Backpropagation Through Time: What It Does and How to Do It

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Backpropagation is now the most widely used tool in the field of artificial neural networks. At the core of backpropagation is a method for calculating derivatives exactly and efficiently in any large system made up of elementary subsystems or calculations which are represented by known, differentiable functions; thus, backpropagation has many applications which do not involve neural networks as such.

This paper first reviews basic backpropagation, a simple method which is now being widely used in areas like pattern recognition and fault diagnosis. Next, it presents the basic equations for backpropagation through time, and discusses applications to areas like pattern recognition involving dynamic systems, systems identification, and control. Finally, it describes further extensions of this method, to deal with systems other than neural networks, systems involving simultaneous equations or true recurrent networks, and other practical issues which arise with this method. Pseudocode is provided to clarify the algorithms. The chain rule for ordered derivatives—the theorem which underlies backpropagation—is briefly discussed.

I. INTRODUCTION

Backpropagation through time is a very powerful tool, with applications to pattern recognition, dynamic modeling, sensitivity analysis, and the control of systems over time, among others. It can be applied to neural networks, to econometric models, to fuzzy logic structures, to fluid dynamics models, and to almost any system built up from elementary subsystems or calculations. The one serious constraint is that the elementary subsystems must be represented by functions known to the user, functions which are both continuous and differentiable (i.e., possess derivatives). For example, the first practical application of backpropagation was for estimating a dynamic model to predict nationalism and social communications in 1974 [1].

Unfortunately, the most general formulation of backpropagation can only be used by those who are willing to work out the mathematics of their particular application. This paper will mainly describe a simpler version of backpropagation, which can be translated into computer code and applied directly by neural network users.

Section II will review the simplest and most widely used form of backpropagation, which may be called “basic backpropagation.” The concepts here will already be familiar to those who have read the paper by Rumelhart, Hinton, and Williams [2] in the seminal book Parallel Distributed Processing, which played a pivotal role in the development of the field. (That book also acknowledged the prior work of Parker [3] and Le Cun [4], and the pivotal role of Charles Smith of the Systems Development Foundation.) This section will use new notation which adds a bit of generality and makes it easier to go on to complex applications in a rigorous manner. (The need for new notation may seem unnecessary to some, but for those who have to apply backpropagation to complex systems, it is essential.)

Section III will use the same notation to describe backpropagation through time. Backpropagation through time has been applied to concrete problems by a number of authors, including, at least, Watrous and Shastri [5], Sawai and Waibel et al. [6], Nguyen and Widrow [7], Jordan [8], Kawato [9], Elman and Zipser, Narendra [10], and myself [1], [11], [12], [15]. Section IV will discuss what is missing in this simplified discussion, and how to do better.

At its core, backpropagation is simply an efficient and exact method for calculating all the derivatives of a single target quantity (such as pattern classification error) with respect to a large set of input quantities (such as the parameters or weights in a classification rule). Backpropagation through time extends this method so that it applies to dynamic systems. This allows one to calculate the derivatives needed when optimizing an iterative analysis procedure, a neural network with memory, or a control system which maximizes performance over time.

II. BASIC BACKPROPAGATION

A. The Supervised Learning Problem

Basic backpropagation is current the most popular method for performing the supervised learning task, which is symbolized in Fig. 1.

In supervised learning, we try to adapt an artificial neural network so that its actual outputs (Y) come close to some target outputs (Y_t) for a training set which contains T patterns. The goal is to adapt the parameters of the network so that it performs well for patterns from outside the training set.

The main use of supervised learning today lies in pattern...
Fig. 1. Schematic of the supervised learning task.

recognition work. For example, suppose that we are trying to build a neural network which can learn to recognize handwritten ZIP codes. (AT&T has actually done this [13], although the details are beyond the scope of this paper.) We assume that we already have a camera and preprocessor which can digitize the image, locate the five digits, and provide a 19 × 20 grid of ones and zeros representing the image of each digit. We want the neural network to input the 19 × 20 image, and output a classification; for example, we might ask the network to output four binary digits which, taken together, identify which decimal digit is being observed.

Before adapting the parameters of the neural network, one must first obtain a training database of actual handwritten digits and correct classifications. Suppose, for example, that this database contains 2000 examples of handwritten digits. In that case, T = 2000. We may give each example a label t between 1 and 2000. For each sample t, we have a record of the input pattern and the correct classification. Each input pattern consists of 380 numbers, which may be viewed as a vector with 380 components; we may call this vector \( X(t) \). The desired classification consists of four numbers, which may be treated as a vector \( Y(t) \). The actual output of the network will be \( \hat{Y}(t) \), which may differ from the desired output \( Y(t) \), especially in the period before the network has been adapted. To solve the supervised learning problem, there are two steps:

- We must specify the "topology" (connections and equations) for a network which inputs \( X(t) \) and outputs a four-component vector \( \hat{Y}(t) \), an approximation to \( Y(t) \). The relation between the inputs and outputs must depend on a set of weights (parameters) \( W \) which can be adjusted.

- We must specify a "learning rule"—a procedure for adjusting the weights \( W \) so as to make the actual outputs \( \hat{Y}(t) \) approximate the desired outputs \( Y(t) \).

Basic backpropagation is currently the most popular learning rule used in supervised learning. It is generally used with a very simple network design—to be described in the next section—but the same approach can be used with any network of differentiable functions, as will be discussed in Section IV.

Even when we use a simple network design, the vectors \( X(t) \) and \( Y(t) \) need not be made of ones and zeros. They can be made up of any values which the network is capable of inputting and outputting. Let us denote the components of \( X(t) \) as \( X_i(t) \) for \( 1 \leq i \leq m \) so that there are \( m \) inputs to the network. Let us denote the components of \( Y(t) \) as \( Y_j(t) \) for \( 1 \leq j \leq n \) so that there are \( n \) outputs. Throughout this paper, the components of a vector will be represented by the same letter as the vector itself, in the same case; this convention turns out to be convenient because \( x(t) \) will represent a different vector, very closely related to \( X(t) \).

Fig. 1 illustrates the supervised learning task in the general case. Given a history of \( X(1) \cdots X(T) \) and \( Y(1) \cdots Y(T) \), we want to find a mapping from \( X \) to \( Y \) which will perform well when we encounter new vectors \( X \) outside the training set. The index "(t)" may be interpreted either as a time index or as a pattern number index; however, this section will not assume that the order of patterns is meaningful.

B. Simple Feedforward Networks

Before we specify a learning rule, we have to define exactly how the outputs of a neural net depend on its inputs and weights. In basic backpropagation, we assume the following logic:

\[
\begin{align*}
x_i &= X_i, \quad 1 \leq i \leq m \\
\text{net}_i &= \sum_{j=1}^{i-1} W_{ij} x_j, \quad m < i \leq N + n \\
x_i &= s(\text{net}_i), \quad m < i \leq N + n \\
Y_i &= x_{i+N}, \quad 1 \leq i \leq n
\end{align*}
\]

where the function \( s \) in (3) is usually the following sigmoidal function:

\[ s(z) = \frac{1}{1 + e^{-z}} \]

and where \( N \) is a constant which can be any integer you choose as long as it is no less than \( m \). The value of \( N + n \) decides how many neurons are in the network (if we include inputs as neurons). Intuitively, \( \text{net}_i \) represents the total level of voltage exciting a neuron, and \( x_i \) represents the intensity of the resulting output from the neuron. (\( x \) is sometimes called the "activation level" of the neuron.) It is conventional to assume that there is a threshold or constant weight \( W_{in} \) added to the right side of (2); however, we can achieve the same effect by assuming that one of the inputs (such as \( x_{in} \)) is always 1.

The significance of these equations is illustrated in Fig. 2. There are \( N + n \) circles, representing all of the neurons in the network, including the input neurons. The first \( m \) circles are really just copies of the inputs \( X_1 \cdots X_m \); they are included as part of the vector \( x \) only as a way of simplifying the notation. Every other neuron in the network—such as neuron number \( n \)—which calculates net, and \( x \)—takes input from every cell which precedes it in the network. Even the last output cell, which generates \( Y_n \), takes input from other output cells, such as the one which outputs \( Y_{n-1} \).

In neural network terminology, this network is "fully connected" in the extreme. As a practical matter, it is usually desirable to limit the connections between neurons. This can be done by simply fixing some of the weights \( W \) to zero so that they drop out of all calculations. For example, most researchers prefer to use "layered" networks, in which all
connection weights $W_{ij}$ are zeroed out, except for those going from one “layer” (subset of neurons) to the next layer. In general, one may zero out as many or as few of the weights as one likes, based on one’s understanding of individual applications. For those who first begin this work, it is conventional to define only three layers—an input layer, a “hidden” layer, and an output layer. This section will assume the full range of allowed connections, simply for the sake of generality.

In computer code, we could represent this network as a Fortran subroutine (assuming a Fortran which distinguishes upper case from lower case):

```fortran
SUBROUTINE NET (X, W, x, Yhat)
REAL X(m), W(N+n,N+n), x(N+n), Yhat(n), net
INTEGER, i, j, m, n, N
C First insert the inputs, as per equation (1)
  DO 1 i = 1, m
1       x(i) = X(i)
C Next implement (2) and (3) together for each value of i
  DO 10 j = 1, N+n
10       net = net + W(i,j)*x(j)
C Finally, calculate the outputs, as per (4)
  DO 2000 i = 1, n
2000     Yhat(i) = 1/(1+exp(-net))

In the pseudocode, note that $X$ and $W$ are technically the inputs to the subroutine, while $x$ and $Y$ are the outputs. $Yhat$ is usually regarded as “the” output of the network, but $x$ may also have its uses outside of the subroutine proper, as will be seen in the next section.

C. Adapting the Network: Approach

In basic backpropagation, we choose the weights $W_{ij}$ so as to minimize square error over the training set:

$$E = \sum_{t=1}^{T} E(t) = \sum_{t=1}^{T} \frac{1}{2} (Y(t) - Y_{\text{TARGET}}(t))^2.$$ (6)

This is simply a special case of the well-known method of least squares, used very often in statistics, econometrics, and engineering; the uniqueness of backpropagation lies in how this expression is minimized. The approach used here is illustrated in Fig. 3.

![Diagram](image)

**Fig. 3.** Basic backpropagation (in pattern learning).

In basic backpropagation, we start with arbitrary values for the weights $W$. (It is usual to choose random numbers in the range from $-0.1$ to $0.1$, but it may be better to guess the weights based on prior information, in cases where prior information is available.) Next, we calculate the outputs $Y(t)$ and the errors $E(t)$ for that set of weights. Then we calculate the derivatives of $E$ with respect to all of the weights; this is indicated by the dotted lines in Fig. 3. If increasing a given weight would lead to more error, we adjust that weight downwards. If increasing a weight leads to less error, we adjust it upwards. After adjusting all the weights up or down, we start all over, and keep on going through this process until the weights and the error settle down. Some researchers iterate until the error is close to zero; however, if the number of training patterns exceeds the number of weights in the network—as recommended by studies on generalization—it may not be possible for the error to reach zero.) The uniqueness of backpropagation lies in the method used to calculate the derivatives exactly for all of the weights in one pass through the system.

D. Calculating Derivatives: Theoretical Background

Many papers on backpropagation suggest that we need only use the conventional chain rule for partial derivatives to calculate the derivatives of $E$ with respect to all of the weights. Under certain conditions, this can be a rigorous approach, but its generality is limited, and it requires great care with the side conditions (which are rarely spelled out); calculations of this sort can easily become confused and erroneous when networks and applications grow complex. Even when using (7) below, it is a good idea to test one’s gradient calculations using explicit perturbations in order to be sure that there is no bug in one’s code.

When the idea of backpropagation was first presented to the Harvard faculty in 1972, they expressed legitimate concern about the validity of the rather complex calculations involved. To deal with this problem, I proved a new chain rule for ordered derivatives:

$$\frac{\partial E}{\partial z_i} = \frac{\partial E}{\partial z_{\text{TARGET}}} + \sum_{j=1}^{p} \frac{\partial E}{\partial z_j} \frac{\partial z_j}{\partial z_i}.$$ (7)

where the derivatives with the superscript represent ordered derivatives, and the derivatives without subscripts represent ordinary partial derivatives. This chain rule is valid only for ordered systems where the values to be calculated can be calculated one by one (if necessary) in the order $z_1$, $z_2$, $z_3$, ... $z_T$, $\text{TARGET}$. The simple partial derivative represents the direct impact of $z_i$ on $z_i$ through the system equations which determines $z_i$. The ordered derivative represents the total impact of $z_i$ on $\text{TARGET}$, accounting for both the direct and indirect effects. For example, suppose that we had a simple system governed by the following two equations, in order:

$$z_2 = 4 \cdot z_1$$
$$z_3 = 3 \cdot z_1 + 5 \cdot z_2.$$

The “simple” partial derivative of $z_2$ with respect to $z_1$ (the direct effect) is 3; to calculate the simple effect, we only look at the equation which determines $z_2$. However, the ordered derivative of $z_2$ with respect to $x_2$ is 23 because of the indirect impact by way of $z_2$. The simple partial derivative measures what happens when we increase $z_1$ (e.g., by 1, in this example) and assume that everything else (like $z_2$) in the equation which determines $z_2$ remains constant. The ordered derivative measures what happens when we increase $z_2$, and also recalculate all other quantities—like
of any ordered derivative simply by perturbing $z$, at the point where we then calculate the derivatives.

For purposes of debugging, one can calculate the true value of any ordered derivative simply by perturbing $z$, at the point in the program where $z$ is calculated; this is particularly useful when applying backpropagation to a complex network.

**E. Adapting the Network: Equations**

For a given set of weights $W$, it is easy to use (1)-(6) to calculate $Y(t)$ and $E(t)$ for each pattern $t$. The trick is in how we then calculate the derivatives.

Let us use the prefix "$F_-$" to indicate the ordered derivative of $F$ with respect to whatever variable the "$F_-$" precedes. Thus, for example,

$$F_{-\dot{Y}}(t) = \frac{\delta F}{\delta \dot{Y}(t)} = \ddot{Y}(t) - Y(t),$$

which follows simply by differentiating (6). By the chain rule for ordered derivatives as expressed in (8),

$$F_{-x_i}(t) = F_{-\dot{Y}}(t) * N_n + \sum_{j=1}^{N_n} W_{j,i} * F_{-y_i}(t),$$

where $s'$ is the derivative of $s(z)$ as defined in (5) and $F_{-y_i}$ is assumed to be zero for $k \leq 0$. Note how (10) requires us to run backwards through the network in order to calculate the derivatives, as illustrated in Fig. 4; this backwards propagation of information is what gives backpropagation its name. A little calculus and algebra, starting from (5), shows us that

$$s'(z) = s(z) * (1 - s(z)), \quad (13)$$

which we can use when we implement (11). Finally, to adapt the weights, the usual method is to set

$$\text{New } W_{ij} = W_{ij} - \text{learning rate} * F_{-W_{ij}} \quad (14)$$

where the learning rate is some small constant chosen on an ad hoc basis. (The usual procedure is to make it as large as possible, up to 1, until the error starts to diverge; however, there are more analytic procedures available.)

**F. Adapting the Network: Code**

The key part of basic backpropagation—(10)-(13)—may be coded up into a "dual" subroutine, as follows.

```plaintext
SUBROUTINE F_NET(F_Yhat, W, x, F_W)
REAL F_Yhat, W(N+N+n, N+n), x(N+n), F_W(N+N+n, F_net(N+n), F_X(n+n)
INTEGER i, j, n, m, N
C Initialize equation (10)
DO 1 i = 1, N
1 F_X(i) = 0
DO 2 i = 1, n
2 F_X(i+N+n) = F_Yhat(i)
C RUN THROUGH (10)-(12) AS A SET,
C FOR I RUNNING BACKWARDS
DO 10 i = N+n, N+1, -1
10 F_W(i, j) = F_net(i)*x(j)
1000 CONTINUE

Note that the array $F_W$ is the only output of this subroutine.

Equation (14) represented "batch learning," in which weights are adjusted only after all $T$ patterns are processed. It is more common to use pattern learning, in which the weights are continually updated after each observation. Pattern learning may be represented as follows:

**C. PATTERN LEARNING**

```plaintext
DO 1000 pass_number = 1, maximum_passes
1000 t = 1, T
CALL NETX(t, W, x, F_Yhat)
C Next Implement equation (9)
DO 9 i = 1, n
9 F_Yhat(i) = Yhat(i) - Y(i)
C Next Implement (10)-(12)
CALL NET(F_Yhat, W, x, F_W)
C Next Implement (14)
C Note how weights are updated within the "DO 100" loop.
```
The key point here is that the weights $W$ are adjusted in response to the current vector $F_w$, which only depends on the current pattern $t$; the weights are adjusted after each pattern is processed. (In batch learning, by contrast, the weights are adjusted only after the “DO 100” loop is completed.)

In practice, maximum_passes is usually set to an enormous number; the loop is exited only when a test of convergence is passed, a test of error size or weight change which can be injected easily into the loop. True real-time convergence through the data and no memory of earlier times is often possible pattern learning or could be real-time learning. Most people using basic backpropagation now use pattern learning rather than real-time learning because, with their data sets, many passes through the data are needed to ensure convergence of the weights.

The reader should be warned that I have not actually tested the code here. It is presented simply as a way of explaining more precisely the preceding ideas. The C implementations which I have worked with have been less transparent, and harder to debug, in part because of the absence of range checking in that language. It is often argued that people “who know what they are doing” do not need range checking and the like; however, people who think they never make mistakes should probably not be writing this kind of code. With neural network code, especially, good diagnostics and tests are very important because bugs can lead to slow convergence and oscillation—problems which are hard to track down, and are easily misattributed to the algorithm in use. If one must use a language without range checking, it is extremely important to maintain a version of the code which is highly transparent and safe, however inefficient it may be, for diagnostic purposes.

III. BACKPROPAGATION THROUGH TIME

A. Background

Backpropagation through time—like basic backpropagation—is used most often in pattern recognition today. Therefore, this section will focus on such applications, using notation like that of the previous section. See Section IV for other applications.

In some applications—such as speech recognition or submarine detection—our classification at time $t$ will be more accurate if we can account for what we saw at earlier times. Even though the training set still fits the same format as above, we want to use a more powerful class of networks to do the classification; we want the output of the network at time $t$ to account for variables at earlier times (as in Fig. 5).
which is programmed just like the subroutine NET, with the modifications one would expect from (15). The output arrays are \( x(0) \) and \( \text{Yhat} \).

When we call this subroutine for the first time, at \( t = 1 \), we face a minor technical problem: there is no value for \( x(-1) \) or \( x(0) \), both of which we need as inputs. In principle, we can use any values we wish to choose; the choice of \( x(-1) \) and \( x(0) \) is essentially part of the definition of our network.

Most people simply set these vectors to zero, and argue that their network will start out with a blank slate in classifying whatever dynamic pattern is at hand, both in the training set and in later applications. (Statisticians have been known to treat these vectors as weights, in effect, to be adapted when one applies the network to new data.)

In this section, I will assume that the data run from an initial time \( t = 1 \) through to a final time \( t = T \), which plays a crucial role in the derivative calculations. Section IV will show how this assumption can be relaxed somewhat.

### C. Adapting the Network: Equations

To calculate the derivatives of \( F \), we use the same equations as before, except that (10) is replaced by

\[
F_x(l) = F_x(l) + \sum_{i=1}^{n} W_{ji} \cdot F_{net}(i) + \sum_{i=m+1}^{n+n} W_{ji} \cdot F_{net}(t+1) + \sum_{i=m+n+1}^{2n+n} W_{ji} \cdot F_{net}(t+2).
\]

\[16\]

Once again, if one wants to fix the \( W \) terms to zero, one can simply delete the rightmost term.

Notice that this equation makes it impossible for us to calculate \( F_x(l) \) and \( F_{net}(l) \) until after \( F_{net}(t+1) \) and \( F_{net}(t+2) \) are already known; therefore, we can only use this equation by proceeding backwards in time, calculating \( F_{net} \) for time \( T \), and then working our way backwards to time \( t \).

To adapt this network, of course, we need to calculate \( F_x(l) \) and \( F_{net}(l) \) as well as \( F_W(l) \):

\[
F_{W}(l) = \sum_{i=1}^{t} F_{net}(t) \cdot x_i \cdot F_x(l) \quad \text{and} \quad F_{W}(l) = \sum_{i=1}^{t} F_{net}(t+1) \cdot x_i.
\]

\[17\]

\[18\]

In all of these calculations, \( F_{net}(t+1) \) and \( F_{net}(t+2) \) should be treated as zero. For programming convenience, I will later define quantities like \( F_{net}(l) = F_{net}(l+1) \), but this is purely a convenience; the subscript "i" and the time argument are enough to identify which derivative is being represented. (In other words, \( F_{net}(l) \) represents a specific quantity \( z_i \) as in (8), and \( F_{net}(l) \) represents the ordered derivative of \( F \) with respect to that quantity.)

### D. Adapting the Network: Code

To fully understand the meaning and implications of these equations, it may help to run through a simple (hypothetical) implementation.

First, to calculate the derivatives, we need a new subroutine, dual to NET2.

```
SUBROUTINE F_NET2(F_Yhat, W, X, F_net, F_W, F_net', F_W', F_W")
REAL F_Yhat(1), W(n+n-1), W"(N+n,N+n)
REAL F_net(N+n), F_net'(N+n), F_net"(N+n)
REAL F_W(N+n,N+n), F_W'(N+n,N+n)

INTEGER i,j,n,m,n

C Initialize equation (16)
DO 1 i = 1, N
  F_X(i) = 0.
DO 2 i = 1, N
  F_X(i) = F_Yhat(i)

C RUN THROUGH (16), (11), AND (12) AS A SET,

C RUNNING BACKWARDS
DO 1000 j = N+n, m+1, -1
  CALL NET2(X(t), W, W", W", X, F_net, F_W, F_net', F_W', F_W")
  REAL F_Yhat(n), W(N+n, N+n), W"(N+n, N+n)
  REAL F_net(n+n), F_net'(n+n), F_net"(n+n)
  REAL F_W(N+n, N+n), F_W'(N+n, N+n)

1000 CONTINUE
```

```c

DO 161 j = i+1, N+n
  F_X(i) = F_X(i) + W(j,i)*F_net(j)
DO 162 j = m+1, N+n
  F_X(i) = F_X(i) + W(j,i)*F_net'(j) + W"(j,i)*F_net"(j)

C next implement (11)
F_net(i) = F_X(i)*x(i)*(1-x(i))

C implement (12), (17), and (18)

(as running sums)
DO 12 j = 1, l-1
  F_W(i) = F_W(i) - F_W(i, j) + F_net(j)*x(j)
DO 1718 j = 1, N+n
  F_W(i) = F_W(i, j) + F_net(j)*x(j)

1718 F_W(i) = F_W(i, j)
1000 CONTINUE
```

Notice that the last two DO loops have been set up to perform running sums, to simplify what follows.
do these calculations in forward time, but exact results
agonation through time, the natural way to adapt the network
weights to zero (or to 1 when we want to force memory) in
order to get the required derivatives. (There are ways to
avoid the calculation of the exponential in (5). Depending
on what kind of processor one has available, this calcula-
tion can multiply run times by a significant factor.
Unfortunately, the second derivatives of this function are not well behaved,
which can affect the efficiency of some applications. Still,
many programmers are now using piecewise linear approxima-
tions to (5), along with lookup tables, which can work
relatively well in some applications. In earlier experiments,
I have also found good uses for a Taylor series approxi-
mation:

\[ s(z) = 1/(1 - z + 0.5 * z^2), \quad z < 0 \]

A. Use of Other Networks

The network shown in (1)-(5) is a very simple, basic net-
work. Backpropagation can be used to adapt a wide variety
of other networks, including networks representing econ-
ometric models, systems of simultaneous equations, etc.
In any case, all of these estimates are

In the neural network field, users are often given a choice of
network “topology.” This simply means that they are
asked to declare which subset of the possible weights con-
nections will actually be used. Every weight removed from
(15) should be removed from (16) as well, along with (12) and
(14) (or whichever apply to that weight); therefore, simpli-
ifying the network by removing weights simplifies all the
other calculations as well. (Mathematically, this is the same
as fixing these weights to zero.) Typically, people will
remove an entire block of weights, such that the limits of the
sums in our equations are all shrunk.

In a truly brain-like network, each neuron in (15)] will
only receive input from a small number of other cells. Ne-
uroscientists do not agree on how many inputs are typical;
some cite numbers on the order of 100 inputs per cell, while
others quote 10,000. In any case, all of these estimates are
small compared to the billions of cells present. To imple-
ment this kind of network efficiently on a conventional
computer, one would use a linked list or a list of offsets to
represent the connections actually implemented for each
cell; the same strategy can be used to implement the back-
wards calculations and keep the connection costs low. Sim-
lar tricks are possible in parallel computers of all types.

Even in the neural network field, many programmers try
to avoid the calculation of the exponential in (5). Depending
on what kind of processor one has available, this calcula-
tion can multiply run times by a significant factor.

This leads to a very simple derivative as well. Unfortunately, the second
derivatives of this function are not well behaved,
which can affect the efficiency of some applications. Still,
many programmers are now using piecewise linear approxi-
mations to (5), along with lookup tables, which can work
relatively well in some applications. In earlier experiments,
I have also found good uses for a Taylor series approxi-
mation:

\[ s(z) = 1/(1 - z + 0.5 * z^2), \quad z < 0 \]
In a similar spirit, it is common to speed up learning by "stretching out" $s(z)$ so that it goes from $-1$ to $1$ instead of $0$ to $1$.

Backpropagation can also be used without using neural networks at all. For example, it can be used to adapt a network consisting entirely of user-specified functions, representing something like an econometric model. In that case, the way one proceeds depends on who one is programming for and what kind of model one has.

If one is programming for oneself and the model consists of a sequence of equations which can be invoked one after the other, then one should consider the tutorial paper [11], which also contains a more rigorous definition of what these "$f_\alpha$" derivatives really mean and a proof of the chain rule for ordered derivatives. If one is developing a tool for others, then one might set it up to look like a standard econometric package (like SAS or TROLL) where the user of the system types in the equations of his or her model; the backpropagation would go inside the package as a way to speed up these calculations, and would mostly be transparent to the user. If one's model consists of a set of simultaneous equations which need to be solved at each time, then one must use more complicated procedures [15]; in neural network terms, one would call this a "doubly recurrent network." (The methods of Pineda [16] and Almeida [17] are special cases of this situation.)

Pearlmutter [18] and Williams [19] have described alternative methods, designed to achieve results similar to those of backpropagation through time, using a different computational strategy. For example, the Williams-Zipser method is a special case of the "conventional perturbation" equation cited in [14], which rejected this as a neural network method on the grounds that its computational costs scale as the square of the network size; however, the method does yield exact derivatives with a time-forward calculation.

Supervised learning problems or forecasting problems which involve memory can also be translated into control problems [15, p. 352], [20], which allows the use of adaptive critic methods, to be discussed in the next section. Normally, this would yield only an approximate solution (or approximate derivatives), but it would also allow time-forward real-time learning. If the network itself contains calculation noise (due to hardware limitations), the adaptive critic approach might even be more robust than backpropagation through time because it is based on mathematics which allow for the presence of noise.

B. Applications Other Than Supervised Learning

Backpropagation through time can also be used in two other major applications: neuroidentification and neurocontrol. (For applications to sensitivity analysis, see [14] and [15].)

In neuroidentification, we try to do with neural nets what econometricians do with forecasting models. (Engineers would call this the identification problem or the problem of identifying dynamic systems. Statisticians refer to it as the problem of estimating stochastic time-series models.)

Our training set consists of vectors $X(t)$ and $u(t)$, not $X(t)$ and $Y(t)$. Usually, $X(t)$ represents a set of observations of the external (sic) world, and $u(t)$ represents a set of actions that we had control over (such as the settings of motors or actuators). The combination of $X(t)$ and $u(t)$ is input to the network at each time $t$. Our target, at time $t$, is the vector $X(t + 1)$.

We could easily build a network to input these inputs, and aim at these targets. We could simply collect the inputs and targets into the format of Section II, and then use basic backpropagation. But basic backpropagation contains no "memory." The forecast of $X(t - 1)$ would depend on $X(t)$, but not on previous time periods. If human beings worked like this, then they would be unable to predict that a ball might roll out the far side of a table after rolling down under the near side; as soon as the ball disappeared from sight [from the current vector $X(t)$], they would have no way of accounting for its existence. (Harold Szu has presented a more interesting example of this same effect: if a tiger chased after such a memoryless person, the person would forget about the tiger after first turning to run away. Natural selection has eliminated such people.) Backpropagation through time permits more powerful networks, which do have a "memory," for use in the same setup.

Even this approach to the neuroidentification problem has its limitations. Like the usual methods of econometrics [15], it may lead to forecasts which hold up poorly over multiple time periods. It does not properly identify where the noise comes from. It does not permit real-time adaptation. In an earlier paper [20], I have described some ideas for overcoming these limitations, but more research is needed. The first phase of Kawato's cascade method [9] for controlling a robot arm is an identification phase, which is more robust over time, and which uses backpropagation through time in a different way; it is a special case of the "pure robust method," which also worked well in the earliest applications which I studied [1], [20].

After we have solved the problem of identifying a dynamic system, we are then ready to move on to controlling that system.

In neurocontrol, we often start out with a model or network which describes the system or plant we are trying to control. Our problem is to adapt a second network, the action network, which inputs $X(t)$ and outputs the control $u(t)$; (In actuality, we can allow the action network to "see" or input the entire vector $x(t)$ calculated by the model network; this allows it to account for memories such as the recent appearance of a tiger.) Usually, we want to adapt the action network so as to maximize some measure of performance or utility $U(X, t)$ summed over time. Performance measures used in past applications have included everything from the energy used to move a robot arm [8], [9] through to net profits received by the gas industry [11]. Typically, we are given a set of possible initial states $x(t)$, and asked to train the action network so as to maximize the sum of utility from time 1 to a final time $T$.

To solve this problem using backpropagation through time, we simply calculate the derivatives of our performance measure with respect to all of the weights in the action network. "Backpropagation" refers to how we calculate the derivatives, not to anything involving pattern recognition or error. We then adapt the weights according to these derivatives, as in (12), except that the sign of the adjustment term is now positive (because we are maximizing rather than minimizing).

The easiest way to implement this approach is to merge
the utility function, the model network, and the action network into one big network. We can then construct the dual to this entire network, as described in 1974 [1] and illustrated in my recent tutorial [11]. However, if we wish to keep the three component networks distinct, then the bookkeeping becomes more complicated. The basic idea is illustrated in Fig. 6, which maps exactly into the approach used by Nguyen and Widrow [7] and by Jordan [8].

Instead of working with a single subroutine, NET, we now need three subroutines:

\[
\text{UTILITY}(x; t; x'^{+}; U) \\
\text{MODEL}(X(t), u(t); x(t); X(t + 1)) \\
\text{ACTION}(x(t); W; x'(t); u(t)).
\]

In each of these subroutines, the two arguments on the right are technically outputs, and the argument on the far right is what we usually think of as the output of the network. We need to know the full vector \( x \) produced inside the model network so that the action network can "see" important memories. The action network does not need to have its own internal memory, but we need to save its internal state \( x' \) so that we can later calculate derivatives. For simplicity, I will assume that MODEL does not contain any lag-two memory terms (i.e., \( W \) weights). The primes after the \( x \)'s indicate that we are looking at the internal states of different networks; they are unrelated to the primes representing lagged values, discussed in Section III, which we will also need in what follows.

To use backpropagation through time, we need to construct dual subroutines for all three of these subroutines:

\[
\text{F-UTILITY}(x'^{+}; t; F.x) \\
\text{F-MODEL}(F.net', F.X', x, F.net, F.u) \\
\text{F-ACTION}(F.u; x'(t); F.W).
\]

The outputs of these subroutines are the arguments on the far right (including \( F.net \)), which are represented by the broken lines in Fig. 4. The subroutine \( \text{F-UTILITY} \) simply reports out the derivatives of \( U(x, t) \) with respect to the variables \( X \). The subroutine \( \text{F-MODEL} \) is like the earlier subroutine \( \text{F-NET2} \), except that we need to output \( F.u \) instead of derivatives to weights. (Again, we are adapting only the action network here.) The subroutine \( \text{F-ACTION} \) is virtually identical to the old subroutine \( F.NET \), except that we need to calculate \( F.W \) as a running sum (as we did in \( F.NET2 \)).

Of these three subroutines, \( \text{F-MODEL} \) is by far the most complex. Therefore, it may help to consider some possible code.

The last small DO loop here assumes that \( u(t) \) was part of the input vector to the original subroutine \( \text{MODEL} \), inserted into the slots between \( x(n + 1) \) and \( x(m) \). Again, a good programmer could easily compress all this; my goal here is only to illustrate the mathematics.

Finally, in order to adapt the action network, we go through multiple passes, each starting from one of the starting values of \( x(1) \). In each pass, we call \( \text{ACTION} \) and then \( \text{MODEL} \), one after the other, until we have built up a stream of forecasts from time 1 up to time \( T \). Then, for each time \( t \) going backwards from \( T \) to 1, we call the \( \text{UTILITY} \) subroutine, then \( \text{F-UTILITY} \), then \( \text{F-MODEL} \), and then \( \text{F-ACTION} \) at the end of the pass, we have the correct array of derivatives \( F.W \), which we can then use to adjust the weights of the action network.

In general, backpropagation through time has the advantage of being relatively quick and exact. That is why I chose it for my natural gas application [11]. However, it cannot account for noise in the process to the controlled. To account for noise in maximizing an arbitrary utility function, we must rely on adaptive critic methods [21]. Adaptive critic methods do not require backpropagation through time in any form, and are therefore suitable for true real-time learning. There are other forms of neurocontrol as well [21] which are not based on maximizing a utility function.

C. Handling Strings of Data

In most of the examples above, I assumed that the training data form one long time series, from \( t = 1 \) to \( t = T \). Thus, in adapting the weights, I always assumed batch learning (except in the code in Section II); the weights were always adapted after a complete set of derivatives was calculated, based on a complete pass through all the data. Mechanically, one could use pattern learning in the backwards pass through time; however, this would lead to a host of problems, and it is difficult to see what it would gain.

Data in the real world are often somewhere between the two extremes represented by Sections II and III. Instead of having a set of unrelated patterns or one continuous time series, we often have a set of time series or strings. For example, in speech recognition, our training set may consist of a set of strings, each consisting of one word or one sen-
tence. In robotics, our training set may consist of a set of strings, where each string represents one experiment with a robot.

In these situations, we can apply backpropagation through time to a single string of data at a time. For each string, we can calculate complete derivatives and update the weights. Then we can go on to the next string. This is like pattern learning, in that the weights are updated incrementally before the entire data set is studied. It requires intermediate storage for only one string at a time. To speed things up even further, we might adapt the net in stages, initially fixing certain weights (like \( w_i \)) to zero or one.

Nevertheless, string learning is not the same thing as real-time learning. To solve problems in neuroidentification and supervised learning, the only consistent way to have internal memory terms and to avoid backpropagation through time is to use adaptive critics in a supporting role [15]. That alternative is complex, inexact, and relatively expensive for these applications; it may be unavoidable for true real-time systems like the human brain, but it would probably be better to live with string learning and focus on other challenges in neuroidentification for the time being.

D. Speeding Up Convergence

For those who are familiar with numerical analysis and optimization, it goes without saying that steepest descent— as in (12)—is a very inefficient method. There is a huge literature in the neural network field on how to speed up backpropagation. For example, Fahlman and Touretzky of Carnegie-Mellon have compiled and tested a variety of intuitive insights which can speed up convergence a hundredfold. Their benchmark problems may be very useful in evaluating other methods which claim to do the same. A few authors have copied simple methods from the field of numerical analysis, such as quasi-Newton methods (BFGS) and Polak-Ribiere conjugate gradients; however, the former works only on small problems (a hundred or so weights) [22], while the latter works well only with batch learning and very careful line searches. The need for careful line searches is discussed in the literature [23], but I have found it to be unusually important when working with large problems, including simulated linear mappings.

In my own work, I have used Shanno’s more recent conjugate gradient method with batch learning; for a dense training set—made up of distinctly different patterns—this method worked better than anything else I tried, including pattern learning methods [12]. Many researchers have used approximate Newton’s methods, without saying that they are using an approximation; however an exact Newton’s method can also be implemented in \( O(N) \) storage, and has worked reasonably well in early tests [12]. Shanno has reported new breakthroughs in function minimization which may perform still better [24]. Still, there is clearly a lot of room for improvement through further research.

Needless to say, it can be much easier to converge to a set of weights which do not minimize error or which assume a simpler network; methods of that sort are also popular, but are useful only when they clearly fit the application at hand for identifiable reasons.

E. Miscellaneous Issues

Minimizing square error and maximizing likelihood are often taken for granted as fundamental principles in large parts of engineering; however, there is a large literature on alternative approaches [12], both in neural network theory and in robust statistics.

These literatures are beyond the scope of this paper, but a few related points may be worth noting. For example, instead of minimizing square error, we could minimize the 1.5 power of error; all of the operations above still go through. We can minimize \( E \) of (5) plus some constant \( k \times \) times the sum of squares of the weights; as \( k \) goes to infinity and the network is made linear, this converges to Kohonen’s pseudoinverse method, a common form of associative memory. Statisticians like Dempster and Efron have argued that the linear form of this approach can be better than the usual least squares methods; their arguments capture the essential insight that people can forecast by analogy to historical precedent, instead of forecasting by a comprehensive model or network. Presumably, an ideal network would bring together both kinds of forecasting [12], [20].

Many authors worry a lot about local minima. In using backpropagation through time in robust estimation, I found it important to keep the “memory” weights near zero at first, and free them up gradually in order to minimize problems. When \( T \) is much larger than \( m \)—as statisticians recommend for good generalization—local minima are probably a lot less serious than rumor has it. Still, with \( T \) larger than \( m \), it is very easy to construct local minima. Consider the example with \( m = 2 \) shown in Table I.

<table>
<thead>
<tr>
<th>( t )</th>
<th>( X(t) )</th>
<th>( Y(t) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>2</td>
<td>1.0</td>
<td>0.1</td>
</tr>
<tr>
<td>3</td>
<td>1.1</td>
<td>0.9</td>
</tr>
</tbody>
</table>

The error for each of the patterns can be plotted as a contour map as a function of the two weights \( w_1 \) and \( w_2 \). (For this simple example, no threshold term is assumed.) Each map is made up of straight contours, defining a fairly sharp trough about a central line. The three central lines for the three patterns form a triangle, the vertices of which correspond roughly to the local minima. Even when \( T \) is much larger than \( m \), conflicts like this can exist within the training set. Again, however, this may not be an overwhelming problem in practical applications [19].

V. Summary

Backpropagation through time can be applied to many different categories of dynamical systems—neural networks, feedforward systems of equations, systems with time lags, systems with instantaneous feedback between variables (as in ordinary differential equations or simultaneous equation models), and so on. The derivatives which it calculates can be used in pattern recognition, in systems identification, and in stochastic and deterministic control. This paper has presented the key equations of backpropagation, as applied to neural networks of varying degrees of complexity. It has also discussed other papers which elaborate on the extensions of this method to more general applications and some of the tradeoffs involved.
REFERENCES


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