FUNCTION APPROXIMATION WITH MULTILAYERED PERCEPTRONS USING $L_1$ CRITERION

by

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April 2003

Thesis submitted in fulfilment of the requirements for the degree of Doctor of Philosophy
ACKNOWLEDGEMENTS

I would like to thank my supervisor, Professor Quah Soon Hoe, for his invaluable help, guidance, understanding, advice, concern, support and encouragement. I am thankful and appreciative that he accepted me to be his Ph.D. student. I would also like to thank Professor Kaj Madsen from The Department of Mathematical Modelling, Technical University of Denmark, Denmark, for allowing me to use his $L_1$ Optimization program.

I am also grateful to Associate Professor Ahmad Abd Majid, Dean of the School of Mathematical Sciences, Universiti Sains Malaysia for his encouragement and for allowing me to use the facilities in the school. I am very thankful to Mr. Wong Ya Ping from Multimedia University, Cyberjaya, for motivating me to look into artificial neural networks and also for helping me in the computer programming.

My appreciation also goes to Dr. Lawrence Chang Hooi Tuang, who helped me to obtain certain papers from the library in the USM Engineering Faculty, to Mr. Tan Ewe Hoe, for translating the abstract into Bahasa Malaysia with his professional touch and to Ms. Catherine Lee Saw Paik, for sacrificing her time and energy to prepare the final version of my thesis for printing.

Above all, I thank God, for His mercy, grace and the opportunity to further my studies.

Ong H.C.
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ABSTRAK

Kaedah ralat kuasa dua terkecil atau kaedah kriteria \( L_2 \) biasanya digunakan bagi persoalan penghampiran fungsian dan pengitlakan di dalam algoritma perambatan balik ralat. Tujuan kajian ini adalah untuk mempersembahkan suatu kriteria ralat mutlak terkecil bagi perambatan balik sigmoid selain daripada kriteria ralat kuasa dua terkecil yang biasa digunakan. Kami membentangkan struktur fungsi ralat untuk diminimumkan serta hasil pembezaan terhadap pemberat yang akan dikemaskinikan. Tumpuan kajian ini ialah terhadap model perseptron multilapisan yang mempunyai satu lapisan tersembunyi tetapi perlaksanaannya boleh dilanjutkan kepada model yang mempunyai dua atau lebih lapisan tersembunyi.

Penyelidikan kami menggunakan hakikat bahawa fungsi perambatan balik sigmoid boleh dibezakan dan menggunakan sifat ini untuk melaksanakan suatu algoritma dua peringkat bagi pengoptimuman \( L_1 \) tak linear oleh Madsen, Hegelund & Hansen (1991) untuk memperoleh keputusan optimum. Algoritma dua-peringkat tersebut merangkumi kaedah tertib pertama yang mengira hampir penyelesaian melalui penggunaan pengaturcaraan linear dan kaedah kuasi-Newton yang menggunakan maklumat tertib kedua yang terhampir untuk menyelesaikan masalah. Kami menunjukkan latar belakang teori dan teorem-teorem penumpuan yang berkaitan dengan algoritma \( L_1 \) dua-peringkat tersebut. Alasan utama bagi menggunakan kriteria ralat mutlak terkecil berbanding dengan kriteria ralat kuasadua terkecil adalah kerana ianya lebih stabil dan tidak mudah dipengaruhi oleh data ekstrim.
Kami menggunakan algoritma $L1$ dua-peringkat untuk mensimulasikan lima fungsi tak linear yang berbeza yang kesemuanya diskalakan supaya sisihan piawai ialah satu dalam satu grid seragam sebesar 2 500 titik pada $[0, 1]^2$. Simulasi tersebut adalah bagi tujuan perbandingan supaya dapat memerhatikan respons pelbagai fungsi dengan menetapkan set latihan pada 225 titik dan menggunakan lapan nod tersembunyi. Kami dapat bahawa algoritma $L1$ dua-peringkat adalah lebih cekap dan efisien berbanding dengan algoritma perambatan balik ralat kuasa dua terkecil dalam kes kelima-lima fungsi tersebut.

Kami mengemukakan penyelesaian teoretis bagi persoalan penghampiran dan pengitlakan dengan menggunakan model perseptron yang mempunyai satu lapisan tersembunyi dan memberi bukti secara membina tentang kemampuannya dalam menghampirkan fungsi. Bagi mengakhiri tesis ini, kami memberi sedikit perbincangan tentang beberapa cara pengitlakan yang mungkin untuk kajian selanjutnya.
ABSTRACT

The least squares error or $L_2$ criterion approach has been commonly used in functional approximation and generalization in the error backpropagation algorithm. The purpose of this study is to present an absolute error criterion for the sigmoidal backpropagation rather than the usual least squares error criterion. We present the structure of the error function to be minimized and its derivatives with respect to the weights to be updated. The focus in the study is on the single hidden layer multilayer perceptron (MLP) but the implementation may be extended to include two or more hidden layers.

Our research makes use of the fact that the sigmoidal backpropagation function is differentiable and uses this property to implement a 2-stage algorithm for nonlinear $L_1$ optimization by Madsen, Hegelund & Hansen (1991) to obtain the optimum result. This is a combination of a first order method that approximates the solution by successive linear programming and a quasi-Newton method using approximate second order information to solve the problem. We show the theoretical background and convergence theorems associated with the 2-stage $L_1$ algorithm. The main reason for using the least absolute error criterion rather than the least squares error criterion is that the former is more robust and less easily affected by noise compared to the latter.

We simulate the 2-stage $L_1$ algorithm over five different non-linear functions which are all scaled so that the standard deviation is one for a large regular grid with 2,500 points on $[0, 1]^2$. The simulation is for comparison purposes. We want to see how the responses are for the various functions keeping each training set fixed at 225 points and using 8 hidden nodes in all the simulation. We find that the 2-stage $L_1$ algorithm
outperforms and is more efficient than the least squares error backpropagation algorithm in the case of all the five functions.

We present a theoretical solution to the generalization and approximation problem using a single hidden layer MLP and give a constructive proof of the density involving the approximation function. To round up the thesis, we give a brief discussion of a few of the many, many possible generalizations to explore.
CHAPTER 1

INTRODUCTION

1.1 A Brief Overview of Neural Network Applications

Artificial neural networks, commonly known as neural networks, have developed rapidly in the late 1980s and are now used widely. Work on artificial neural networks has been motivated right from its inception by the recognition that the brain computes in an entirely different way from the way the conventional digital computer computes. The human brain has the ability to generalize from abstract ideas, recognizes patterns in the presence of noise, quickly recall memories and withstand localized damage. From a statistical perspective, neural networks are interesting because of their potential use in prediction and classification problems. Biological neural networks inspire artificial neural networks (ANNs) as patterns in biological data contain knowledge, if only we can discover it. ANNs find applications in many diverse fields. Possibilities abound in geological, climatic, meteorological, personality, cultural, historical, spectral, electromagnetic, medical, satellite scan data and other data. It remains only for the researcher to glean the essentials and begin to explore the classification and recognition of patterns in data that will lead to discoveries of associations and cause-effect relationships. In short, its usefulness is almost inexhaustive. Haykin (1999), Looney (1997) and Callan (1999) cover the basic concepts and gives a comprehensive foundation of neural networks recognizing the multidisciplinary nature of the subject.

Neural networks have been used for a wide variety of applications where statistical methods are traditionally employed. ANNs have in recent years developed into
powerful tools for solving optimization problems within, for example, classification, estimation and forecasting (Cheng & Titterington, 1994, Ripley, 1994 and Schioler & Kulezyeki, 1997). ANNs models and learning techniques appear to provide applied statisticians with a rich and interesting class of new methods applicable to any regression problem requiring some sort of flexible functional form. ANNs can be constructed without any assumptions concerning the functional form of the relationship between the predictors and response. ANNs, if trained on large data sets and suitably tested, can be quite successful in purely predictive problems covering various forms of pattern recognition problems (Stern, 1996). This serves as an advantage in predictive settings but this can be a disadvantage to applied statisticians investigating an understanding of the relationship between the variables. In such settings, investigators can frequently specify a plausible probability model with meaningful parameters. The advantages of ANNs are likely to outweigh the disadvantages in purely predictive problems but not for problems in which we are interested in determining the relationship between the various quantities.

Advantages to the approach using ANNs include: no need to start from an a priori mathematical form for the solution, no programming required to get a solution and easily extended solution including more variables to obtain more accuracy (Roy, Lee, Mentele & Nau, 1993). For example, longitudinal data could be added as an additional input variable and the ANN could be retrained. However, too many neurons bias the fit to the details of the data rather than the underlying pattern. Ever since the advent of renewed interest in ANNs with improved computer systems in the late 1980s, many papers have been published researching into various aspects of the ANNs. For example, there are attempts to explain and interprete how the ANNs work (Benitez, Castro &
Requena, 1997), efforts to provide a system that will outperform the sigmoidal backpropagation network models (Van der Walt, Barnard & Van Deventer, 1995) and various issues affecting the ANN models.

ANN models have several important characteristics, which are important:

(i) The major problem of developing a realistic system is obviated because of the ability of the ANN models to be trained using data records for the particular system of interest.

(ii) The ANN models possess the ability to learn nonlinear relationships with limited prior knowledge about the process structure.

(iii) ANN models, by their very nature, have many inputs and outputs and so can be applied to multivariable systems.

(iv) The structure of the ANN model is highly parallel in nature. This is likely to give rise to three benefits: Very fast parallel processing, fault tolerance and robustness.

1.2 Types of Artificial Neural Network (ANN) Models

1.2.1 Classification of Neural Networks

The ability of the ANN models to learn from data with or without a teacher has endowed them with a powerful property. In one form or another, the ability of the ANN models to learn from examples has made them invaluable tools in such diverse applications. The new wave of ANN models came into being because learning could be
performed at multiple levels, learning pervades every level of the intelligent machine in an increasing number of applications.

The desired behaviour of the network provides another approach to distinguishing networks. For example, the desired function of the ANN model may be specified by enumeration of a set of stable network states or by identifying a desired network output as a function of the network inputs and current state. Popular examples of classifying ANN models by processing objectives are:

(i) **The pattern associator.** This ANN functionality relates patterns which may be vectors. Commonly, it is implemented using feedforward networks. This type of network includes the multilayer perceptron (MLP), Adaline/Madaline units and networks and Hebbian or correlation based learning (Rumelhart, Hinton & Williams, 1986 and Karhunen & Joutsensalo, 1995).

(ii) **The content-addressable or associative memory model (CAM or AM).** This model is based on the implementation of association and is best exemplified by the Hopfield model (Hopfield, 1982 and Paik & Katsaggelos, 1992).

(iii) **Self-organizing networks.** These networks exemplify neural implementation of unsupervised learning in the sense that they typically self-organize input patterns into classes or clusters based on some form of similarity. Examples include learning vector quantization (LVQ) algorithm (which is similar to the Kohonen SOFM) formulation and the adaptive resonance architectures (Kohonen, 1988 and Linde, Buzo & Gray, 1980).
1.2.2 A Comparison Between the Multilayer Perceptron (MLP) and the Radial Basis Function (RBF)

In this section, we briefly introduce and compare two network structures which are used for the problem of function approximation, namely the multilayer perceptron (MLP) and the radial basis function (RBF) network. Both are feedforward structures with hidden layers and training algorithms, which may be used for mapping. The training of an MLP is usually accomplished by using a backpropagation (BP) algorithm that involves two phases, that is, the forward phase and the backward phase which will be described in Chapter 2 (Rumelhart, Hinton & Williams, 1986). The RBF networks use memory based learning for their design (Park & Sandberg, 1993). RBF networks differ from multilayer perceptrons in some fundamental respects:

(i) RBF networks are local approximators whereas MLP are global approximators.

(ii) RBF networks have a single hidden layer whereas an MLP can have any number of hidden layers.

(iii) The output of a RBF network is always linear whereas the output in a MLP can be linear or nonlinear.

(iv) The activation function of the hidden layer in a RBF network computes the Euclidean distance between the input signal vector and parameter vector of the network, whereas the activation function of a MLP computes the inner product between the input signal vector and the pertinent synaptic weight vectors.

1.3 Regression and Neural Network

Many ideas in statistics can be expressed in neural network notation. They include regression models from simple linear regression to projection pursuit regression,
nonparametric regression, generalized additive models and others. Also, included are many approaches to discriminant analysis such as logistic regression, Fisher’s linear discriminant function and classification trees, as well as methods for density estimation of both parametric and nonparametric types. Regression is used to model a relationship between variables. By explaining neural networks in relation to regression analysis some deeper understanding can be achieved.

The general form of the regression model is

$$\eta = \sum_{i=0}^{N} \beta_i x_i$$

with $\eta = h(\mu)$ and $E(z) = \mu$,

where $h(.)$ is the link function, $\beta_i$ are the coefficients, $N$ is the number of covariate variables and $\beta_0$ is the intercept. A random component of the response variable $z$ in the model has mean $\mu$ and variance $\sigma^2$ while a systematic component relates the stimuli $x_i$ to a linear predictor $\eta = \sum_{i=0}^{N} \beta_i x_i$.

The generalized linear model reduces to the multiple linear regression model if we believe that the random component has a normal distribution with mean zero and variance $\sigma^2$ and we specify the link function $h(.)$ as the identity function. The model is then:

$$z_p = \beta_0 + \sum_{i=1}^{N} \beta_i x_{pi} + \epsilon_p$$

where $\epsilon_p \sim N(0, \sigma^2)$.

The objective of this regression problem is to find the coefficients $\beta_i$ that minimize the sum of squared errors $E = \sum_{p=1}^{n} \left( z_p - \sum_{i=1}^{n} \beta_i x_{pi} \right)^2$. 
To find the coefficients, we must have a data set that includes the independent variables and associated known values of the dependent variable (akin to a training set in supervised learning in neural networks).

The problem is equivalent to a two layer feedforward neural network (zero hidden layer) as shown in Figure 1.1. The independent variables correspond to the inputs of the neural network and the response variable $z$ to the output. The coefficients, $\beta_i$, correspond to the weights in the neural network. The activation function is the identity function. To find the weights in the neural network, we would use backpropagation. A difference in the two approaches is that multiple linear regression has a closed form solution for the coefficients, while neural network uses an iterative process.

![Figure 1.1](image)

In general, any generalized linear model can be mapped onto an equivalent two layer neural network. The activation function is selected to match the inverse of the link function and the cost function is selected to match the deviance, which is based on maximum likelihood theory and is determined by the distribution of the random component. The generalized linear model attempts to find coefficients to minimize the deviance. The theory for these problems already exist and the neural networks as
presented only produce similar results, adding nothing to the theory. These examples only give some insight and understanding of neural networks.

In regression models, a functional form is imposed on the data. In the case of multiple linear regression, this assumption is that the outcome is related to a linear combination of the independent variables. If this assumed model is not correct, it will lead to error in the prediction. An alternate approach is to not assume any functional relationship and let the data define the functional form. In a sense, we let the data speak for itself. This is the basis of the power of the neural networks. As will be mentioned in the later chapters, a three layered, feedforward network (a single hidden layer) with sigmoid activation functions is a universal approximator because it can approximate any continuous function to any degree of accuracy. A neural network is extremely useful when we do not have any idea of the functional relationship between the dependent and independent variables. If we have an idea of the functional relationship, we are better off using a regression model.

1.4 Format of Presentation

Having briefly explained the increasingly important role that ANNs play in the various fields of research, the purpose of this thesis is to present an absolute error criterion for sigmoidal backpropagation rather than the usual least squares error criterion. The research makes use of the fact that the sigmoidal backpropagation function is differentiable and uses this property to implement a 2-stage algorithm for nonlinear $L_1$ optimization by Hald (1981) to obtain the optimum result. The main reason for using the least absolute criterion rather than the least squares error criterion is that the former
is more robust and less easily affected by noise compared to the latter (Namba, Kamata & Ishida, 1996 and Taji, Miyake & Tamura, 1999). Cadzow (2002) mentioned that in cases where the data being analyzed contain a few data outliers, a minimum $L_1$ approximate is preferable to approximation using other $L_p$ norms with $p > 1$ since it tends to ignore bad data points.

For the important problem of approximating a function to data that might contain some wild points, minimization of the $L_1$ norm residual is superior to minimization using other $L_p$ norms with $p > 1$ (Hald & Madsen, 1985 and Portnoy & Koenker, 1997). The larger the value of $p$, the more the focus is placed on the data points with largest deviations from the approximating function. Going to $L_\infty$, the maximum deviation will be minimized. The minimum $L_\infty$ norm is used when the maximum error vector is minimized among all possible vectors corresponding to the desired vector (Ding & Tso, 1998). In other words, the $L_\infty$ norm is relevant in sensitivity analysis, particularly in robotics. However, for the function approximation problem here, the $L_1$ norm is preferable. Depending on the type of approximating function, the optimization problem will be linear or nonlinear in the variable function parameters (El-Attar, Vidyasagar & Dutta, 1979).

In Chapter 2, we will present the structure and derive the algorithm for the single hidden layer backpropagation with the least squares criterion together with the reasons for using only the single hidden layer for our application. However, the algorithm and implementation may be extended to include two or more hidden layers. The report of deVilliers & Barnard (1992) states that while optimally trained MLPs of one and two
hidden layers perform no different statistically (there is no significant difference between them on the average), the network with a single hidden layer performs better on recognition. Other findings were that networks with two hidden layers are often more difficult to train and are affected more by the choice of an initial weight set and the architecture. This is then followed by a discussion of the various issues in the implementation of the multilayer perceptrons (MLPs).

We then present the 2-stage nonlinear $L1$ method and the sigmoidal backpropagation function for optimization in Chapter 3. We then look into some convergence theorems related to the 2-stage $L1$ algorithm for the error backpropagation function. The performance of the 2-stage nonlinear $L1$ method is then tested and discussed in Chapter 4 with reference to five different functions, namely, a simple interaction function, a radial function, a harmonic function, an additive function and a complicated interaction function. 225 training sets were used for the five functions and the results were generated graphically using MATLAB. A comparison of the performance is then graphically shown between the 2-stage $L1$ method of Chapter 3 and the least squares error algorithm explained in Chapter 2. Having discussed both methods mentioned in Chapters 2 and 3, we present the theoretical solution for the generalization and approximation problem in Chapter 5. In Chapter 6, we conclude with a discussion of the results obtained and possible ways to generalize and explore in future research.
CHAPTER 2

THE ERROR BACKPROPA GATION FOR THE MULTILAYER PERCEPTRON (MLP)

2.1 An Overview of the Multilayer Perceptron

The perceptron is the simplest form of a neural network and it consists of a single neuron with adjustable synaptic weights and bias. The perceptron built around a single neuron is limited to performing pattern classification with only two classes. The multilayer feedforward networks, an important class of neural networks, consist of sensory neurons or nodes that constitute the input layer, one or more hidden layers of computation nodes and an output layer of computation nodes. The signal is propagated from the input layer in a forward direction, on a layer-by-layer basis. These neural networks are commonly referred to as multilayered perceptrons (MLPs). The MLPs have been successfully applied to diverse problems by training them in a supervised manner with a highly popular algorithm known as the error backpropagation algorithm.

The error backpropagation algorithm defines two sweeps of the network. In the forward sweep from the input layer to the output layer, an activity pattern (input vector) is applied to the input sensory nodes of the network and propagates layer by layer during which the synaptic weights are fixed. This produces an output as a response from the network. The error signal is then calculated as the difference between the response of the network and the actual (desired) response. During the backward sweep, the synaptic weights are adjusted according to an error correction rule. The synaptic weights are adjusted to make the network output closer to the desired output. The model of each neuron includes a smooth nonlinear activation function, which is
differentiable. The commonly used form of nonlinear activation function is the sigmoidal function with a bias or the hyperbolic tangent function.

It is well known that a two layered feedforward neural network, that is, one that does not have any hidden layers, is not capable of approximating generic nonlinear continuous functions (Widrow, 1990). On the other hand, four or more layered feedforward neural networks are rarely used in practice (Scarselli & Tsoi, 1998). Hence, almost all the work deals with the most challenging issue of the approximation capability of the three layered feedforward neural network (or the single hidden layered MLP). Hornik, Stinchcombe & White (1989), Cybenko (1989), Leshno, et al. (1993) and Funahashi (1989) showed that it is sufficient to use a single hidden layered MLP in universal approximations. Petrushev (1998) showed that a feedforward neural network with one hidden layer of computational nodes given by certain sigmoidal functions have the same efficiency of approximation as other more traditional methods of multivariate approximations such as polynomials, splines or wavelets. Even though these results show that one hidden layer is sufficient, it may be more parsimonious to use two or more hidden layers. Our study will focus only on the single hidden layer MLP but can be generalized to include two or more layers.

2.2 Derivation of the Error Backpropagation Algorithm

An MLP with a single hidden layer is shown in Figure 2.1. There are \( I \) neurons in the input layer, \( J \) neurons in the hidden layer and \( K \) neurons in the output layer. \( K \) is normally taken to be one for the case of functional approximation. The interconnection weights from the input to the hidden layer are denoted by \( \{w_{ij}\} \) while that from the
hidden layer to the output is denoted by \( \{u_{jk}\} \). The sigmoidal activation function for the hidden and output layers are \( h(\cdot) \) and \( g(\cdot) \) respectively. Each exemplar vector \( x^{(q)} \) is mapped into an output \( z^{(q)} \) from the network and compared to the target output \( t^{(q)} \), where \( q = 1, 2, \ldots, Q \) is the number of exemplars or training sets.

![Diagram of a single hidden layer neural network for training.](image)

Figure 2.1 Single hidden layer neural network for training.

The error backpropagation algorithm with unipolar sigmoids and its derivatives can be derived for the network in Figure 2.1. Let

\[
    r_j = \sum_{i=1}^{l} w_{ij} x_i \quad \text{and} \quad y_j = h \left( \sum_{i=1}^{l} w_{ij} x_i \right) = \frac{1}{1 + \exp \left( -a_1 \sum_{i=1}^{l} w_{ij} x_i + b_1 \right)}
\]

and

\[
    s_k = \sum_{j=1}^{J} u_{jk} y_j \quad \text{and} \quad z_k = g \left( \sum_{j=1}^{J} u_{jk} y_j \right) = \frac{1}{1 + \exp \left( -a_2 \sum_{j=1}^{J} u_{jk} y_j + b_2 \right)}
\]

for \( h(r_j) = \frac{1}{1 + \exp \left( -a_1 r_j + b_1 \right)} \) and \( g(s_k) = \frac{1}{1 + \exp \left( -a_2 s_k + b_2 \right)} \), where \( a_1, a_2 \) are rate factors and \( b_1, b_2 \) are biases.
Total sum of squared errors = $E(w, u)$

$$= \sum_{q=1}^{Q} \left\| t^{(q)} - z^{(q)} \right\|^2,$$

where $q = 1, 2, \cdots, Q$ are the number of exemplars

$$= \sum_{q=1}^{Q} \left[ \sum_{k=1}^{K} (t_k^{(q)} - z_k^{(q)})^2 \right]$$

$$= \sum_{q=1}^{Q} \left[ \sum_{k=1}^{K} \left( t_k^{(q)} - g \left( \sum_{j=1}^{J} u_{jk} y_j \right) \right)^2 \right]$$

$$= \sum_{q=1}^{Q} \left[ \sum_{k=1}^{K} \left( t_k^{(q)} - g \left( \sum_{j=1}^{J} u_{jk} \left( \sum_{i=1}^{I} w_{yi} x_i \right) \right) \right)^2 \right].$$

To minimize $E(w, u)$, the necessary conditions are

$$\frac{\partial E(w, u)}{\partial w_{ij}} = 0 \text{ and } \frac{\partial E(w, u)}{\partial u_{jk}} = 0.$$

The steepest linear gradient descent for iterative updating of individual weights in the error function during the $r$th step takes a generalized Newtonian form, i.e.,

$$w_{ij}^{(r+1)} = w_{ij}^{(r)} - m_1 \frac{\partial E(w, u)}{\partial w_{ij}} \text{ and } u_{jk}^{(r+1)} = u_{jk}^{(r)} - m_2 \frac{\partial E(w, u)}{\partial u_{jk}},$$

where $m_1$ and $m_2$ are the step sizes.

2.2.1 Derivation of the Weight Increment in the Outer Layer $u_{jk}$

$$\frac{\partial E}{\partial u_{jk}} = \frac{\partial E}{\partial s_k} \cdot \frac{\partial s_k}{\partial u_{jk}} \quad \text{since } s_k = s_k(u_{jk}) \text{ functionally}$$

$$= \frac{\partial E}{\partial z_k} \cdot \frac{\partial z_k}{\partial s_k} \cdot \frac{\partial s_k}{\partial u_{jk}} \quad \text{since } z_k = g(s_k) \text{ functionally}.$$

14
\[
\frac{\partial}{\partial z_k} \left( \sum_{k=1}^{K} (t_k - z_k)^2 \right) = \frac{\partial g(s_k)}{\partial s_k} \cdot \frac{\partial s_k}{\partial u_{jk}} \\
= -2(t_k - z_k) \cdot g'(s_k) \cdot \frac{\partial}{\partial u_{jk}} \left( \sum_{j=1}^{J} u_{jk} y_j \right) \\
= -2(t_k - z_k) \cdot g'(s_k) \cdot y_j.
\]

\[
g'(s_k) = \frac{\partial}{\partial s_k} \left[ 1 + \exp(-a_2 s_k + b_2) \right]^{-1}
\]

\[
= a_2 \left[ 1 + \exp(-a_2 s_k + b_2) \right]^{-2} \exp(-a_2 s_k + b_2) \\
= a_2 \left[ 1 + \exp(-a_2 s_k + b_2) \right]^{-2} \left[ 1 + \exp(-a_2 s_k + b_2) - 1 \right] \\
= a_2 z_k^2 \left[ \frac{1}{z_k} - 1 \right] \quad \text{where} \quad z_j = \left[ 1 + \exp(-a_2 s_k + b_2) \right]^{-1} \\
= a_2 z_k (1 - z_k).
\]

\[
\frac{\partial E}{\partial u_{jk}} = -2a_2 (t_k - z_k) \cdot z_k (1 - z_k) \cdot y_j \\
\therefore \quad u_{jk}^{(r+1)} = u_{jk}^{(r)} + m_2 (t_k - z_k) z_k (1 - z_k) y_j \\
\tag{2.1}
\]

where \( -2 \) and \( a_2 \) are absorbed into the step gain \( m_2 \).

2.2.2 Derivation of the Weight Increment in the Hidden Layer \( w_{ij} \)

\[
\frac{\partial E}{\partial w_{ij}} = \frac{\partial E}{\partial r_j} \cdot \frac{\partial r_j}{\partial w_{ij}} \quad \text{since} \quad r_j = r_j(w_{ij}) \quad \text{functionally} \\
= \frac{\partial E}{\partial y_j} \cdot \frac{\partial y_j}{\partial r_j} \cdot \frac{\partial r_j}{\partial w_{ij}} \quad \text{since} \quad y_j = h(r_j) \quad \text{functionally} \\
= \frac{\partial}{\partial y_j} \left( \sum_{k=1}^{K} (t_k - z_k)^2 \right) \cdot \frac{\partial h(r_j)}{\partial r_j} \cdot \frac{\partial}{\partial w_{ij}} \left( \sum_{i=1}^{I} w_{ij} x_i \right)
\]
Having derived the backpropagation training equation to minimize the partial sum of squared errors function \( E^{(q)} \) for any fixed \( q \)th exemplar pair \((x^{(q)}, t^{(q)})\), an implementation of the error backpropagation can be done using the flowchart in Figure 2.2. After all the \( Q \) exemplar pairs have been trained in each cycle, an epoch has been completed.
input/read data \((x^{(g)}, y^{(g)})\)
input no. of epochs \(R\),
set parameters \(a_i, b_i, m_i, i = 1, 2\)

generate initial weights \(w_{ij}^{(1)}, u_{jk}^{(1)}\)

set number of epochs \(r = 1\)

set number of exemplars, \(q = 1\)

compute output \(z^{(g)}\)

update weights
\[
\begin{align*}
    w_{ij}^{(r+1)} &= w_{ij}^{(r)} + m_1 \frac{\partial E(w, u)}{\partial w_{ij}} \\
    u_{jk}^{(r+1)} &= u_{jk}^{(r)} + m_2 \frac{\partial E(w, u)}{\partial u_{jk}}
\end{align*}
\]

Figure 2.2 Backpropagation flowchart for the single hidden layer MLP.
2.3 Extension of the Error Backpropagation Algorithm to Two Hidden Layers

Relatively little is known concerning the advantages and disadvantages of using a single hidden layer with many units (neurons) over many hidden layers with fewer units. Some authors see little theoretical gain in considering more than one hidden layer since a single hidden layer model suffices for density or a good approximation (Pinkus, 1999). One important advantage of the multiple (rather than single) hidden layer has to do with the existence of a locally supported or at least 'localized' function in the two hidden layer model. 

An extended architecture from Figure 2.1 is shown in Figure 2.3 with two hidden layers. The interconnection weights for the input/output training pairs \{(x^{(q)}, t^{(q)}): q = 1, 2, \ldots, Q\} are \(v_{\ell i}, w_{ij}\) and \(u_{jk}\) with \(\ell = 1, 2, \ldots, L, i = 1, 2, \ldots, I, j = 1, 2, \ldots, J\) and \(k = 1, 2, \ldots, K\) (where \(k\) is normally taken to be 1 for the approximation problem). We derive the computational formulae for updating the weights via steepest descent in the extended backpropagation algorithm using similar sigmoidal functions as in the case of the single hidden layer.

![Figure 2.3 The extended two hidden layer neural network.](image-url)
The partial derivatives in the weight updates are as follows:

\[ v_{li}^{(r+1)} = v_{li}^{(r)} + m_0 \frac{\partial E(v_{li}, w_{ij}, u_{jk})}{\partial v_{li}}, \]

\[ w_{ij}^{(r+1)} = w_{ij}^{(r)} + m_1 \frac{\partial E(v_{li}, w_{ij}, u_{jk})}{\partial w_{ij}}, \]

\[ u_{jk}^{(r+1)} = u_{jk}^{(r)} + m_2 \frac{\partial E(v_{li}, w_{ij}, u_{jk})}{\partial u_{jk}}. \]

We use the same parameter names and indices for the output layers in Figures 2.1 and 2.3. The second hidden layer is the hidden layer adjacent to the output layer. Using Eq.(2.1) for the increment on \( u_{jk} \), the weights from the second hidden layer to the output layer, we have

\[ u_{jk}^{(r+1)} = u_{jk}^{(r)} + m_2(t_k - z_k)z_k(1 - z_k)y_j. \]

The difference in the variable designations here as compared to the case of a single hidden layer in Section 2.2 is the input \( x_t = (x_1, \cdots, x_L) \) and the additional first hidden layer weights \( c_i = (c_1, \cdots, c_T). \) The corresponding weights from the input to the first hidden layer are \( v_{li}. \) The updates in the weights from the first to the second hidden layer, upon making changes to Eq.(2.2) are:

\[ w_{ij}^{(r+1)} = w_{ij}^{(r)} + m_1 \sum_{k=1}^{K} (t_k - z_k) \cdot z_k \cdot (1 - z_k) u_{jk}^{(r)} \cdot y_j \cdot (1 - y_j) \cdot c_i. \]

Let \( p \) be the sigmoidal activation function for the first hidden layer. The value of the neuron in the first hidden layer

\[ c_i = p(n_i) = p \left( \sum_{t=1}^{L} v_{li} x_t \right) = \frac{1}{1 + \exp \left( - a_0 \sum_{t=1}^{L} v_{li} x_t + b_0 \right)}. \]
for \( n_i = \sum_{t=1}^{L} v_{li} x_t \) where \( a_0 \) and \( b_0 \) are the corresponding rate factor and bias respectively.

The derivation of the weight updates at the first hidden layer is more tedious and we suppress the \( q \) subscripts as compared to the derivation in Section 2.2.

The total sum of squared errors

\[
E = E(v_{li}, w_{ij}, u_{jk})
\]

\[
= \sum_{k=1}^{K} (t_k - z_k)^2
\]

\[
= \sum_{k=1}^{K} (t_k - g(s_k))^2
\]

\[
= \sum_{k=1}^{K} \left( t_k - g \left( \sum_{j=1}^{J} u_{jk} y_j \right) \right)^2
\]

\[
= \sum_{k=1}^{K} \left( t_k - g \left( \sum_{j=1}^{J} u_{jk} h(r_j) \right) \right)^2
\]

\[
= \sum_{k=1}^{K} \left( t_k - g \left( \sum_{j=1}^{J} u_{jk} h \left( \sum_{l=1}^{I} w_{lj} c_l \right) \right) \right)^2
\]

\[
= \sum_{k=1}^{K} \left( t_k - g \left( \sum_{j=1}^{J} u_{jk} h \left( \sum_{l=1}^{I} w_{lj} p(n_l) \right) \right) \right)^2
\]

\[
= \sum_{k=1}^{K} \left( t_k - g \left( \sum_{j=1}^{J} u_{jk} h \left( \sum_{l=1}^{I} w_{lj} p \left( \sum_{t=1}^{L} v_{li} x_t \right) \right) \right) \right)^2
\]

It can be seen from Figure 2.3 and Eq.(2.3) that each weight \( v_{li} \) at the first hidden layer affects not only every difference \( t_k - z_k \) but also the value in every unit in the second
hidden layer. To derive \( \frac{\partial E}{\partial v_{tl}} \), we need to sum the total error adjustments over all the units from \( j = 1, \ldots, J \) and \( k = 1, \ldots, K \). Thus, we use

\[
E_{jk} = \left( t_k - g \left( u_{jk} h \left( \sum_{l=1}^{L} w_{lj} p \left( \sum_{t=1}^{L} v_{tl} x_t \right) \right) \right) \right)^2.
\]

(2.4)

From Eq.(2.4), we can write down the chain rule in terms of dependent variables that start at the output layer and we work backwards to the first hidden layer. This leads to the partial derivative

\[
\frac{\partial E_{jk}}{\partial v_{ti}} = \left( \frac{\partial E_{jk}}{\partial z_k} \right) \left( \frac{\partial z_k}{\partial s_k} \right) \left( \frac{\partial s_k}{\partial t_i} \right)
\]

\[
= \left( \frac{\partial E_{jk}}{\partial z_k} \right) \left( \frac{\partial z_k}{\partial s_k} \right) \left( \frac{\partial s_k}{\partial t_j} \right) \left( \frac{\partial u_{jk}}{\partial v_{tl}} \right)
\]

\[
= \left[ (-2)(t_k - z_k) \right] \left[ g'(s_k)u_{jk} \right] \left[ h'(r_j)w_{lj} \right] p'(n_t) x_t.
\]

(2.5)

The derivatives above all have the same form of \( \theta(1-\theta) \). Substituting the derivatives into Eq.(2.5) and absorbing the 2 into \( m_0 \) just as in Section 2.2, we obtain the \((r+1)\)st iterate
We sum up for $k = 1, \ldots, K$ and $j = 1, \ldots, J$ to obtain the final update

$$v_{ti}^{(r+1)} = v_{ti}^{(r)} + m_{0} \sum_{k=1}^{K} [(t_{k} - z_{k}) \cdot z_{k} \cdot (1 - z_{k})] u_{jk} \cdot y_{j} \cdot (1 - y_{j}) \cdot w_{ij} \cdot c_{i} (1 - c_{i}) \cdot x_{t}.$$ 

To summarize the learning equation for updating the weights in the two hidden layers MLP, we have

$$u_{jk}^{(r+1)} \leftarrow u_{jk}^{(r)} + m_{2} (t_{k} - z_{k}) z_{k} (1 - z_{k}) y_{j}$$

$$w_{ij}^{(r+1)} \leftarrow w_{ij}^{(r)} + m_{3} \sum_{k=1}^{K} [(t_{k} - z_{k}) \cdot z_{k} \cdot (1 - z_{k})] u_{jk}^{(r)} \cdot y_{j} \cdot (1 - y_{j}) \cdot c_{i}$$

$$v_{ti}^{(r+1)} \leftarrow v_{ti}^{(r)} + m_{0} \sum_{k=1}^{K} [(t_{k} - z_{k}) \cdot z_{k} \cdot (1 - z_{k})] \sum_{j=1}^{J} [u_{jk}^{(r)} \cdot y_{j} \cdot (1 - y_{j}) \cdot w_{ij}^{(r)}] c_{i} (1 - c_{i}) \cdot x_{t}$$

### 2.4 Issues Related to the Implementation of the MLP

The gradient training of MLPs encounters much problematic behaviour unfortunately. The problems encountered in the use of the backpropagation method as a new tool led to some failures which were actually due to one or more of these problems. Some works give bounds on the number of nodes needed to realize a desired approximation. In general, the bounds depend also on the method used to measure the complexity of the approximated function and on whether the proof is constructive. Thus, a satisfactory comparison is difficult (Scarselli & Tsoi, 1998). In addition to an optimum number of hidden nodes, there is also the problem of testing irrelevant hidden units. The work of Davies (1977) appears relevant in testing the irrelevant hidden unit. The number of nodes and the convergence rate are related. If we want to approximate a sufficiently smooth function $f$ using the form $z$, with $n$ neurons and using a bounded activation function (for example, the sigmoidal function), Jones (1992) showed that the
convergence for $L^2$ approximation is $O(1/\sqrt{n})$. A heuristic reason why the projection method in the MLP might work well with a modest number of hidden units is that the first stage allows a projection onto a subspace of much lower dimensionality within which the approximation can be performed.

The quality of an artificial neural network estimate depends on the number of subjects or data available with the estimate improving as the number of subjects increases (Brown, Branford & Moran, 1997). They also showed that the results obtained when neural networks are applied to survival data depend critically on the treatment of censoring in the data. There may be a large amount of noisy and missing data. The learning requires careful and generalized training with validation and verification testing being a crucial stage.

Early stopping using a validation set or early stopping using a fixed training level error is one of the methods that aim to prevent overtraining due to too powerful a model class, noisy training examples or a small training set. Cataltepe, Abu-Mostafa & Magdon-Ismail (1999) analyzed early stopping at a certain training error minimum and showed that one should minimize the training error as much as possible when all the information available about the target is the training set. They also demonstrated that when additional information is available early stopping could help. When the number of training data becomes infinitely large and they are unbiased, the network parameter converges to one of the local minima of the empirical risk function (expected loss) to be minimized. When the number of training data is finite, the true risk function (generalization error) is different from the empirical risk function. Thus, since the
training examples are biased, the network parameters converge to a biased solution. This is known as overfitting or overtraining because the parameter values fit too well the speciality of the biased training example and are not optimal in the sense of minimizing the generalization error given the risk function. There are a number of methods to avoid overfitting like, for example, model selection methods by Murata, Yoshizawa & Amari (1994). For optimum performance when the number of parameters, \( m \), is large, we should use almost all the \( t \) examples in the training set and only \( (t / \sqrt{2m}) \% \) for cross validation (Amari et al., 1997). In the asymptotic phase where \( t \) is sufficiently large, the asymptotic theory of statistics is applicable and the estimated parameters are approximately normally distributed around the true values.

There are several ways to improve the generalization capability of a neural network. One is to collect more training data. This may be difficult for some applications and it is not clear when the amount of training data is enough. The other approach is to constrain the neural network so that it fits the problem. Jean & Wang (1994) incorporate a smoothing constraint into the objective function of the backpropagation to seek a solution with smoother connection weights. They develop a weight-smoothing algorithm for that purpose and the resulting network generalized better as confirmed by simulations.

For capacity and universality in application to functional approximation, it is apparently better for the number of hidden nodes to be as large as possible. Lawrence, Giles & Tsoi (1997) found that the backpropagation fails to find an optimal solution in many cases. Furthermore, networks with more weights than might be expected can result in lower training and generalization errors in certain cases as MLP trained with backpropagation