the
previous approaches break down for multi-degrees-of-freedom systems with high nonlinearity and a new approach based on measured data such as locally linear embedding method is needed. The paper finishes with some overall conclusion and future work.

\section{Principal component analysis}

Principal Component Analysis takes a multivariate data set and maps it onto a new set of variables called “principal components”, which are linear combinations of the old variables. The first principal component will account for the highest amount of the variance in the data set and the second principal component will account for the second highest variance in the data set independent of the first, and so on. The importance of the method arises from the fact that, in terms of mean-squared-error of reconstruction, it is the optimal linear tool for compressing data of high dimension into data of lower dimension. The unknown parameters of the transformation can be computed directly from the raw data set and, once all parameters are derived, compression and decompression are small operations based on matrix algebra \cite{10, 11, 12}. One has,

\begin{equation}
[X] = [K][Y] \tag{1}
\end{equation}

Where \([Y]\) represents the original input data with size \(p \times n\), with \(p\) number of variables and \(n\) the number of data sets, \([X]\) is the scores matrix of reduced dimension \(q \times n\) where \(q < p\) contains the transformed variables and \([K]\) is called the loading matrix. The columns of \([K]\) are the eigenvectors corresponding to the largest eigenvalues of the covariance matrix of \([Y]\). The covariance matrix is equal to:

\begin{equation}
[S] = E \left[ (\{Y\} - \{\bar{Y}\}) (\{Y\} - \{\bar{Y}\})^T \right] \tag{2}
\end{equation}

where \(E\) is the expectation operator and \(\bar{Y}\) is the mean value.

The original data reconstruction is performed by the inverse of equation (1):

\begin{equation}
[Y] = [K]^T[X] \tag{3}
\end{equation}

The information loss of the mapping procedure is calculated in the reconstruction error matrix:

\begin{equation}
[E] = [Y] - [\hat{Y}] \tag{4}
\end{equation}

For further information on PCA, readers are referred to any text book on multivariate analysis (examples being references \cite{10, 11}).

\section{Kernel independent component analysis}

Independent component analysis (ICA) is a tool that recovers a latent random vector \(\{x\} = (x_1, ..., x_m)\) from measurements of \(m\) unknown linear functions of that vector. The components of \(\{x\}\) are required to be mutually independent. As a result an observation \(\{y\} = (y_1, ..., y_m)\) is modelled as \cite{13, 14, 15}:

\begin{equation}
\{y\} = [A]\{x\} \tag{5}
\end{equation}

where \([A]\) is an \(m \times m\) matrix of parameters.

If \([W] = [A]^{-1}\) is the parameter matrix inverse then the estimate of \([\hat{W}]\) can be calculated by giving an estimate of the latent independent components such as:
\[
\{ \hat{x} \} = [W]\{y\} \quad (6)
\]

It can be shown [13, 14, 15] that minimising the mutual information between the components of (6) is essentially a contrast function minimisation.

Contrast functions are statistical functions that are capable of separating or extracting independent components from a data mixture [15]. If a contrast function is derived by the \(F\)-correlation statistics, it can be defined as the maximum correlation between the tested random variables \(f_1\) and \(f_m\) [15] and can be written as:

\[
pf = \max_{f_1, f_m \in f} \text{corr}(f_1(x_1), f_m(x_m)) = \max_{f_1, f_m \in f} \frac{\text{cov}(f_1(x_1), f_m(x_m))}{\left(\text{var}(f_1(x_1))\right)^{1/2} \left(\text{var}(f_m(x_m))\right)^{1/2}} \quad (7)
\]

for each \(i\ldots m\), of estimated source vectors such as \(\{x\} = (x_1, \ldots, x_m)\). This contrast function is equal to zero only if the variables are independent.

Different methods have been introduced in the literature regarding ICA that make use of different nonlinear contrast functions [13, 14, 15]. The nonlinear ICA method that is used in this study is kernel independent component analysis (KICA) which makes use of the “kernel trick” which is an algorithm that uses a multiple nonlinear functions but through an entire function space of a family of candidate nonlinearities. The “kernel trick” is basically forcing the functions to work in a reproducing kernel Hilbert space.

Given the nature of the current paper a full description of the complicated algorithm is not possible but for further information on ICA and Kernel ICA, readers are referred to [13, 14, 15].

Briefly the general outline of algorithm is as follows:

If one assumes \([y] = (\{y_1\}, \ldots, \{y_m\})\) of data vectors and the parameter matrix \([W]\) of equation (6), and set \(\{x\} = [W]\{y\}\) then one can derive a set of estimated source vectors such as \(\{x\} = (\{x_1\}, \ldots, \{x_m\})\). The \(m\) components of these vectors lead to a set of \(m\) centered kernel Gram matrices, \([K_1], \ldots, [K_m]\).

Briefly, a Gram matrix can be generally defined as, \(K_{ij} = K(x_i, x_j)\), which is positive semidefinite Kernel matrix [15]. This kernel \([K]\) matrix is accompanied by a mapping of a function \(\Phi\) to an \(F\)-distribution such as:

\[
K(x, y) = \langle \Phi(x), \Phi(y) \rangle \quad (8)
\]

This kernel can be then used to compute the inner product in the \(F\)-distribution space. This is often called the kernel trick. These kernel matrices can then be used in order to define a contrast function [15]:

\[
C(W) = \hat{I}_{pf}([K_1], \ldots, [K_m]) \quad (9)
\]

where \(\hat{I}_{pf}\) is a contrast function given by:

\[
\hat{I}_{pf} = -\frac{1}{2} \log \left(1 - \max_{f_1, f_m \in f} \text{corr}(f_1(x_1), f_m(x_m))\right) \quad (10)
\]

This valid contrast function is derived by \(F\)-correlation statistics and is defined as the maximum correlation between the tested random variables \(f_1\) and \(f_m\) [15].

It has very useful properties as it is nonnegative and equal to zero only if the variables are independent. The kernel ICA algorithm involves minimising this function \(C(W)\) with respect to the matrix \([W]\). And this is called the kernelised canonical correlation algorithm (KCCA) [15] (which is mainly used in this study). Canonical correlation analysis (CCA) is a multivariate method similar in nature to PCA. The main difference
is that while PCA works with a single random vector and maximises the variance of projections of the
observations, CCA works with a set of $m$ random vectors by maximising the correlation between sets of
projections [15]. One needs to remember that PCA solves an eigenvector problem, CCA solves a generalized
eigenvector problem.

Another contrast function which can be defined is via the kernel generalised variance (KGV) algorithm which
suggests defining a corresponding quantity for kernelized canonical correlation analysis [15]. For further
information readers are referred to [15].

The basic concept that one has to remember is that ICA can remove correlations and higher order dependences
between the variables compared to PCA (which can only go up to second order statistics).

4 An example

The system of interest will be a nonlinear two-DOF lumped parameter system (see Fig.1). Data were simulated
using a fixed-step $4^{th}$-order Runge-Kutta algorithm and the excitation was chosen to be a Gaussian white
noise sequence with zero mean and unit variance and the associated displacements were extracted. The model
parameters adopted were: $m = 0.1$, $c_1 = 0.005$, $c_2 = 0.01$, $k_1 = 50$, $k_2 = 100$, $k_{nl} = 10^4$. The nonlinearity
that is assumed is cubic. It has to be noted that the damping is proportional, so the underlying linear system
uncouples.

The method that is used in order to calculate the power spectral densities (PSDs) which follow is the Welch
method based on time averaging over short, modified periodograms which could decolour the effect of
different random excitation inputs [16]. The signals are split into sections and the periodograms of each
section are averaged. Through the Welch method these data sections are overlapped and a window, such as
the Hanning window is applied in order to filter each section. The overlapping of the signal sections is usually
either 50% (as in this paper) or 75%.

Fig.2 shows the results of PSDs for the simulated physical variables. Both modes are present in the PSDs
for the transformed coordinates which shows that the system is clearly not uncoupled. For all the graphs the
vertical axe is the PSD of displacement and the frequency is in Hz.

As can be seen in Fig.3, PCA fails in decoupling the nonlinear system (standard linear modal analysis) but
kernel ICA, as seen in Fig.4, is successfully decoupling the nonlinear system into two SDOF systems due to
the removal of the higher order statistical dependence. Standard linear modal analysis is equivalent to PCA in
this case as the mass matrix is diagonal.

![Figure 1: Nonlinear two-DOF lumped parameter system.](image-url)
Figure 2: PSDs for physical variables.

Figure 3: PSDs for transformed variables: standard linear modal analysis (PCA).
5 A three-degree-of-freedom system

In order to validate the results further, a more complicated system in terms of degrees of freedom and increased cubic nonlinearity is discussed (see Fig.5). As can be seen in Figs.6-8, both PCA and kernel ICA lack in efficiency and performance in decoupling the nonlinear modes of the system. This is the reason that a novel approach to structural dynamics is introduced next in the form of the local linear embedding method. The system of interest will be a nonlinear three-DOF lumped parameter system. Data were simulated using a fixed-step 4th-order Runge-Kutta algorithm and the excitation was chosen to be a Gaussian white noise sequence with zero mean and unit variance. The model parameters adopted were: \( m = 0.1, c_1 = 0.01, c_2 = 0.02, c_3 = 0.03, k_1 = 50, k_2 = 150, k_3 = 300, k_{nl} = 10^5 \). The nonlinearity that is assumed is cubic.

5.1 Nonlinear manifold learning via locally linear embedding

As can be seen in the previous section the combination of stronger nonlinearity with multi-degree of freedom systems makes the performance of both the PCA and ICA algorithm very weak. Neither of them can decouple successfully the nonlinear modes. This is the reason that a quick and effective method of nonlinear manifold learning such as locally linear embedding is introduced in nonlinear modal analysis here [17, 18].
Figure 6: PSDs for physical variables.

Figure 7: PSDs for transformed variables: standard linear modal analysis (PCA).
Other very strong methods can be applied in such complex nonlinear manifolds such as nonlinear principal component analysis via the usage of auto-associative neural networks [11, 19, 20]. The usage of such methods is SHM can be seen in [21]. For the current study LLE is used as it is a novel introduction into nonlinear modal analysis and a much simpler tool.

An extensive overview of the algorithm can be found in [17, 18]. Briefly and for the purposes of this paper a short description is discussed.

The LLE method is based on simple geometric intuition. If the observations consist of \( N \) real-valued vectors \( \{ x_i \} \) with dimensions \( D \) and they are sampled from a smooth underlying nonlinear manifold, then each data point and its neighbours is expected to lie on or close to a locally formed patch of the manifold. This local geometries can be characterised by finding linear coefficients that can reconstruct each data point with respect to each set of neighbours.

If one establishes \( K \) nearest neighbours per data point then the reconstruction error is given by the cost function:

\[
error(W) = \sum_i \left| \{ y_i \} - \sum_j [W_{ij}] \{ x_j \} \right|^2
\]

(11)

where \([W_{ij}]\) is the weight contribution of the \( j_{th} \) data point to the \( i_{th} \) reconstruction. In order to compute these weights the cost function has to be minimised under the following constraints. The reconstruction errors that are subject to the constrained weights should be invariant to rotations and rescaling. In turn, in order that the LLE algorithm preserve this invariant manifold idea as a final step of the method, each measurement \( \{ x_i \} \) should be mapped to lower dimensional vector \( \{ y_i \} \) that minimises the cost function:

\[
error(Y) = \sum_i \left| \{ y_i \} - \sum_j [W_{ij}] \{ y_j \} \right|^2
\]

(12)
The main difference with the previous cost function is that here the weights are fixed but the $\{Y_i\}$ co-ordinates are optimised.

In Fig. 9 the LLE method is shown to successfully decoupling the modes as it was able to unfold and learn the underlying nonlinear manifold.

![Figure 9: PSDs for transformed variables: Local linear embedding.](image)

6 Conclusion

The purpose of this paper is to highlight the key utility of some machine learning methods, not only for dynamic analysis of structure but as well as a method of reduction for nonlinear mechanical systems. The main benefit of the approach taken here is that complicated algebraic analysis is not necessary. Furthermore, the physical equations of the system are not needed.

The biggest advantage of these methods is that one can build for several datasets the nonlinear subspace manifold only once and it then can be used for future testing datasets. As a result, this machine learning approach is suited to experimental investigation of nonlinear systems using only the measured output responses. A further work in the form of a journal article is under preparation where other multi-degree of freedom systems are investigated as well as experimental validation of the methods.

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