Applying Neural Network Models to Prediction and Data Analysis in Meteorology and Oceanography

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ABSTRACT

Empirical or statistical methods have been introduced into meteorology and oceanography in four distinct stages: 1) linear regression (and correlation), 2) principal component analysis (PCA), 3) canonical correlation analysis, and recently 4) neural network (NN) models. Despite the great popularity of the NN models in many fields, there are three obstacles to adapting the NN method to meteorology–oceanography, especially in large-scale, low-frequency studies: (a) nonlinear instability with short data records, (b) large spatial data fields, and (c) difficulties in interpreting the nonlinear NN results. Recent research shows that these three obstacles can be overcome. For obstacle (a), ensemble averaging was found to be effective in controlling nonlinear instability. For (b), the PCA method was used as a prefilter for compressing the large spatial data fields. For (c), the mysterious hidden layer could be given a phase space interpretation, and spectral analysis aided in understanding the nonlinear NN relations. With these and future improvements, the nonlinear NN method is evolving to a versatile and powerful technique capable of augmenting traditional linear statistical methods in data analysis and forecasting; for example, the NN method has been used for El Niño prediction and for nonlinear PCA. The NN model is also found to be a type of variational (adjoint) data assimilation, which allows it to be readily linked to dynamical models under adjoint data assimilation, resulting in a new class of hybrid neural–dynamical models.

1. Introduction

The introduction of empirical or statistical methods into meteorology and oceanography can be broadly classified as having occurred in four distinct stages: 1) Regression (and correlation analysis) was invented by Galton (1885) to find a linear relation between a pair of variables \(x\) and \(z\). 2) With more variables, the principal component analysis (PCA), also known as the empirical orthogonal function (EOF) analysis, was introduced by Pearson (1901) to find the correlated patterns in a set of variables \(x_1, \ldots, x_n\). It became popular in meteorology only after Lorenz (1956) and was later introduced into physical oceanography in the mid-1970s. 3) To linearly relate a set of variables \(x_1, \ldots, x_n\) to another set of variables \(z_1, \ldots, z_m\), the canonical correlation analysis (CCA) was invented by Hotelling (1936), and became popular in meteorology and oceanography after Barnett and Preisendorfer (1987). It is remarkable that in each of these three stages, the technique was first invented in a biological–psychological field, long before its adaptation by meteorologists and oceanographers many years or decades later. Not surprisingly, stage 4, involving the neural network (NN) method, which is just starting to appear in meteorology–oceanography, also had a biological–psychological origin, as it developed from investigations into the human brain function. It can be viewed as a generalization of stage 3, as it nonlinearly relates a set of variables \(x_1, \ldots, x_n\) to another set of variables \(z_1, \ldots, z_m\).

The human brain is one of the great wonders of nature—even a very young child can recognize people and objects much better than the most advanced artificial intelligence program running on a supercom-
puter. The brain is vastly more robust and fault tolerant than a computer. Though the brain cells are continually dying off with age, the person continues to recognize friends last seen years ago. This robustness is the result of the brain’s massively parallel computing structure, which arises from the neurons being interconnected by a network. It is this parallel computing structure that allows the neural network, with typical neuron “clock speeds” of a few milliseconds, about a millionth that of a computer, to outperform the computer in vision, motor control, and many other tasks.

Fascinated by the brain, scientists began studying logical processing in neural networks in the 1940s (McCulloch and Pitts 1943). In the late 1950s and early 1960s, F. Rosenblatt and B. Widrow independently investigated the learning and computational capability of neural networks (Rosenblatt 1962; Widrow and Sterns 1985). The perceptron model of Rosenblatt consists of a layer of input neurons, interconnected to an output layer of neurons. After the limits of the perceptron model were found (Minsky and Papert 1969), interests in neural network computing waned, as it was felt that neural networks (also called connectionist models) did not offer any advantage over conventional computing methods.

While it was recognized that the perceptron model was limited by having the output neurons linked directly to the neurons, the more interesting problem of having one or more “hidden” layers of neurons (Fig. 1) between the input and output layer was mothballed due to the lack of an algorithm for finding the weights interconnecting the neurons. The hiatus ended in the second half of the 1980s, with the rediscovery of the back-propagation algorithm (Rumelhart et al. 1986), which successfully solved the problem of finding the weights in a model with hidden layers. Since then, NN research rapidly became popular in artificial intelligence, robotics, and many other fields (Crick 1989). Neural networks were found to outperform the linear Box–Jenkins models (Box and Jenkins 1976) in forecasting time series with short memory and at longer lead times (Tang et al. 1991). During 1991–92, the Santa Fe Institute sponsored the Santa Fe Time Series Prediction and Analysis Competition, where all prediction methods were invited to forecast several time series. For every time series in the competition, the winner turned out to be an NN model (Weigend and Gershenfeld 1994).

Among the countless applications of NN models, pattern recognition provides many intriguing examples; for example, for security purposes, the next generation of credit cards may carry a recorded electronic image of the owner. Neural networks can now correctly verify the bearer based on the recorded image, a nontrivial task as the illumination, hair, glasses, and mood of the bearer may be very different from those in the recorded image (Javidi 1997). In artificial speech, NNs have been successful in pronouncing English text (Sejnowski and Rosenberg 1987). Also widely applied to financial forecasting (Trippi and Turban 1993; Gately 1995; Zirilli 1997), NN models are now increasingly being used by bank and fund managers for trading stocks, bonds, and foreign currencies. There are well over 300 books on artificial NNs in the subject guide to Books in Print, 1996–97 (published by R.R. Bowker), including some good texts (e.g., Bishop 1995; Ripley 1996; Rojas 1996).

By now, the reader must be wondering why these ubiquitous NNs are not more widely applied to meteorology and oceanography. Indeed, the purpose of this paper is to examine the serious problems that can arise when applying NN models to meteorology and oceanography, especially in large-scale, low-frequency studies, and the attempts to resolve these problems. Section 2 explains the difficulties encountered with NN models, while section 3 gives a basic introduction to NN models. Sections 4–6 examine various solutions to overcome the difficulties mentioned in section 2. Section 7 applies the NN method to El Niño forecasting, while section 8 shows how the NN can be used as nonlinear PCA. The NN model is linked to variational (adjoint) data assimilation in section 9.

Fig. 1. A schematic diagram illustrating a neural network model with one hidden layer of neurons between the input layer and the output layer. For a feed-forward network, the information only flows forward starting from the input neurons.
2. Difficulties in applying neural networks to meteorology—oceanography

Elsner and Tsonis (1992) applied the NN method to forecasting various time series, and comparing with forecasts by autoregressive (AR) models. Unfortunately, because of the computer bug noted in their corrigendum, the advantage of the NN method was not as conclusive as in their original paper. Early attempts to use neural networks for seasonal climate forecasting were also at best of mixed success (Derr and Slutz 1994; Tang et al. 1994), showing little evidence that exotic, nonlinear NN models could beat standard linear statistical methods such as the CCA method (Barnett and Preisendorfer 1987; Barnston and Ropelewski 1992; Shabbar and Barnston 1996) or its variant, the singular value decomposition (SVD) method (Bretherton et al. 1992), or even the simpler principal oscillation pattern (POP) method (Penland 1989; von Storch et al. 1995; Tang 1995).

There are three main reasons why NNs have difficulty being adapted to meteorology and oceanography: (a) nonlinear instability occurs with short data records, (b) large spatial data fields have to be dealt with, and (c) the nonlinear relations found by NN models are far less comprehensible than the linear relations found by regression methods. Let us examine each of these difficulties.

(a) Relative to the timescale of the phenomena one is trying to analyze or forecast, most meteorological and oceanographic data records are short, especially for climate studies. While capable of modeling the nonlinear relations in the data, NN models have many free parameters. With a short data record, the problem of solving for many parameters is ill conditioned; that is, when searching for the global minimum of the cost function associated with the problem, the algorithm is often stuck in one of the numerous local minima surrounding the true global minimum (Fig. 2). Such an NN could give very noisy or completely erroneous forecasts, thus performing far worse than a simple linear statistical model. This situation is analogous to that found in the early days of numerical weather prediction, where the addition of the nonlinear advec-tive terms to the governing equations, instead of improving the forecasts of the linear models, led to disastrous nonlinear numerical instabilities (Phillips 1959), which were overcome only after extensive research on how to discretize the advec-tive terms correctly. Thus the challenge for NN models is how to control nonlinear instability, with only the relatively short data records available.

(b) Meteorological and oceanographic data tend to cover numerous spatial grids or stations, and if each station serves as an input to an NN, then the NN will have a very large number of inputs and associated weights. The optimal search for many weights over the relatively short temporal records would be an ill-conditioned problem. Hence, even though there had been some success with NNs in seasonal forecasting using a small number of predictors (Navone and Ceccatto 1994; Hastenrath et al. 1995), it was unclear how the method could be generalized to large data fields in the manner of CCA or SVD methods.

(c) The interpretation of the nonlinear relations found by an NN is not easy. Unlike the parameters from a linear regression model, the weights found by an NN model are nearly incomprehensible. Furthermore, the “hidden” neurons have always been a mystery.

![Cost function diagram](image-url)

**Fig. 2.** A schematic diagram illustrating the cost function surface, where depending on the starting condition, the search algorithm often gets trapped in one of the numerous deep local minima. The local minima labeled 2, 4, and 5 are likely to be reasonable local minima, while the minimum labeled 1 is likely to be a bad one (in that the data was not well fitted at all). The minimum labeled 3 is the global minimum, which could correspond to an overfitted solution (i.e., fitted closely to the noise in the data) and may, in fact, be a poorer solution than the minima labeled 2, 4, and 5.
Over the last few years, the University of British Columbia Climate Prediction Group has been trying to overcome these three obstacles. The purpose of this paper is to show how these obstacles can be overcome:

For obstacle a, ensemble averaging was found to be effective in controlling nonlinear instability. Various penalty, pruning, and nonconvergent methods also helped. For b, the PCA method was used to greatly reduce the dimension of the large spatial data fields. For c, new measures and visualization techniques helped in understanding the nonlinear NN relations and the mystery of the hidden neurons. With these improvements, the NN approach has evolved to a technique capable of augmenting the traditional linear statistical methods currently used.

Since the focus of this review is on the application of the NN method to low-frequency studies, we will only briefly mention some of the higher-frequency applications. As the problem of short data records is no longer a major obstacle in higher-frequency applications, the NN method has been successfully used in areas such as satellite imagery analysis and ocean acoustics (Lee et al. 1990; Badran et al. 1991; IEEE 1991; French et al. 1992; Peak and Tag 1992; Bankert 1994; Peak and Tag 1994; Stogryn et al. 1994; Krasnopolsky et al. 1995; Butler et al. 1996; Marzban and Stumpf 1996; Liu et al. 1997).

3. Neural network models

To keep within the scope of this paper, we will limit our survey of NN models to the feed-forward neural network. Figure 1 shows a network with one hidden layer, where the $j$th neuron in this hidden layer is assigned the value $y_j$, given in terms of the input values $x_i$ by

$$y_j = \tanh \left( \sum_i w_{ij} x_i + b_j \right),$$

where $w_{ij}$ and $b_j$ are the weight and bias parameters, respectively, and the hyperbolic tangent function is used as the activation function (Fig. 3). Other functions besides the hyperbolic tangent could be used for the activation function, which was designed originally to simulate the firing or nonfiring of a neuron upon receiving input signals from its neighbors. If there are additional hidden layers, then equations of the same form as (1) will be used to calculate the values of the next layer of hidden neurons from the current layer of neurons. The output neurons $z_k$ are usually calculated by a linear combination of the neurons in the layer just before the output layer, that is,

$$z_k = \sum_j \tilde{w}_{jk} y_j + \tilde{b}_k.$$  

(2)

To construct a NN model for forecasting, the predictor variables are used as the input, and the predictands (either the same variables or other variables at some lead time) as the output. With $z_{dk}$ denoting the observed data, the NN is trained by finding the optimal values of the weight and bias parameters ($w_{ij}$, $\tilde{w}_{jk}$, $b_j$, and $\tilde{b}_j$), which will minimize the cost function:

$$J = \sum (z_k - z_{dk})^2,$$

(3)

where the rhs of the equation is simply the sum-squared error of the output. The optimal parameters can be found by a back-propagation algorithm (Rumelhart et al. 1986; Hertz et al. 1991). For the reader familiar with variational data assimilation methods, we would point out that this back propagation is equivalent to the backward integration of the adjoint equations in variational assimilation (see section 9).

The back-propagation algorithm has now been superseded by more efficient optimization algorithms,
such as the conjugate gradient method, simulated annealing, and genetic algorithms (Hertz et al. 1991). Once the optimal parameters are found, the training is finished, and the network is ready to perform forecasting with new input data. Normally the data record is divided into two, with the first piece used for network training and the second for testing forecasts. Due to the large number of parameters and the great flexibility of the NN, the model output may fit the data very well during the training period yet produce poor forecasts during the test period. This results from overfitting; that is, the NN fitted the training data so well that it fitted to the noise, which of course resulted in the poor forecasts over the test period (Fig. 4). An NN model is usually capable of learning the signals in the data, but as training progresses, it often starts learning the noise in the data; that is, the forecast error of the model over the test period first decreases and then increases as the model starts to learn the noise in the training data. Overfitting is often a serious problem with NN models, and we will discuss some solutions in the next section.

Consider the special case of a NN with no hidden layers—inputs being several values of a time series, output being the prediction for the next value, and the input and output layers connected by linear activation functions. Training this simple network is then equivalent to determining an AR model through least squares regression, with the weights of the NN corresponding to the weights of the AR model. Hence the NN model reduces to the well-known AR model in this limit.

In general, most NN applications have only one or two hidden layers, since it is known that to approximate a set of reasonable functions \( f_j({x_j}) \) to a given accuracy, at most two hidden layers are needed (Hertz et al. 1991, 142). Furthermore, to approximate continuous functions, only one hidden layer is enough (Cybenko 1989; Hornik et al. 1989). In our models for forecasting the tropical Pacific (Tangang et al. 1997; Tangang et al. 1998a; Tangang et al. 1998b; Tang et al. 1998, manuscript submitted to J. Climate, hereafter THMT), we have not used more than one hidden layer. There is also an interesting distinction between the nonlinear modeling capability of NN models and that of polynomial expansions. With only a few parameters, the polynomial expansion is only capable of learning low-order interactions. In contrast, even a small NN is fully nonlinear and is not limited to learning low-order interactions. Of course, a small NN can learn only a few interactions, while a bigger one can learn more.

When forecasting at longer lead times, there are two possible approaches. The iterated forecasting approach trains the network to forecast one time step forward, then use the forecasted result as input to the same network for the next time step, and this process is iterated until a forecast for the \( n \)th time step is obtained. The other is the direct or “jump” forecast approach, where a network is trained for forecasting at a lead time of \( n \) time steps, with a different network for each value of \( n \). For deterministic chaotic systems, iterated forecasts seem to be better than direct forecasts (Gershenfeld and Weigend 1994). For noisy time series, it is not clear which approach is better. Our experience with climate data suggested that the direct forecast approach is better. Even with “clearning” and continuity constraints (Tang et al. 1996), the iterated forecast method was unable to prevent the forecast errors from being amplified during the iterations.

4. Nonlinear instability

Let us examine obstacle a; that is, the application of NNs to relatively short data records leads to an ill-conditioned problem. In this case, as the cost function is full of deep local minima, the optimal search would likely end up trapped in one of the local minima (Fig. 2). The situation gets worse when we either (i) make the NN more nonlinear (by increasing the number of neurons, hence the number of parameters), or
(ii) shorten the data record. With a different initialization of the parameters, the search would usually end up in a different local minimum.

A simple way to deal with the local minima problem is to use an ensemble of NNs, where their parameters are randomly initialized before training. The individual NN solutions would be scattered around the global minimum, but by averaging the solutions from the ensemble members, we would likely obtain a better estimate of the true solution. Tangang et al. (1998a) found the ensemble approach useful in their forecasts of tropical Pacific sea surface temperature anomalies (SSTA).

An alternative ensemble technique is “bagging” (abbreviated from bootstrap aggregating) (Breiman 1996; THMT). First, training pairs, consisting of the predictor data available at the time of a forecast and the forecast target data at some lead time, are formed. The available training pairs are separated into a training set and a forecast test set, where the latter is reserved for testing the model forecasts only and is not used for training the model. The training set is used to generate an ensemble of NN models, where each member of the ensemble is trained by a subset of the training set. The subset is drawn at random with replacement from the training set. The subset has the same number of training pairs as the training set, where some pairs in the training set appear more than once in the subset, and about 37% of the training pairs in the training set are unused in the subset. These unused pairs are not wasted, as they are used to determine when to stop training. To avoid overfitting, THMT stopped training when the error variance from applying the model to the set of “unused” pairs started to increase. By averaging the output from all the individual members of the ensemble, a final output is obtained.

We found ensemble averaging to be most effective in preventing nonlinear instability and overfitting. However, if the individual ensemble members are severely overfitting, the effectiveness of ensemble averaging is reduced. There are several ways to prevent overfitting in the individual NN models of the ensemble, as presented below.

“Stopped training” as used in THMT is a type of nonconvergent method, which was initially viewed with much skepticism as the cost function does not in general converge to the global minimum. Under the enormous weight of empirical evidence, theorists have finally begun to rigorously study the properties of stopped training (Finnoff et al. 1993). Intuitively, it is not difficult to see why the global minimum is often not the best solution. When using linear methods, there is little danger of overfitting (provided one is not using too many parameters); hence, the global minimum is the best solution. But with powerful nonlinear methods, the global minimum would be a very close fit to the data (including the noise), like the solid curve in Fig. 4. Stopped training, in contrast, would only have time to converge to the dashed curve in Fig. 4, which is actually a better solution.

Another approach to prevent overfitting is to penalize the excessive parameters in the NN model. The ridge regression method (Chauvin, 1990; Tang et al. 1994; Tangang et al. 1998a, their appendix) modifies the cost function in (3) by adding weight penalty terms, that is,

$$J = \sum \left( z_k - z_{dk} \right)^2 + c_1 \sum w_{ij}^2 + c_2 \sum w_{jk}^2,$$

with $c_1, c_2$ positive constants, thereby forcing unimportant weights to zero.

Another alternative is network pruning, where insignificant weights are removed. Such methods have oxymoronic names like “optimal brain damage” (Le Cun et al. 1990). With appropriate use of penalty and pruning methods, the global minimum solution may be able to avoid overfitting. A comparison of the effectiveness of nonconvergent methods, penalty methods, and pruning methods is given by Finnoff et al. (1993).

In summary, the nonlinearity in the NN model introduces two problems: (i) the presence of local minima in the cost function, and (ii) overfitting. Let $D$ be the number of data points and $P$ the number of parameters in the model. For many applications of NN in robotics and pattern recognition, there is almost an unlimited amount of data, so that $D >> P$, whereas in contrast, $D \sim P$ in low-frequency meteorological–oceanographic studies. The local minima problem is present when $D >> P$, as well as when $D \sim P$. However, overfitting tends to occur when $D \sim P$ but not when $D >> P$. Ensemble averaging has been found to be effective in helping the NNs to cope with local minima and overfitting problems.

5. Prefiltering the data fields

Let us now examine obstacle b, that is, the spatially large data fields. Clearly if data from each spatial grid
is to be an input neuron, the NN will have so many parameters that the problem becomes ill conditioned. We need to effectively compress the input (and output) data fields by a prefiltering process. PCA analysis is widely used to compress large data fields (Preisendorfer 1988).

In the PCA representation, we have

\[ x_i(t) = \sum_n a_n(t)e_n, \tag{5} \]

and

\[ a_n(t) = \sum_i x_i(t)e_n, \tag{6} \]

where \( e_n \) and \( a_n \) are the \( n \)th mode PCA and its time coefficient, respectively, with the PCAs forming an orthonormal basis. PCA maximizes the variance of \( a_i(t) \) and then, from the residual, maximizes the variance of \( a_j(t) \), and so forth for the higher modes, under the constraint of orthogonality for the \( \{ e_n \} \). If the original data \( x_i(i = 1, \ldots, I) \) imply a very large number of input neurons, we can usually capture the main variance of the input data and filter out noise by using the first few PCA time series \( a_n(n = 1, \ldots, N) \), with \( N \ll I \), thereby greatly reducing the number of input neurons and the size of the NN model.

In practice, we would need to provide information on how the \( \{ a_n \} \) have been evolving in time prior to making our forecast. This is usually achieved by providing \( \{ a_n(t) \}, \{ a_n(t-\Delta t) \}, \ldots, \{ a_n(t-m\Delta t) \} \), that is, some of the earlier values of \( \{ a_n \} \), as input. Compression of input data in the time dimension is possible with the extended PCA or EOF (EPCA or EEOF) method (Weare and Nasstrom 1982; Graham et al. 1987).

In the EPCA analysis, copies of the original data matrix \( \mathbf{X}_i = x_j(t) \) are stacked with time lag \( \tau \) into a larger matrix \( \mathbf{X}_j \),

\[ \mathbf{X}^T = (\mathbf{X}_j^T, \mathbf{X}_{j+\Delta t}^T, \ldots, \mathbf{X}_{j+n\Delta t}^T), \tag{7} \]

where the superscript \( T \) denotes the transpose and \( \mathbf{X}_{j+n\Delta t} = x(t_j + n\Delta t) \). Applying the standard PCA analysis to \( \mathbf{X}^T \) yields the EPCAs, with the corresponding time series \( \{ a'_n \} \), which because time-lag information has already been incorporated into the EPCAs, could be used instead of \( \{ a_n(t) \}, \{ a_n(t-\Delta t) \}, \ldots, \{ a_n(t-m\Delta t) \} \) from the PCA analysis, thereby drastically reducing the number of input neurons.

This reduction in the input neurons by EPCAs comes with a price, namely the time-domain filtering automatically associated with the EPCA analysis, which results in some loss of input information. Monahan et al. (1998, manuscript submitted to Atmos.–Ocean) found that the EPCAs could become degenerate if the lag \( \tau \) was close to the integral timescale of standing waves in the data. While there are clearly trade-offs between using PCAs and using EPCAs, our experiments with forecasting the tropical Pacific SST found that the much smaller networks resulting from the use of EPCAs tended to be less prone to overfitting than the networks using PCAs.

We are also studying other possible prefiltering processes. Since CCA often uses EPCA to prefilter the predictor and predictand fields (Barnston and Ropelewsuki 1992), we are investigating the possibility of using the NN model with the CCA as a prefilter, that is, using the CCA modes instead of the PCA or EPCA modes. Alternatively, we may argue that all these prefilters are linear processes, whereas we should use a nonlinear prefilter for a nonlinear method such as NN. We are investigating the possibility of using NN models as nonlinear PCAs to do the prefiltering (section 8).

6. Interpreting the NN model

We now turn to obstacle \( c \), namely, the great difficulty in understanding the nonlinear NN model results. In particular, is there a meaningful interpretation of those mysterious neurons in the hidden layer?

Consider a simple NN for forecasting the tropical Pacific wind stress field. The input consists of the first 11 EPCA time series of the wind stress field (from The Florida State University), plus a sine and cosine function to indicate the phase with respect to the annual cycle. The single hidden layer has three neurons, and the output layer the same 11 EPCA time series one month later. As the values of the three hidden neurons can be plotted in 3D space, Fig. 5 shows their trajectory for selected years. From Fig. 5 and the trajectories of other years (not shown), we can identify regions in the 3D phase space as the El Niño warm event phase and its precursor phase and the cold event and its precursor. One can issue warm event or cold event forecasts whenever the system enters the warm event precursor region or the cold event precursor region, respectively. Thus the hidden layer spans a 3D phase space for the El Niño–Southern Oscillation (ENSO) system and is, thus, a higher-dimension generalization.
of the 2D phase space based on the POP method (Tang
1995).

Hence we interpret the NN as a projection from the
input space onto a phase space, as spanned by the neu-
rons in the hidden layer. The state of the system in the
phase space then allows a projection onto the output
space, which can be the same input variables some
time in the future, or other variables in the future. This
interpretation provides a guide for choosing the appro-
priate number of neurons in the hidden layer; namely,
the number of hidden neurons should be the same as
the embedding manifold for the system. Since the
ENSO system is thought to have only a few degrees
of freedom (Grieger and Latif 1994), we have limited
the number of hidden neurons to about 3–7 in most of
our NN forecasts for the tropical Pacific, in contrast
to the earlier study by Derr and Slutz (1994), where
30 hidden neurons were used. Our approach for gen-
erating a phase space for the ENSO attractor with an
NN model is very different from that of Grieger and
Latif (1994), since their phase space was generated
from the model output instead of from the hidden
layer. Their approach generated a 4D phase space from
using four PCA time series as input and as output,
whereas our approach did not have to limit the num-
ber of input or output neurons to generate a low-
dimensional phase space for the ENSO system.

In many situations, the NN model is compared with a linear
model, and one wants to know how nonlinear the NN model
really is. A useful diagnostic tool to measure nonlinearity is
spectral analysis (Tangang et al. 1998b). Once a network has
been trained, we replace the N
input signals by artificial sinu-
soidal signals with frequencies
ω,
ω,
..., ω,
which were care-
fully chosen so that the nonlin-
ear interactions of two signals of
frequencies ω,
and ω,
would
generate frequencies ω + ω
and |
ω − ω |
not equal to any of
the original input frequencies.
The amplitude of a sinusoidal
signal is chosen so as to yield
the same variance as that of the
original real input data. The
output from the NN with the
sinusoidal inputs is spectrally
analyzed (Fig. 6). If the NN is basically linear, spec-
tral peaks will be found only at the original input fre-
quencies. The presence of an unexpected peak at
frequency ω′, equaling ω + ω
or |
ω − ω |
for some
i, j, indicates a nonlinear interaction between the i
and the jth predictor time series. For an overall mea-
sure of the degree of nonlinearity of an NN, we can
calculate the total area under the output spectrum, ex-
cluding the contribution from the original input fre-
quencies, and divide it by the total area under the
spectrum, thereby yielding an estimate of the portion
of output variance that is due to nonlinear interactions.
Using this measure of nonlinearity while forecasting
the regional SSTA in the equatorial Pacific, Tangang
et al. (1998b) found that the nonlinearity of the NN
tended to vary with forecast lead time and with geo-
graphical location.

7. Forecasting the tropical Pacific SST

Many dynamical and statistical methods have been
applied to forecasting the ENSO system (Barnston
et al. 1994). Tangang et al. (1997) forecasted the SSTA
in the Niño 3.4 region (Fig. 7) with NN models using
several PCA time series from the tropical Pacific wind
stress field as predictors. Tangang et al. (1998a) com-
pared the relative merit of using the sea level pressure (SLP) as predictor versus the wind stress as predictor, with the SLP emerging as the better predictor, especially at longer lead times. The reason may be that the SLP is less noisy than the wind stress; for example, the first seven modes of the tropical Pacific SLP accounted for 81% of the total variance, versus only 54% for the seven wind stress modes. In addition, Tangang et al. (1998a) introduced the approach of ensemble averaging NN forecasts with randomly initialized weights to give better forecasts. The best forecast skills were found over the western-central and central equatorial regions (Niño 4 and 3.4) with lesser skills over the eastern regions (Niño 3, P4, and P5) (Fig. 7).

To further simplify the NN models, Tangang et al. (1998b) used EPCAs instead of PCAs and a simple pruning method. The spectral analysis method was also introduced to interpret the nonlinear interactions in the NN, showing that the nonlinearity of the networks tended to increase with lead time and to become stronger for the eastern regions of the equatorial Pacific Ocean.

THMT compare the correlation skills of NN, CCA, and multiple linear regression (LR) in forecasting the Niño 3 SSTA index. First the PCAs of the tropical Pacific SSTA field and the SLP anomaly field were calculated. The same predictors were chosen for all three methods, namely the first seven SLP anomaly PCA time series at the initial month, and 3 months, 6 months, and 9 months before the initial month (a total of 7 × 4 = 28 predictors); the first 10 SSTA PCA time series at the initial month; and the Niño 3 SSTA at the initial month. These 39 predictors were then further compressed to 12 predictors by an EPCA, and cross-validated model forecasts were made at various lead times after the initial month. Ten CCA modes were used in the CCA model, as fewer CCA modes degraded the forecasts. Figure 8 shows that the NN has better forecast correlation skills than CCA and LR at all lead times, especially at the 12-month lead time (where NN has a correlation skill of 0.54 vs 0.49 for CCA). Figure 9 shows the cross-validated forecasts of the Niño 3 SSTA at 6-month lead time (THMT). In other tropical regions, the advantage of NN over CCA was smaller, and in Niño 4, the CCA slightly outperformed the NN model. In this comparison, CCA had an advantage over the NN and LR: While the predictand for the NN and LR was Niño 3 SSTA, the predictands for the CCA were actually the first 10 PCA modes of the tropical Pacific SSTA field, from which the regional Niño 3 SSTA was then calculated. Hence during training, the CCA had more information available by using a much broader predictand field than the NN and LR models, which did not have predictand information outside Niño 3.

As the extratropical ENSO variability is much more nonlinear than in the Tropics (Hoerling et al. 1997), it is possible that the performance gap between the nonlinear NN and the linear CCA may widen for forecasts outside the Tropics.
8. Nonlinear principal component analysis

PCA is popular because it offers the most efficient linear method for reducing the dimensions of a dataset and extracting the main features. If we are not restricted to using only linear transformations, even more powerful data compression and extraction is generally possible. The NN offers a way to do nonlinear PCA (NLPCA) (Kramer 1991; Bishop 1995).

In NLPCA, the NN outputs are the same as the inputs, and data compression is achieved by having relatively few hidden neurons forming a “bottleneck” layer (Fig. 10). Since there are few bottleneck neurons, it would in general not be possible to reproduce the inputs exactly by the output neurons. How many hidden layers would such an NN require in order to perform NLPCA? At first, one might think only one hidden layer would be enough. Indeed with one hidden layer and linear activation functions, the NN solution should be identical to the PCA solution. However, even with nonlinear activation functions, the NN solution is still basically that of the linear PCA solution (Bourlard and Kamp 1988). It turns out that for NLPCA, three hidden layers are needed (Fig. 10) (Kramer 1991).

The reason is that to properly model nonlinear continuous functions, we need at least one hidden layer between the input layer and the bottleneck layer, and another hidden layer between the bottleneck layer and the output layer (Cybenko 1989). Hence, a nonlinear function maps from the higher-dimension input space to the lower-dimension space represented by the bottleneck layer, and then an inverse transform maps from the bottleneck space back to the original higher-dimensional space represented by the output layer, with the requirement that the output be as close to the input as possible. One approach chooses to have only one neuron in the bottleneck layer, which will extract a single NLPCA mode. To extract higher modes, this first mode is subtracted from the original data, and the procedure is repeated to extract the next NLPCA mode.

Using the NN in Fig. 10, A. H. Monahan (1998, personal communication) extracted the first NLPCA mode for data from the Lorenz (1963) three-component chaotic system. Figure 11 shows the famous Lorenz attractor for a scatterplot of data in the \( x - z \) plane. The first PCA mode is simply a horizontal line, explaining 60% of the total variance, while the first NLPCA is the U-shaped curve, explaining 73% of the variance. In general, PCA models data with lines, planes, and hyperplanes for higher dimensions, while the NLPCA uses curves and curved surfaces.

Malthouse (1998) pointed out a limitation of the Kramer (1991) NLPCA method, with its three hidden layers. When the curve from the NLPCA crosses itself, for example forming a circle, the method fails. The reason is that with only one hidden layer between the input and the bottleneck layer, and again one hidden layer between the bottleneck and the output, the non-
linear mapping functions are limited to continuous functions (Cybenko 1989), which of course cannot identify 0° with 360°. However, we believe this failure can be corrected by having two hidden layers between the input and bottleneck layers, and two hidden layers between the bottleneck and output layers, since any reasonable function can be modeled with two hidden layers (Hertz et al. 1991, 142). As the PCA is a cornerstone in modern meteorology–oceanography, a nonlinear generalization of the PCA method by NN is indeed exciting.

9. Neural networks and variational data assimilation

Variational data assimilation (Daley 1991) arose from the need to use data to guide numerical models [including coupled atmosphere–ocean models (Lu and Hsieh 1997, 1998a, 1998b)], whereas neural network models arose from the desire to model the vast empirical learning capability of the brain. With such diverse origins, it is no surprise that these two methods have evolved to prominence completely independently. Yet from section 3, the minimization of the cost function (3) by adjusting the parameters of the NN model is exactly what is done in variational (adjoint) data assimilation, except that here the governing equations are the neural network equations, (1) and (2), instead of the dynamical equations (see appendix for details).

Functionally, as an empirical modeling technique, NN models appear closely related to the familiar linear empirical methods, such as CCA, SVD, PCA, and POP, which belong to the class of singular value or eigenvalue methods. This apparent similarity is somewhat misleading, as structurally, the NN model is a variational data assimilation method. An analogy would be the dolphin, which lives in the sea like a fish, but is in fact a highly evolved mammal, hence the natural bond with humans. Similarly, the fact that the NN model is a variational data assimilation method allows it to be bonded naturally to a dynamical model under a variational assimilation formulation. The dynamical model equations can be placed on equal footing with the NN model equations, with both the dynamical model parameters and initial conditions and the NN parameters found by minimizing a single cost function. This integrated treatment of the empirical and dynamical parts is very different from present forecast systems such as Model Output Statistics (Wilks 1995; Vislocky and Fritsch 1997), where the dynamical model is run first before the statistical method is applied.

What good would such a union bring? Our present dynamical models have good forecast skills for some variables and poor skills for others (e.g., precipitation, snowfall, ozone concentration, etc.). Yet there may be sufficient data available for these difficult variables that empirical methods such as NN may be useful in improving their forecast skills. Also, hybrid coupled models are already being used for El Niño forecasting (Barnett et al. 1993), where a dynamical ocean model is coupled to an empirical atmospheric model. A combined neural–dynamical approach may allow the NN to complement the dynamical model, leading to an improvement of modeling and prediction skills.

10. Conclusions

The introduction of empirical or statistical methods into meteorology and oceanography has been broadly classified as having occurred in four distinct stages: 1) linear regression (and correlation analysis), 2) PCA analysis, 3) CCA (and SVD), and 4) NN. These four stages correspond respectively to the evolving needs of finding 1) a linear relation (or correlation) between two variables x and z; 2) the correlated patterns within a set of variables $x_1, \ldots, x_r$; 3) the linear relations between a set of variables $x_1, \ldots, x_n$ and another set of variables $z_1, \ldots, z_m$; and 4) the nonlinear
relations between $x_1, \ldots, x_n$ and $z_1, \ldots, z_m$. Without exception, in all four stages, the method was first invented in a biological–psychological field, long before its adaptation by meteorologists and oceanographers many years or decades later.

Despite the great popularity of the NN models in many fields, there are three obstacles in adapting the NN method to meteorology–oceanography: (a) nonlinear instability, especially with a short data record; (b) overwhelmingly large spatial data fields; and (c) difficulties in interpreting the nonlinear NN results. In large-scale, low-frequency studies, where data records are in general short, the potential success of the unstable nonlinear NN model against stable linear models, such as the CCA, depends critically on our ability to tame nonlinear instability.

Recent research shows that all three obstacles can be overcome. For obstacle a, ensemble averaging is found to be effective in controlling nonlinear instability. Penalty and pruning methods and nonconvergent methods also help. For b, the PCA method is found to be an effective prefilter for greatly reducing the dimension of the large spatial data fields. Other possible prefilters include the EPCA, rotated PCA, CCA, and nonlinear PCA by NN models. For c, the mysterious hidden layer can be given a phase space interpretation, and a spectral analysis method aids in understanding the nonlinear NN relations. With these and future improvements, the nonlinear NN method is evolving to a versatile and powerful technique capable of augmenting traditional linear statistical methods in data analysis and in forecasting. The NN model is a type of variational (adjoint) data assimilation, which further allows it to be linked to dynamical models under adjoint data assimilation, potentially leading to a new class of hybrid neural–dynamical models.

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Appendix: Connecting neural networks with variational data assimilation

With a more compact notation, the NN model in section 3 can be simply written as

$$ z = N(x, q), \quad (A1) $$

Fig. 11. Data from the Lorenz (1963) three-component ($x, y, z$) chaotic system were used to perform PCA and NLPCA (using the NN model shown in Fig. 10), with the results displayed as a scatterplot in the $x$–$z$ plane. The horizontal line is the first PCA mode, while the curve is the first NLPCA mode (A. H. Monahan 1998, personal communication).
where $\mathbf{x}$ and $\mathbf{z}$ are the input and output vectors, respectively, and $\mathbf{q}$ is the parameter vector containing $w_{ij}^d$, $\tilde{w}_{ij}^d$, $b_j$, and $\tilde{b}_c$. A Lagrange function, $L$, can be introduced,

$$L = J + \sum \mu \star (\mathbf{z} - \mathbf{N}), \quad (A2)$$

where the model constraint (A1) is incorporated in the optimization with the help of the Lagrange multipliers (or adjoint variables) $\mu$. The NN model has now been cast in a standard adjoint data assimilation formulation, where the Lagrange function $L$ is to be optimized by finding the optimal control parameters $\mathbf{q}$, via the solution of the adjoint equations for the adjoint variables $\mu$ (Daley 1991). Thus, the NN jargon “back-propagation” is simply “backward integration of the adjoint model” in adjoint data assimilation jargon.

Often, the predictands are the same variables as the predictors, only at a different time; that is, (A1) and (A2) become

$$\mathbf{z}(t + \Delta t) = \mathbf{N}(\mathbf{z}_d(t), \mathbf{q}), \quad (A3)$$

$$L = J + \sum \mu(t) \star [\mathbf{z}(t + \Delta t) - \mathbf{N}(\mathbf{z}_d(t), \mathbf{q})], \quad (A4)$$

where $\Delta t$ is the forecast lead time. For notational simplicity, we have ignored some details in (A3) and (A4): for example, the forecast made at time $t$ could use not only $\mathbf{z}_d(t)$, but also earlier data; and $\mathbf{N}$ may also depend on other predictor or forcing data $\mathbf{x}_d$.

There is one subtle difference between the feedforward NN model (A4) and standard adjoint data assimilation—the NN model starts with the data $\mathbf{z}_d$ at every time step, whereas the dynamical model takes the data as initial condition only at the first step. For subsequent steps, the dynamical model takes the model output of the previous step and integrates forward. So in adjoint data assimilation with dynamic models, $\mathbf{z}_d(t)$ in Eq. (A4) is replaced by $\mathbf{z}(t)$, that is,

$$L = J + \sum \mu(t) \star [\mathbf{z}(t + \Delta t) - \mathbf{N}(\mathbf{z}(t), \mathbf{q})], \quad (A5)$$

where only at the initial time $t = 0$ is $\mathbf{z}(0) = \mathbf{z}_d(0)$. We can think of this as a strong constraint of continuity (Daley 1991), since it imposes that during the data assimilation period [0,$T$], the solution has to be continuous. In contrast, the training scheme of the NN has no constraint of continuity.

However, the NN training scheme does not have to be without a continuity constraint. Tang et al. (1996) proposed a new NN training scheme, where the Lagrange function is

$$L = J + \alpha \sum [\tilde{\mathbf{z}}(t) - \mathbf{z}_d(t)]^2 + \beta \sum [\tilde{\mathbf{z}}(t) - \mathbf{z}(t)]^2 + \sum \mu(t) \star [\mathbf{z}(t + \Delta t) - \mathbf{N}(\tilde{\mathbf{z}}(t), \mathbf{q})]. \quad (A6)$$

where, $\tilde{\mathbf{z}}$, the input to the NN model, is also adjusted in the optimization process, along with the model parameter vector $\mathbf{q}$. The second term on the right-hand side of (A6), the relative importance of which is controlled by the coefficient $\alpha$, is a constraint to force the adjustable inputs to be close to the data. The third term, whose relative importance is controlled by the coefficient $\beta$, is a constraint to force the inputs to be close to the outputs of the previous step. However, this term does not dictate that the inputs have to be the outputs of the previous step. It is thus a weak constraint of continuity (Daley 1991). Note that $\alpha$ and $\beta$ are scalar constants, whereas $\mu(t)$ is a vector of adjoint variables. The weak continuity constraint version (A6) can thus be thought of as the middle ground between the strong continuity constraint version (A5) and no continuity constraint version (A4).

Let us now couple an NN model to a dynamical model under an adjoint assimilation formulation. This kind of a hybrid model may benefit a system where some variables are better simulated by a dynamical model, while other variables are better simulated by an NN model.

Suppose we have a dynamical model with governing equations in discrete form,

$$\mathbf{u}(t + \Delta t) = \mathbf{M}(\mathbf{u}, \mathbf{v}, \mathbf{p}, t), \quad (A7)$$

where $\mathbf{u}$ denotes the vector of state variables in the dynamical model, $\mathbf{v}$ denotes the vector of variables not modeled by the dynamical model, and $\mathbf{p}$ denotes a vector of model parameters and/or initial conditions. Suppose the $\mathbf{v}$ variables, which could not be forecasted well by a dynamical model, could be forecasted with better skills by an NN model, that is,

$$\mathbf{v}(t + \Delta t) = \mathbf{N}(\mathbf{u}, \mathbf{v}, \mathbf{q}, t), \quad (A8)$$

where the NN model $\mathbf{N}$ has inputs $\mathbf{u}$ and $\mathbf{v}$, and parameters $\mathbf{q}$.

If observed data $\mathbf{u}_d$ and $\mathbf{v}_d$ are available, then we can define a cost function,
where the superscript $T$ denotes the transpose and $\mathbf{U}$ and $\mathbf{V}$ the weighting matrices, often computed from the inverses of the covariance matrices of the observational errors. For simplicity, we have omitted the observational operator matrices and terms representing a priori estimates of the parameters, both commonly used in actual adjoint data assimilation. The Lagrange function $L$ is given by

$$
L = J + \sum \lambda(t)^T \left[ \mathbf{u}(t + \delta t) - \mathbf{M} \right] \\
+ \sum \mu(t)^T \left[ \mathbf{v}(t + \Delta t) - \mathbf{N} \right],
$$

(A10)

where $\lambda$ and $\mu$ are the vectors of adjoint variables. This formulation places the dynamical model $\mathbf{M}$ and the neural network model $\mathbf{N}$ on equal footing, as both are optimized (i.e., optimal values of $\mathbf{p}$ and $\mathbf{q}$ are found) by minimizing the Lagrange function $L$—that is, the adjoint equations for $\lambda$ and $\mu$ are obtained from the variation of $L$ with respect to $\mathbf{u}$ and $\mathbf{v}$, while the gradients of the cost function with respect to $\mathbf{p}$ and $\mathbf{q}$ are found from the variation of $L$ with respect to $\mathbf{p}$ and $\mathbf{q}$. Note that without $\mathbf{v}$ and $\mathbf{N}$, Eqs. (A7), (A9), and (A10) simply reduce to the standard adjoint assimilation problem for a dynamical model; whereas without $\mathbf{u}$ and $\mathbf{M}$, Eqs. (A8), (A9), and (A10) simply reduce to finding the optimal parameters $\mathbf{q}$ for the NN model.

Here, the hybrid neural-dynamical data assimilation model (A10) is in a strong continuity constraint form. Similar hybrid models can be derived for a weak continuity constraint or no continuity constraint.

References


