Statistical Learning with Artificial Neural Network Applied to Health and Environmental Data

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Statistical Learning with Artificial Neural Network Applied To
Health and Environmental Data

by

Taysseer Sharaf

A dissertation submitted in partial fulfillment
of the requirements for the degree of
Doctor of Philosophy
Mathematics & Statistics
College of Arts and Sciences
University of South Florida

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Dan Shen, Ph.D.

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Keywords: Statistical Learning, Bayesian Learning for ANN, Artificial Neural Network,
Cancer Survival, Global Warming

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Dedication

This doctoral dissertation is dedicated to my parents (Mahmoud Sharaf and Samiaa Hassan), my wife (Sara Mahmoud), my Children (Lujyeen Sharaf, Adam Sharaf and Ryan Sharaf), and my brothers and sisters.
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I would like to express my deepest gratitude to my advisor Professor Chris P. Tsokos for his endless support, motivation, encouragement and advice during my graduate study. I will be owing him for my success as a researcher and teacher in years to come.

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Finally, I could not reach this point without the endless support of my beloved wife. She was always by my side supporting me and encouraging me to do my best. Thank you Sara for being there for me and taking care of our lovely children when I was not there.
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The current study illustrates the utilization of artificial neural network in statistical methodology. More specifically in survival analysis and time series analysis, where both holds an important and wide use in many applications in our real life. We start our discussion by utilizing artificial neural network in survival analysis. In literature there exist two important methodology of utilizing artificial neural network in survival analysis based on discrete survival time method. We illustrate the idea of discrete survival time method and show how one can estimate the discrete model using artificial neural network. We present a comparison between the two methodology and update one of them to estimate survival time of competing risks.

To fit a model using artificial neural network, you need to take care of two parts; first one is the neural network architecture and second part is the learning algorithm. Usually neural networks are trained using a non-linear optimization algorithm such as quasi Newton Raphson algorithm. Other learning algorithms are base on Bayesian inference. In this study we present a new learning technique by using a mixture of the two available methodologies for using Bayesian inference in training of neural networks. We have performed our analysis using real world data. We have used patients diagnosed with skin cancer in the United states from SEER database, under the supervision of the National Cancer Institute.

The second part of this dissertation presents the utilization of artificial neural to time series analysis. We present a new method of training recurrent artificial neural network with Hybrid Monte Carlo Sampling and compare our findings with the popular auto-regressive integrated moving average (ARIMA) model. We used the carbon dioxide monthly average emission to apply our comparison, data collected from NOAA.
Chapter 1

Introduction

Artificial intelligence (A.I) is an important academic field study that studies methods of making machines and software that possess human intelligence and even more. For example, AI is used to develop robots that can do duties making human life more easier. The development in the field of AI is growing exponentially and includes the combination of several subjects such as; Mathematics, Physics, Engineering, Health Sciences, etc. Recently Statistics was found one of the important subjects that AI should include. Peter Norvig said:

AI had some early success by ignoring probabilities, but once we started to run up against the real-world, we had to re-tool and move away from logic towards probability, statistics, and game theory [1]. In the same article we refer to another quote by Yee Whye Teh:

Twenty years ago the dominant ideas in AI were logic-based reasoning. Now the exciting ideas are around intelligent behaviour from noisy examples, often based around statistical principles [1].

The attention in AI has been drawn to Statistical principles lately, and one of the important tools in AI utilizing statistical methods is Artificial Neural Networks (ANN). ANN is a mapping of human biological nervous system. It learns by example like humans and may also be given the opportunity of self learning.

In this dissertation we study the use of ANN in statistical methods such as survival analysis and time series analysis, giving an example for making machines think statistically and have the capabilities of solving real world problems that our society faces on a daily bases. ANN is popular for capturing non-linear complex patterns within a set of data, especially if we are facing huge data sets, which makes ANN a strong competitive among other methods in dealing with BIG DATA. ANN has applications in several fields in life.
such as engineering, predicting time for traffic signals, and in economics, forecasting stock markets and home appraisals saving time on buyers and loaners inspection among other vital fields in life. We highlight in the next section the use of ANN in medical sciences. In which there exist a rich literature materials covering the significant finding of using ANN in solving medical scientific problems.

1.1 The Use of Artificial Neural Network in Health Sciences

In medical sciences, most of the proposed applications of ANN were on prognostic models. For example, one of the most paramount research entities is cancer. Classifying a tumor as malignant or benign is important in cancer research. Chen et al in 2002 used ANN to diagnose breast cancer tumors, [2]. Ercal in 1994 presented an ANN model to distinguish between three benign skin cancer categories and malignant melanoma, [3]. But fitting a complex non-linear modeling such as ANN in regression problems is less prevalent. Determining the risk factors that cause cancer or modeling the survival time of a patient once he/she is diagnosed with cancer using ANN is less common.

We are interested in utilizing ANN in survival time modeling of skin cancer (melanoma) patients. Soong et. al. [4] in 2010 developed a statistical model to predict the survival time of localized melanoma patients. They used the proportional hazard model developed by Cox [5], but the assumptions of hazard function proportionality may not be applicable to a different set of data. Moreover, they did not study the effect of interaction terms. Thus, applying ANN is more applicable and efficient, especially when the data does not satisfy Cox PH assumptions. One of the basic approaches in utilizing ANN in survival analysis is by classification, whether a patient will survive over a fixed time interval or not [6]. However, the latter classification method lacks the information about the survival probability function estimates. In 1995, P. Lapuerta et. al. proposed the use of multiple neural networks one for each time interval [7]. This model predicts the survival probability of each time period based on a neural network trained on the observations of the same time
period only. The pitfall of this approach is the large number of networks that will be trained if one studies the survival time over immense time intervals.

Another methods of ANN applied to survival times are by D. Faraggi and R. Simon in 1995 [8] and another by Machado in 1996 [9]. We consider in this dissertation the approach presented by E. Biganzoli et. al. in 1998 [10] named \textit{Partial Logistic Artificial Neural Network (PLANN)}. PLANN is a modification of an earlier study done by P. Ravdin and G. Clark in 1992 [11]. The second approach presented by Mani et. al. in 1999 [12] (we called it \textit{Discrete Hazard Artificial Neural Network (DHANN)}). In Chapter 3, we present a comparison between the two models and discuss relevant findings for predicting the survival time of melanoma patients.

\subsection{Skin Cancer (Melanoma)}

Melanoma is the most fatal type of skin cancer. It is ranked first in death among skin cancer diseases. Melanoma is a malignant tumor associated with skin cancer. If melanoma is detected at a late stage, it can spread to other parts of the body at the point of being a lethal form of cancer. More general information about melanoma can be found in (Markovic & et. Al, 2007) [13] and (Mackle & et. Al, 2009)[14]. Over the last decades, the incidence of melanoma has been rapidly increasing in the United States. It appears more in white populations than other races. According to clinical studies, risk factors of melanoma are but not limited to, ultraviolet light exposure, moles, light hair, freckling and family history of melanoma. Some of the statistical analyses done on the risk factors are shown in (Sara Gandini & et. Al, 2005, Luigi Naldi & et. al, 2000, and Eunyoung Cho & et al., 2005)[15–18].

We focus in the current study on survival time for the melanoma patients, which is the time it takes a patient once he/she is diagnosed with melanoma till death occurs. The study includes questioning the effect of several risk factors that we believe contribute to the survival time of melanoma patient using artificial neural network trained with Bayesian
inference. Using Bayesian inference has several advantages that we summarize in the next chapter; one of the advantages is that it helps in finding the relative importance of risk factors towards the outcome. Among those risk factors are: age at diagnosis, tumor thickness and tumor behaviour (invasive or non-invasive). Other factors are gender and sequence number (a number that indicates how many tumors the patient had prior to being diagnosed with melanoma). Seng-jaw Soong, et.al, 2010, developed an electronic prediction tool based on the AJCC melanoma database, to predict survival outcome of localized melanoma [4]. Other predictive models of survival for localized melanoma have been developed in the United States and other countries (Clark & et al, 1989, Mackie & et al, 1995, Bamhill et al, 1996, Schuchter & et al., 1996, Sahin & et al., 1997)[19–23]. Soong, et.al, used the Cox survival function model, which considers the survival time as a continuous random variable, where in fact most of survival times are recorded in discrete form as number of month or years. They used the same three risk factors in their analysis beside the primary melanoma site and primary tumor ulceration. We performed an initial study that address the concern of having a common model for male and female patient or should gender be treated separately [24]. We used discrete survival time methodology and came up to the conclusion that male and female patients should have separate models of predicting survival. Additionally, more variables need to be included with the four variables used by Soong for predicting survival time.

One of our goals in future is to model the incident of melanoma. Detecting the contributing variables to the incident of melanoma, in order to have a better understanding of these variables and to make decision regrading the prevention of be being diagnosed by melanoma.

1.3 Survival Analysis

Survival analysis is concerned with statistical methods that investigate time to event data, in which, the variable of interest is time until an event takes place (known as survival time).
This arises in various applications such as in clinical studies; time until death, time until disease incidence or recurrence. In engineering; time until machine malfunction, time until electronic component fails, etc. In some cases, one may be investigating time until the occurrence of more than one event, the statistical method is referred to as either a competing risks (for example: death from several causes) or recurrent events (for example: tumor recurrence after treatment). There exist several statistical methods that estimate the probability function that characterize the behavior of survival time (base line survival function) or conditional on some explanatory variables (risk factors). Some of these models treat survival time as a discrete random variable such as in Allison 1982 [25] or in Sharaf & Tsokos 2014 [24], others consider the distribution of the outcome (i.e., the time to event) to be specified in terms of unknown parameters (Kleinbaum & Klein, 2012 ) [26]. In the meantime, the non-parametric Kaplan-Meier (Kaplan & Meier, 1958) [27] method is the most widely used method to estimate time to event. In case of competing risks Kaplan Meier method tends to overestimate event rates (Southern, et al., 2006) [28]. Another popular method is the Cox proportional hazard model (Cox, 1972) [29], that can also estimate the survival time given specification of a set of explanatory variables (risk factors). However, Cox PH has some limitations similar to most of conventional statistical methods , which is a set of assumptions that need to be satisfied by the data before applying Cox PH to it.

1.3.1 Discrete Survival Time

In most data, survival time is recorded in discrete form either as number of months or number of years. Another methodology for survival analysis is used, that consider time as discrete. Haiyi Xie, et.al (2003) [30] , summarized the advantages of using discrete-time survival analysis, as first, useful for many longitudinal studies in clinical settings where data are often collected at discrete time periods. Secondly, it facilitates the examination of the shape of the hazard function. Third, is simple and convenient to use, because it is a modification of the logistic regression model. Lastly, and most important advantage,
one can include the time-varying covariates easily in the model. After Cox presented the
discrete time survival model, two basic versions of logistic models were introduced, the
ordinal version and the dichotomous version. The dichotomous version (Allison,1982,
Singer & Willett, 1993, Xie, Mchugo, Drake, and Sengupta, 2003)[25, 30, 31] where each
survival time is represented as a set of indicators of whether or not an individual failed at
each time point until a person either experiences the event or is censored.

A discrete survival time method was proposed in [25], and in [31]. The method starts by
dividing the continuous time into an infinite sequence of contiguous time \((0, t_1), (t_1, t_2), \ldots, (t_{k-1}, t_k), \ldots \) and so on. Let \(k\) represent the number of time intervals. For example, consider a
time variable recorded in months over 20 years period. Then, time is divided into 20 inter-
vals each consists of 12 months; \((1, 12), (12, 24), \ldots, (228, 240)\). So, If a subject survival time
is 7 months, then this subject’s event is classified as taking place during the 1st time inter-
val, if another subject survival time is 50 months then is classified as taking place during
the 5th time interval.

To estimate the survival function, we start with the discrete-time hazard model given by:

\[
h_{ik} = \frac{1}{1 + \exp\{- (\alpha_1 T_{1ik} + \ldots + \alpha_J T_{Jik}) - (\beta_1 X_{1ik} + \ldots + \beta_p X_{pik})\}} \tag{1.1}
\]

where, \([T_{1ik}, T_{2ik}, \ldots, T_{Jik}]\) are sequence of dummy variables, with values
\([t_{1ik}, t_{2ik}, \ldots, t_{Jik}]\) indexing time periods, where J refers to the last time period observed
for any individual in the sample. If individual i was observed (experienced the event or
ensored) in fourth period, then \(J = 4\), the time periods dummy variables are defined iden-
tically for each individual; \(t_{1ik} = 1\), when \(j = 1\) and \(0\) when \(j\) takes any other value. The
coefficients \((\alpha_1, \alpha_2, \ldots, \alpha_J)\) Act as the intercept parameters for the baseline hazard in each
time period, and the coefficients \((\beta_1, \beta_2, \ldots, \beta_p)\) describes the effect of the predictors on
the baseline hazard in the logit scale. Singer and Willett, (1993) discussed briefly the pro-
cedures to construct the likelihood function (in terms of the discrete hazard function) used
to estimates the latter intercepts and slope parameters. The likelihood function presented
by Singer and Willett is given by:

\[
L = \prod_{i=1}^{n} \prod_{k=1}^{j_i} h_{ik}^{y_{ik}} \{1 - h_{ik}\}^{(1-y_{ik})} \tag{1.2}
\]

Where, \( Y_{ik} \) is a sequence of dummy variables that records the event history for subject \( i \), whose values are defined as:

\[
Y_{ik} = \begin{cases} 
1 & \text{if the } i \text{th subject experinced the event in period } k \\
0 & \text{if the } i \text{th subject did not experince the event in period } k
\end{cases}
\]

The likelihood function in (1.2) is identical to the likelihood function to a sequence of \( N = (k_1 + k_2 + + k_n) \) independent Bernoulli trials with parameters \( h_{ik} \). Using results by Allison (1982), we can consider the \( Y_{ik} \) values as the outcome variable in a logistic regression analysis, which provides a simple model to obtain the maximum likelihood estimate rather than finding the solution by maximizing equation (1.2).

For discrete event history data, each record consists of the information for one subject such as; survival time, age and whether or not the subject time is censored. In order to apply the logistic model discussed previously, the data need to be converted into new person-period data, in which each subject will have multiple records, one per time period of observation. As shown by Singer and Willett, the new person-period data will contain the information about the \( k^{th} \) time period as follows:

- The time indicators: the set of dummy variables \([T_{1ik}, T_{2ik}, \cdots, T_{Jik}]\).
- The predictors. The covariates that are under study, where we have the ability of using the time-varying covariates have values that differ from time period to time period.
- The event indicator (response variable in the logistic model). This variable records whether the event of interest occurred in period \( j \) or not. The variable takes value 1 if the event occurred, takes 0 if did not.
Table 1: Information of three melanoma patients

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Survival Time</th>
<th>Age Diagnosis</th>
<th>Tumor</th>
<th>Ext. of Tumor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>49</td>
<td>84</td>
<td>120</td>
<td>10</td>
</tr>
<tr>
<td>2</td>
<td>3</td>
<td>66</td>
<td>230</td>
<td>30</td>
</tr>
<tr>
<td>3</td>
<td>86</td>
<td>61</td>
<td>134</td>
<td>30</td>
</tr>
</tbody>
</table>

For illustration purposes consider a record of three melanoma patients as shown in table 1. By picking first subject we can see that his survival time is equal to 49 months, which means that this subject information will repeated for 5 time intervals whether the event took place or the subject is censored.

In the new data setting the first patient will have five records, record for corresponding to every time period (from the first to the fifth). The event indicator variable will take 0 for the first four records and 1 in the fifth record where the event took place. The transformation of these 3 subjects information in presented on table 2. The variables $D_1, D_2, \cdots, D_T$ represents the $T$ dummy variables of time intervals. Each row shows subject $i$ information during each time interval until the event occurs or he/she is censored. The first column in Table 2 (Indc.) represents the indicator variable which takes 0 if the event did not take place during the current time period interval, and takes 1 if the event occurs during the time period interval. The setting of the data shown in table 2 can allow us to use time varying covariates easily, which is one of the advantages of using discrete time survival method. On the other hand, repeating subject’s information causes redundancy forcing variables to be correlated (which also is not practical with today’s huge amount of data). As we will show in chapter three that the likelihood function in (1.2) can be maximized using artificial neural network (PLANN) to predict (1.1). As readers will find on chapter 3 that the ANN method based on the discrete time method (explained above), does not perform better compared to the second method DHANN.
Table 2: Information of three melanoma patients

<table>
<thead>
<tr>
<th>Indc.</th>
<th>ID</th>
<th>ST</th>
<th>Ag</th>
<th>TS</th>
<th>D_1</th>
<th>D_2</th>
<th>D_3</th>
<th>D_4</th>
<th>D_5</th>
<th>D_6</th>
<th>D_7</th>
<th>D_8</th>
<th>...</th>
<th>D_T</th>
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<tr>
<td>0</td>
<td>1</td>
<td>49</td>
<td>84</td>
<td>120</td>
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<td>0</td>
<td>1</td>
<td>49</td>
<td>84</td>
<td>120</td>
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The rest of the dissertation goes as follows, on next chapter we highlight some parts of neural network theory used throughout the remainder of the dissertation. To make the reader familiar with the terms used, with neural network structures, and neural network training algorithms. On Chapter three we present a comparison between PLANN and DHANN on data collected from SEER. We discuss the procedure of developing both models and illustrate how different neural network structures can be used for same purpose. Additionally we talk about significant findings regarding the survival time of male and female melanoma patients.

In Chapter four we introduce new artificial neural network for modeling survival time in the presence of competing risk. This new neural network is an update on DHANN.
Moreover, we study two different Bayesian learning methods for ANN on the DHANN for competing risks, and propose a new method for Bayesian learning. In Chapter five, we study a different structure of neural network used in forecasting time series data. We examine Bayesian learning techniques on predicting the monthly carbon dioxide emission in the atmosphere in the United States.
Artificial neural networks (ANN) are inspired by the human nervous system. In 1943, Warren McCulloch and Walter Pitts [32] presented the first artificial neurons used to simulate human biological nervous system. Since that time ANN were developed in several fields independently. It is now an important tool of artificial intelligence and are used extensively in machine learning. In statistics, ANN is considered as a non-linear modeling tool, that possesses high accuracy in prediction and a strong identifier of complex patterns within a set of data.

A neural network consists of several layers of interconnected units called neurons. The way those neurons are connected differs according to the application ANN is used for. Fig-
Figure 1, shows an ANN with three layers: Input layer, hidden layer and output layer. Each unit in the input layer is directly connected to every unit in the hidden layers. Those connections are done by the weights (which acts as the synaptic connection in the human nervous system). A definition of Neural Network by D. Kriesel [33] as follows: “A neural network is a sorted triple \((N, V, \omega)\) with two sets \(N, V\) and a function \(\omega\), where \(N\) is the set of neurons and \(V\) a set \(\{(i, j) \mid i, j \in N\}\) whose elements are called connections between neuron \(i\) and neuron \(j\)”. The set of weights acts as the connection that transfer data between neurons. Each neuron has main two functions; one function to transfer the input of the neuron and other responsible for the output of the neuron. The first function normally called combination function or sometimes propagation function \((W_j)\) transfer the inputs of the neuron to one value and the most popular function is the weighted sum which for a given neuron \(j\) is given by:

\[
W_j = \sum_{i=1}^{I} a_i * \omega_i
\]

(2.1)

where \(a_i\) is the output of the previous layer, \(I\): is the number of neurons in the previous layer.

The second function is called the output function or sometimes the activation function. There exist several functions used as activation functions, some of the popular functions are linear function, sigmoid (logistic function) given by:

\[
O_i = \frac{1}{1 + \exp^{-x}}
\]

(2.2)

and the hyperbolic tangent given by:

\[
O_i = \tanh x = \frac{1 - \exp^{-2x}}{1 + \exp^{-2x}}
\]

(2.3)

The choice of which function to use depends on the application ANN is used for. For instance, if ANN is used for linear regression problem then the proper activation function
for the output neuron would be the linear function. If ANN is used for predicting Hazard probability (as we will show next chapter) then the logistic function is used for hidden and output neurons.

Due to wide areas and fields which ANN is utilized in, there exist several ANN architecture (Structure or topology). In this chapter we are going to give a brief introduction on two different ANN architecture. In addition discuss the training algorithms and methods used in the training purpose of ANN.

2.1 Types of Artificial Neural Networks

In this section we are going to explore some of the popular networks types in terms of it’s design (topology/architecture). We will start with the most popular and widely used design, the Feedforward Network.

2.1.1 Feedforward Networks

Feedforward ANN is the most widely used network design in classification and regression problems. Feedforward networks typically consist of one input layer, $H$ hidden layers ($H$ refers to number of hidden layers), and one output layer. A Feedforward network with one hidden layer is called three layer feedforward network, which is represented by Figure 1. On the left side is the input layer, representing the starting point of data flow. Input layer represents covariates(input variables). Neurons of the input layer feed every neuron in the hidden layer (not opposite), then data are processed by each neuron of the hidden layer and then feed to the third layer (output layer). In some case neurons in the first layer can feed neurons in third or above layers (known as shortcut connections), but flow of data/information remains same going from left to right.

To illustrate how feedforward networks can be used to model in linear regression will consider the next model:
\[ Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 \] (2.4)

The previous model can be estimated using two layer feedforward network(Figure 2). One input layer representing two independent variables \( X_1 \) and \( X_2 \) and one output layer with one neuron representing the response variable \( Y \).

![Two Layer Feedforward Network](image)

**Figure 2.:** Two Layer Feedforward Network for estimating the Linear regression model in equation 2.4

First layer of the neural network in Figure 2, consist of two nodes for the two independent variables \( X_1 \) and \( X_2 \) and the orange colored node for the bais term. Then we have one neuron in the output layer with weighted sum as combination function and linear function as the activation function. The output of such network is given by:

\[ Y = f(b + \omega_1 X_1 + \omega_2 X_2) \] (2.5)

where \( f() \) is a linear function. equation 2.5 is equivalent to equation 2.4, and how the parameters \( b, \omega_1, \) and \( \omega_2 \) are estimated will be discussed later in this chapter. More on how we can use Feedforward networks in modeling linear and generalized linear regression can be found in C. Bishop book [34].
2.1.2 Recurrent (Recursive) Networks

Neurons in a Feedforward networks feeds only neurons of the next layers. Other types of networks are formed such that neurons are feeding themselves or neurons of the preceding layers or neurons of same layer, such networks are called recurrent networks. There are several types of recurrent networks, such as the most two popular types are Hopfield neural networks [35] and Bi-directional networks [36]. In some cases recurrent networks do not have explicit definition of input and output layers (as in figure 3).

Figure 3.: Recurrent neural network, where doted lines represent indirect recurrences

Speech recognition and language learning are two of popular application of recurrent networks [37–39]. Other important use of recurrent networks (Known as Real-Time Recurrent Networks) is in the modeling of time series data as we will show later in chapter five. Figure 4 displays feedforward network with output going back as one of the inputs which is used in modeling non-linear auto-regressive models.

In this dissertation we will see the utilization of feedforward neural network and recurrent neural networks (the one in Figure 4) in performing statistical modeling such as survival analysis and time series analysis.
2.2 Learning method of Neural Networks

Neural networks are inspired by the human nervous system making them learn in the same way as human learns, which is by example. This type of learning is called supervised learning, that is, neural network is given set of examples (data). Consider some data \((X, t)\), where \(X\) is a matrix of information (covariates) and \(t\) is vector of targets, response variable, that acts as a teacher forcing neural network to find the best approximated output to the target values. Another type of learning for neural network is called unsupervised learning, in which a network decide on its own what would be the best output fit for a set of data without external help [40].

In this section we summarize the idea of training neural network from C. Bishop books “Neural Network for Pattern Recognition” [41] and “Pattern Recognition and Machine Learning” [34]. To train a neural network we need to estimate the weights that minimize error function chosen for the certain task of the neural network. One of the most popular algorithms to do so is called back-propagation, and it took this name as it is based on propagating the errors backward from output neurons to neurons of first layer. For example, consider similar structure of neural network as in Figure 1, that will be used in a linear
regression problem. We have data in the form of set of covariates and targets \( \{X_s, t_s\} \), where \( s = 1, 2, \ldots, N \) number of observations. The network activation function for the output unit in this case would be the linear (identity) function, and therefore the best choice of error function \( E \) will be the mean square error given by:

\[
E = \frac{1}{2} \sum_{s=1}^{N} (y(x_s, \omega) - t_s)^2
\]  

(2.6)

where \( y(x_s, \omega) \) is the network output for observation \( s \), which can be expressed in terms of probability. For instance (in regression problem), if our target follows a Gaussian distribution then network output can be expressed as:

\[
p(t \mid x, \omega) \approx \mathcal{N}(t \mid y(x, \omega), \frac{1}{\beta})
\]  

(2.7)

where \( \beta \) is the inverse variance of the Gaussian (noise) distribution. From equation (2.7) one can find the likelihood function for the whole \( N \) observations on the form \( \{X_s, t_s\} \) as:

\[
p(t \mid X, \omega, \beta) = \prod_{s=1}^{N} p(t_s \mid x_s, \omega, \beta)
\]  

(2.8)

So one can estimate network weights \( \omega \) by maximizing the negative logarithm of (2.8), which in neural networks literature is equivalent to minimizing the error function in (2.6). The most popular algorithm used in minimizing the error function is \textit{back-propagation} that consists mainly of two parts.

\subsection*{2.2.1 Evaluating derivatives of Error function}

The first part of training neural networks through back-propagation is to evaluate the derivatives of the error function with respect to the network weights.

To evaluate the derivatives we will introduce some notations first:

- The first function for a given neuron \( j \) as we explained earlier is the weighted sum given
by:

\[ a_j = \sum_i \omega_{ji} z_i \tag{2.9} \]

where \( \omega_{ji} \) is the weight passing information from neuron \( i \) to neuron \( j \). And, \( z_i \) is the output of neuron \( i \) of the preceding layer.

- The output of each neuron \( z_i \) is computed by the activation function as:

\[ z_i = h(a_j) \tag{2.10} \]

where \( h() \) is the activation function (as mentioned earlier it can be; identity function, sigmoid or softmax).

- Recalling equation (2.6), the error function is a function of the network weights through the summed inputs \( a_j \), so by applying the chain rule we obtain:

\[ \frac{\partial E}{\partial \omega_{ji}} = \frac{\partial E}{\partial a_j} \frac{\partial a_j}{\partial \omega_{ji}} \tag{2.11} \]

By using (2.9) we obtain \( \frac{\partial a_j}{\partial \omega_{ji}} = z_j \), then (2.11) is simplified to:

\[ \frac{\partial E}{\partial \omega_{ji}} = \delta_j z_j \tag{2.12} \]

where \( \delta_j \) is given by:

\[ \delta_j = h'(a_t) \frac{\partial E}{\partial y_t} \tag{2.13} \]

for neurons in the output layer. Where \( t \) is number of neurons in output unit, and \( y_t \) is the output of neurons in the output layer. The neurons in hidden layers \( \delta_j \) is given by:

\[ \delta_j = h'(a_t) \sum_t \omega_{tj} \delta_t \tag{2.14} \]
equation (2.14) represents the back-propagation formula as the error derivatives are propagation backward from neurons of output layer to neurons of hidden layers.

Once the derivatives of (2.12) are calculated then it is the time to turn these derivatives to learning rule to change network weights. Updating the weights can be done after each observation (called on-line learning) or after all data have been passed through the network (batch learning) by summing the derivatives over all observations $N$

$$
\frac{\partial E}{\partial \omega_{ji}} = \sum_{s=1}^{N} \frac{\partial E_s}{\partial \omega_{ji}} \tag{2.15}
$$

### 2.2.2 Optimization Algorithm

The second part of training neural networks by the back-propagation algorithm is to use the derivatives of the error function (2.12) and turn them to learning rule to adjust the network weights. This is done by involving a non-linear optimization algorithm like gradient descent or quasi-Newton method among others. We are going to discuss the quasi-Newton method as it was the one used in training our neural networks in next chapter. Our goal is to find weight vector $\omega$ that minimize the error function $E(\omega)$. Imagine this problem graphically as error surface sitting on weight space, so we need to find the gradient of the error in the weight space that will lead us to minimum point in the space ($\nabla E = 0$). The gradient of error using quasi-Newton method is given by:

$$
graident = \nabla E = \mathbf{H}(\mathbf{w} - \mathbf{w}^*) \tag{2.16}$$

where $\mathbf{H}$ is the hessian matrix that can be obtained by the second derivatives of the error function with respect to the weights ($\mathbf{H} = \frac{\partial^2 E}{\partial \omega_i \partial \omega_j}$). And, the weight vector $\mathbf{w}^*$ satisfies

$$
\mathbf{w}^* = \mathbf{w} - \mathbf{H}^{-1} \nabla E \tag{2.17}
$$
where $H^{-1} \nabla E$ is known as Newton’s direction/step that automatically determines the size and the direction of the step towards the error minimum. Since the direct calculation of the Hessian matrix is computationally expensive (which is known by Newton’s method), the alternative is by building up an approximation to the inverse Hessian over a number of steps (which is known by quasi-Newton method) [41]. The approximation of the Hessian matrix involves generating a matrix $G^{(k)}$ at step $k$ as follows:

$$G^{k+1} = G^{(k)} + \frac{pp^T}{p^Tv} - \frac{(G^{(k)}v)v^T G^{(k)}}{v^T G^{(k)}v} + (v^T G^{(k)}v) uu^T \quad (2.18)$$

where we define:

$$p = w^{k+1} - w^{(k)} \quad (2.19)$$

$$v = g^{(k+1)} - g^{(k)} \quad (2.20)$$

g: is the gradient in (2.16). And,

$$u = \frac{p}{p^Tv} - \frac{G^{(k)}v}{v^T G^{(k)}v} \quad (2.21)$$

initial choice of $G$ would be the identity matrix then we can replace Newton’s step $-Hg$ in (2.17) by $-Gg$. However, as Bishop [41] mentioned full Newton’s step in (2.17) may take the search outside the approximation valid area and thus the use of line-search algorithm is applied and (2.17) becomes:

$$w^{k+1} = w^{(k)} + \varsigma^{(k)} G^{(k)} g^{(T)} \quad (2.22)$$

where $\varsigma^{(k)}$ is found by line minimization.

The draw back of quasi-Newton method is the need of more storage to update matrix $G$ as the number of network weights increases. Moreover, quasi-Newton method follows
always a descent direction in the space so it can be trapped in local minimum as going out from that point requires increase in the error function which is not allowed. Neural networks popular draw back in general is the excess time it may take to approximate function based on certain case not only because of the optimization algorithm used but also if weights were allowed to take large values during training, which is similar to the performance of human brains when large information are processed through it in a certain time. In another words, If network weights are allowed to take on large values, conversion of the optimization algorithm becomes slower. The solution for the previous issue is solved by introducing additional term called weight decay $\alpha$, that controls the variance of the weights and penalize large weight values. Weight decay is based on Bayesian method that will be discussed in the following section. The Error function for networks working with weight decay becomes:

$$E = E_D + \alpha E_W$$

(2.23)

where $E_D$ is the error due to approximation of network outcomes to targets given by (2.6) and

$$E_W = \frac{1}{2} \sum_{i=1}^{W} \omega_i^2$$

(2.24)

### 2.3 Bayesian Neural Networks

In this section we will discuss the use of Bayesian inference methods in neural networks. David MacKay [42–44] proposed the use of Bayesian in neural network and after him, R. Neal [45] updated his work, so more details can be found in their papers. The use of Bayesian in neural networks have several advantages:

1. Helps in determining the optimal values for the regularization parameters (like weight decay) by estimating them online from the training data.

2. Regularization can be given a natural interpretation in the Bayesian framework.
3. Parameter uncertainty can be accounted for in the network predictions and hence over-fitting is not an issue.

4. Helps in comparing between several neural network architectures using training data only.

5. The relative importance of input variables can be determined using automatic relevance determination [45, 46]

In the previous section we talked about training neural network through the maximum likelihood. In Bayesian setting we find the conditional probability of network outputs through posterior distribution of the network weights. Through Bayesian inference the posterior distribution of a vector of weights $\omega$ given a set of data $\mathcal{D}$ (where $\mathcal{D}$ contains both set of variables information $\mathbf{X}$ and its corresponding targets $\mathbf{t}$) is given by:

$$
p(\omega | \mathcal{D}) = \frac{p(\mathcal{D} | \omega) \, p(\omega)}{p(\mathcal{D})} \tag{2.25}
$$

where $p(\mathcal{D} | \omega)$ is the data likelihood, $p(\omega)$ is the prior distribution of the weights, and $p(\mathcal{D})$ in the normalization factor (also called the evidence). So once the posterior distribution in (2.25) is calculated then network prediction output for a given input $\mathbf{x}^\prime$ can be obtained by:

$$
p(y | \mathbf{x}^\prime, \mathcal{D}) = \int p(y | \mathbf{x}^\prime, \omega) p(\omega | \mathcal{D}) \tag{2.26}
$$

as we know in Bayesian, the integral of (2.26) in most cases is analytically intractable and is approximated or we use simulations to evaluate it. Two methods for evaluating the integral of (2.26) will be discussed, the first is the evidence procedure [43] and the second uses the Hybrid Monte Carlo [45]. Before we discuss the two methods we will talk first about the choice of prior of the network weights.
2.3.1 Neural Network Weights: Choice of Priors

The choice of priors rises from the need of having small weights so we can obtain smooth network performance and to avoid mapping with large curvature. This need for small weights suggest a Gaussian prior with zero mean, given by:

\[ p(\omega) = \frac{1}{Z_W(\alpha)} \exp\left( -\frac{\alpha}{2} \sum_{i=1}^{W} \omega_i^2 \right) \]  

(2.27)

where \( \alpha \) is the inverse variance of the weight distribution and \( Z_W(\alpha) \) is the normalizing constant given by:

\[ Z_W(\alpha) = \left( \frac{2\pi}{\alpha} \right)^{\frac{W}{2}} \]  

(2.28)

after ignoring the normalizing constant of (2.27) and taking the negative log, we will find that the prior is equivalent to the error term of (2.24). This error term regularizes the weights by penalizing over large values. The weights distribution is assumed to be a Gaussian distribution with mean zero and variance \( \frac{1}{\alpha} \), so the value of the hyperparameter \( \alpha \) controls the weights of fanning out from each input variable and hence large alpha values means small variance so inputs are of less relevant to output. This approach can be generalized by assuming different hyperparameters for each input variable. This procedure is known as automatic relevance determination (ARD). In the following we will discuss the two methods for evaluating the integral of (2.26)

2.3.2 The Evidence Procedure

The evidence procedure introduced by MacKay [43] as an iterative algorithm for determining optimal values for network weights and hyperparameters. It is not considered as fully Bayesian as it searches for optimal values rather than integrating out the whole vector of weights in (2.26). By introducing the hyperparameters, the posterior probability distribu-
tion of the network weights is given by:

\[ p(\omega \mid \mathcal{D}) = \int \int p(\omega \mid \alpha, \beta, \mathcal{D}) p(\alpha, \beta \mid \mathcal{D}) \, d\alpha \, d\beta \]  

(2.29)

To evaluate the latter integral which is at least double integral and usually will be more (depending on the number of different hyperparameters), the evidence procedure approximate the latter integral to:

\[ p(\omega \mid \mathcal{D}) \approx p(\omega \mid \alpha_{MP}, \beta_{MP}, \mathcal{D}) \]  

(2.30)

where \( \alpha_{MP} \) and \( \beta_{MP} \) are the most probable values of the hyperparameters. These values are considered to be the values where the posterior density of the hyperparameters is peaked around. Which means that first we need to estimate the most probable hyperparameters, then next step involves integrating the posterior distribution of the weights (2.30) to obtain network outputs. So, first step in the evidence procedure is to evaluate the posterior distribution of hyperparameters by approximating it to the most probable values of hyperparameters. Using Bayesian inference, the posterior distribution of the hyperparameters is given by:

\[ p(\alpha, \beta \mid \mathcal{D}) = \frac{p(\mathcal{D} \mid \alpha, \beta) \, p(\alpha, \beta)}{p(\mathcal{D})} \]  

(2.31)

where \( p(\mathcal{D}) \) can be ignored since in the evidence procedure we only need the peaks this density. \( p(\alpha, \beta) \) is the prior density of the hyperparameters in which there exist a wide range of choices for the prior based on the applied problem we will be using the neural network on. For the meantime we will assume uniform distribution as the prior of the hyperparameters so it can be ignored in the rest of the analysis. This leave for us the likelihood \( p(\mathcal{D} \mid \alpha, \beta) \) which we need to maximize by finding the peek values of \( \alpha \) and \( \beta \). \( p(\mathcal{D} \mid \alpha, \beta) \) is also known as the evidence that can be used to compare between different
neural network structures. The evidence is given by:

\[
p(D \mid \alpha, \beta) = \frac{1}{Z_D(\beta)} \frac{1}{Z_W(\alpha)} \int \exp(-S(\omega))d\omega
\]  

(2.32)

and to find the values of \(\alpha\) and \(\beta\) we take partial derivative for the log of (2.32). In the evidence procedure the Hessian matrix \(A\) of the error function \(S(\omega)\) is evaluated at \(A = H + \alpha I\) (2.34), so by differentiating log of (2.32) with respect to \(\alpha\) and equating to zero we get:

\[
\alpha_{new} = \frac{\gamma}{2E_W}
\]  

(2.33)

where \(\gamma = \sum_{i=1}^{W} \frac{\alpha}{\lambda_i + \alpha}\), so that \(\lambda_i + \alpha\) are eigenvalues of \(A\). Following same for \(\beta\) we obtain:

\[
\beta_{new} = \frac{N - \gamma}{2E_D}
\]  

(2.34)

Now it is turn to find the weights posterior distribution, recall the goal in Feedforward networks is to find the set of weights that minimize the error function. MacKay used another approximation for the error function (also known as the cost function) as a second Taylor-expansion that have local minimum at \(\omega_{MP}\). The expansion of error function \(S(\omega)\) is given by:

\[
S(\omega) \approx S(\omega_{MP}) + \frac{1}{2}(\omega - \omega_{MP})^T A(\omega - \omega_{MP})
\]  

(2.35)

where \(A\) is the Hessian matrix of the total error function:

\[
A = \nabla^2 S(\omega_{MP}) = \beta H + \alpha I
\]  

(2.36)

where \(H = \nabla^2 E_D(\omega_{MP})\), and since it is preferable that the error \(S(\omega)\) to have Gaussian distribution, it follows that the posterior distribution of (2.30) is:

\[
p(\omega \mid D) = \frac{1}{Z_S} \exp \left(-S(\omega_{MP}) - \frac{1}{2}(\omega - \omega_{MP})^T A(\omega - \omega_{MP})\right)
\]  

(2.37)
where $Z_S$ is the normalizing factor of the Gaussian distribution for the error function.

Finally we can summarize the evidence procedure in the following algorithm:

1. Choose initial values for hyperparameters $\alpha$ and $\beta$, and network weights and bias.

2. Train neural network with appropriate optimization algorithm to minimize the error function $S(\omega)$.

3. Then using the Gaussian approximation for the hyperparameters, new values of hyperparameters can be found using (2.33) and (2.34).

4. Repeat steps 2 and 3 until convergence is reached.

Note that evidence procedure can be implemented easily but choosing different initial $\alpha$ values for each of the input variables.

### 2.3.3 Hybrid Monte Carlo Method for Neural Networks

The idea that Mackay presented for applying Bayesian inference in learning of neural network was improved by Neal [45]. The origin of hybrid Monte Carlo sampling returns to Duane [47]. Neal enhanced Bayesian neural networks by calculating the integral of (2.26) using Hybrid Monte Carlo sampling, to sample from the weights posterior distribution. In Hybrid Monte Carlo method the integral in (2.26) is approximated by:

$$p(y \mid x, D) \simeq \frac{1}{N} \sum_{c=1}^{C} p(y \mid x, \omega_c)$$  \hspace{1cm} (2.38)

where $\{\omega_c\}$ represent a sample generated from the posterior distribution of the weights.

The hybrid Monte Carlo combines stochastic sampling techniques with the Metropolis-Hastings algorithm. Since in neural network we can easily evaluate the gradient information using the back-propagation technique, this gradient information can be used to direct the direction of MH algorithm. Though Neal have shown that his approach is better than
using the evidence procedure but still have some back flaws. First, Neal uses initial val-
ues for the hyperparameters that does not change with respect to the data, and as it is well
known that the Bayesian prediction is as good as the choice of the prior. Unless we are sure
about the priors belief, using the Hybrid Monte Carlo sampling by Neal will require a huge
number of trails to test for different values of hyperparameters. Secondly, it is highly timely
consuming method, especially with today’s massive data used to predict certain problem.

In Chapter four we shall introduce a solution for the latter problem by applying a mixture
of the evidence procedure and the Hybrid Monte Carlo sampling. Furthermore, we are
going to show in chapter four that we did not only obtain the lowest error by using this
new approach, but we also were able to obtain neural network models in much shorter
computational time.
Chapter 3
Prediction of Survival Time for Melanoma Patients Using Artificial Neural Network

In Chapter 1, we mentioned several models that were introduced to model survival time using artificial neural networks. However, in literature we did not see any comparisons among these models. In this chapter a comparison between two of these methods that utilize artificial neural network in predicting survival time is performed. The first method is called the partial logistic artificial neural network (PLANN), [10], and the second method we call it discrete hazard artificial neural network (DHANN), [12]. The comparison is done using real world data on skin cancer patient specifically malignant melanoma, the most deadly type of skin cancer. Also, we study the difference between the survival time of male and female melanoma patients.

3.1 Melanoma Patient Data

In the present study we use 130,006 patients diagnosed with melanoma between the years 2000 to 2009 in the US. Data accumulated from 13 registers of the Surveillance, Epidemiology, and end results program (SEER) [48]. We filter out this large data set to contain only consummate information with respect to the patient’s age at diagnosis, tumor thickness, stage of cancer and ulceration. Soong et al. [4] in 2010 used these four variables, but their study did not consider the difference between male and female survivals. We found that there exists a significant difference between the median survival time of males and females based on a 5% level of significance using the Kruskal-Wallis test. Thus, studying the effect of gender on survival time by making one model for both males and females is
not statistically correct, as the survival time for male and females does not have the same probability distribution. Figure 5, below, represents a schematic diagram of the distribution of the complete data with respect to gender and cancer stage. We train neural network model separately for males and females.

Figure 5: Distribution of complete information of melanoma patients

Figure 5, also includes the total number of patients with complete information who were diagnosed with melanoma between the years 2000 to 2009. Patients were either alive at the end of the 10 year period (censored) or lost follow up during the ten year interval (censored). In addition, patients who died because of melanoma (uncensored) and percentage of censored patients split as 95% of male patients were censored and 98.1% of female patients were censored. We omitted patients with incomplete information. For training purposes, each of the gender category data was divided into six groups; five will be used in the cross validation technique to train and validate the neural network model, while the last group is used for prediction and accomplishing the comparison between the two modeling approaches. More details about the cross validation will follow in the next section. In our modeling procedure, we used age at diagnosis and tumor thickness (in millimeter) as
quantitative variables along with three dummy variables representing stage 1 (referring to a localized tumor), stage 2 (referring to a regional tumor), and stage 3 (referring to a distant tumor). The base level is referred to as in situ tumor.

3.2 Methodology

In this section we are going to discuss the two method PLANN and DHANN. Both of the two methods used three layer Feedforward neural network.

3.2.1 Partial Logistic Artificial Neural Network (PLANN)

Partial logistic artificial neural network (PLANN) is the approach that was introduced by E. Biganzoli et. al. [10]. PLANN is a three layer feed forward artificial neural network with one output unit in the output layer. The activation function used in the hidden and output layer is the sigmoid (logistic) function given by (2.2). PLANN estimates the conditional hazard function that is based on the discrete survival method (discussed in chapter one). Recall that the discrete hazard probability function for time period $k$ given a vector of covariates $x_i$ is given by

$$h_k(x_i, t_k) = \frac{1}{1 + \exp(-\alpha_k t_k - \beta^T x_i)}$$

(3.1)

where $t_k$ represents the time input for time period $k$. To estimate the conditional hazard in (3.1), PLANN uses three layer Feedforward network with an activation function for the hidden and output layers given by (2.1). The output of the network with an $H$ number of hidden units is given by:

$$\hat{h}(x_i, t_k) = f\left[ b + \sum_{h=1}^{H} \omega_h f(a_h + \sum_{p=1}^{P} \omega_{ph} x_{ip}) \right]$$

(3.2)
where $\omega_{ph}$ and $\omega_h$ are the weights of the ANN to be estimated for the first layer and second layer respectively, also $a_h$ and $b$ are the weights for the bias connection with the hidden units and with the output unit respectively. The target of this network is the censoring indicator $c_{ik}$, which takes the value one if the event occurred for subject $i$, and zero otherwise. The cost function used in PLANN is the cross-entropy function which is appropriate for binary classification problems, [41]. The weights of PLANN can be estimated by minimizing the cost function given by

$$E = -\sum_{i=1}^{n} \sum_{k=1}^{k_i} \left\{ c_{ik} \log \left[ \hat{h}(x_i, t_k) \right] + (1 - c_{ik}) \log \left[ 1 - \hat{h}(x_i, t_k) \right] \right\} \quad (3.3)$$

Once the network weights are estimated, the monotone survival probabilities can be easily found by converting the discrete hazard rate estimates obtained from the network output by using the following equation:

$$S(t_k) = \prod_{l=1}^{k} \{1 - h(t_l)\} \quad (3.4)$$

The advantage of this approach is that the time dependent covariates can easily be introduced in the model, as the individual records are available for each time period. However, for large data sets or studies conducted over a long period of time this approach is inaccessible due to the immense number of replication requisites, [49]. Figure 6 on next page, shows the architecture of the PLANN introduced by Biganzoli. The first layer contains the bias and one node for the time period and the rest of the nodes for the covariates. PLANN uses one input for the time to estimate smooth discrete hazard rates. However, we have used 10 nodes for the time (one for each time period) to be able to compare it with the second method (DHANN).
Figure 6.: Feedforward ANN for partial logistic artificial neural network, with three layers. The input layer has p covariates and hidden layer with H hidden units and one output unit in the output layer. Activation function used in both hidden and output layer is the logistic function (2.1).

### 3.2.2 Discrete Hazard Artificial Neural Network

Mani [12] developed an approach that predicts the hazard discrete function similar to an approach by Street [50], that predicts the survival function using a neural network with K outputs, where K is the number of time periods. He trained his network utilizing a target vector derived by Kaplan-Meier survival curves [51]. Mani used same neural architecture as Street, but rather estimate the hazard function not the survival function. In order to estimate the hazard function, each individual or subject would have a training vector (1 by K) target of hazard probabilities $h_{ik}$ as follows:

$$
h_{ik} = \begin{cases} 
0 & \text{for } 1 \leq k \leq K \\
1 & \text{for } t \leq k \leq K \text{ and event } = 1 \\
\frac{r_k}{n_k} & \text{for } t \leq k \leq K \text{ and event } = 0
\end{cases}
$$
here, $h_{ik} = 0$ for each time period patient $i$ survived. $h_{ik} = 1$ from time interval $t$ to $K$ if patient died because of melanoma at duration $t$ within the study time. And, for those patients who lost follow up during the study of duration $t < K$ their hazards are equal to the ratio $\frac{r_k}{n_k}$, which is the Kaplan-Meier hazard estimate for time interval $k$. Also, $r_k$ is the number of patients who died because of melanoma in time period $k$, and $n_k$ is the number of patients that are at risk in time interval $k$. For training the neural network, Mani used the logistic sigmoid function given by (2.1). The network weights are estimated by minimizing the cost function, which is the cross entropy function. Figure 7, shows the neural network architecture utilized by DHANN. The number of units $p$ of the input layer are equivalent to the number of independent variables or risk factors. The $K$ output units of the output layer learns to estimate the hazard probability of each individual. Once the ANN is trained and the hazard estimates are predicted, we convert those hazard estimates to the survival estimates by using (3.4), for each method. We have trained the weights of the ANN for both methods using the quasi-newton algorithm discussed in chapter 2.

Figure 7.: DHANN: Three layer network with $K$ output units in the output layer where $K$ is equal to the number of time intervals.
3.3 Model Selection

Now, we are concerned in identifying the optimal number of hidden units in the hidden layer that will give us the best neural network model. There are several methods in literature that we can use to select the best neural network. The most popular method is the v-fold cross validation method as it does not rely on any probabilistic assumptions and help in determining when over fitting occurs. Other statistical methods like hypothesis testing or information criteria that were introduced and examined by Ulrich Anders and Olaf Korn in 1999, [52] for neural network model selection, they suggested that these statistical methods should take part during the process of developing neural network models. However, since their proposed methods were based on certain probabilistic assumptions, it may not be always applicable in modeling real phenomena. In order to do our comparison we took the best neural network for each method and then tested their performance using the same set of data, this set of data was not used in the training process to identify the model. In the current study, we have used 5-fold cross validation to select the best model for each method, PLANN & DHANN. We divide the male and female data sets into six groups. Five were used in the training and validation, and the last group was used for comparing the best models from the two methods together (hold out data set). In addition, we use the weight decay that helps avoid over fitting and penalize large weight solutions to help in generalization.

As mentioned by B. Ripley [53, 54] a weight decay value between $\alpha = 0.01$ and $\alpha = 0.1$ would be more appropriate depending on the degree of fit that is expected. We have used the cross validation method along with four different values of weight decay $\alpha = \{0.025, 0.05, 0.075, 0.1\}$, to pick the best model. The same procedure of trying different weight decay values were used in, [10]. The cross validation method will help us in finding the optimal number of hidden nodes. In addition, we consider the model with the lowest prediction error when applied to a new data set. Therefore, for each method we picked...
the best model, with lowest cross validation error, then compare their performance on the
hold out data set. We repeated our comparison for the four values of weight decay, since
for same data two factors affects ANN performance (Number of hidden units and Weight
decay value).

3.4 Results

In our analysis, we used ten time intervals (12 month each) and in order to do the compari-
son between the PLANN and DHANN method we used 10 inputs for the ten time intervals
in PLANN instead of one, so that we can compare the output of the PLANN with the sec-
ond method. After training, the cross validation method resulted in choosing the networks
with 52 hidden units (number of hidden nodes seems to be large but by taking into account
the number of output units, we have 10 parallel networks with five hidden nodes each)
for both methods as the best model. We obtain similar results from ANN trained with the
four different values of weight decay. Still, we want to examine the prediction accuracy
for all eight available models to choose our best-fit model. Table 1 shows the comparison
between the eight models for male melanoma patients and Table 2 shows the comparison
for the female melanoma patients.

Table 3: Average prediction error for eight competing neural network models for estimat-
ing the survival time of Male melanoma patients

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>Stdv</th>
<th>Count</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PLANN results</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \alpha = 0.025 )</td>
<td>-0.3776</td>
<td>1.7876</td>
<td>85460</td>
</tr>
<tr>
<td>( \alpha = 0.050 )</td>
<td>-0.4113</td>
<td>1.9388</td>
<td>85460</td>
</tr>
<tr>
<td>( \alpha = 0.075 )</td>
<td>-0.308</td>
<td>1.4295</td>
<td>85460</td>
</tr>
<tr>
<td>( \alpha = 0.1 )</td>
<td>-0.3366</td>
<td>1.5329</td>
<td>85460</td>
</tr>
<tr>
<td><strong>DHANN results</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \alpha = 0.025 )</td>
<td>-0.0236</td>
<td>0.2029</td>
<td>85460</td>
</tr>
<tr>
<td>( \alpha = 0.050 )</td>
<td>-0.0215</td>
<td>0.1710</td>
<td>85460</td>
</tr>
<tr>
<td>( \alpha = 0.075 )</td>
<td>-0.0197</td>
<td>0.1539</td>
<td>85460</td>
</tr>
<tr>
<td>( \alpha = 0.1 )</td>
<td>-0.0197</td>
<td>0.1485</td>
<td>85460</td>
</tr>
</tbody>
</table>

It is clear from Table 1, that using DHANN method yields better predictive neural net-
work model than the PLANN. Among the four competing models of DHANN method we have chosen the model with weight decay $\alpha = 0.1$ as the best fit model for predicting survival times of male melanoma patients, that yield smaller mean error and standard deviation. While the results in table 2 supports the decision we have made for male melanoma patients that DHANN method has a better predictive accuracy than that of the PLANN. But the best model for predicting the female melanoma patient’s survival time is the model with weight decay value $\alpha = 0.075$.

Table 4: Average prediction error for eight competing neural network models for estimating the survival time of Female melanoma patients

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>Stdv</th>
<th>Count</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PLANN results</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\alpha = 0.025$</td>
<td>-0.3384</td>
<td>2.1326</td>
<td>69130</td>
</tr>
<tr>
<td>$\alpha = 0.050$</td>
<td>-0.2274</td>
<td>1.4451</td>
<td>69130</td>
</tr>
<tr>
<td>$\alpha = 0.075$</td>
<td>-0.1882</td>
<td>1.1672</td>
<td>69130</td>
</tr>
<tr>
<td>$\alpha = 0.1$</td>
<td>-0.2011</td>
<td>1.2199</td>
<td>69130</td>
</tr>
<tr>
<td><strong>DHANN results</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\alpha = 0.025$</td>
<td>-0.0208</td>
<td>0.2310</td>
<td>69130</td>
</tr>
<tr>
<td>$\alpha = 0.050$</td>
<td>-0.0183</td>
<td>0.1849</td>
<td>69130</td>
</tr>
<tr>
<td>$\alpha = 0.075$</td>
<td>-0.0167</td>
<td>0.1606</td>
<td>69130</td>
</tr>
<tr>
<td>$\alpha = 0.1$</td>
<td>-0.0246</td>
<td>0.3022</td>
<td>69130</td>
</tr>
</tbody>
</table>

### 3.5 Survival Time Findings

It is clear to us that male and female melanoma patients need to be treated differently as its shown by the survival plots in figure 8, which displays the surface plot of survival probability of male (plot on the left) and female (plot on the right). In figure 8, the survival is estimated as a function of Age at diagnosis and time in years. The tumor thickness is 0.58 mm and ulceration variable is set to 0 and considering the patient was diagnosed in the initial stage. Male infant patients have less survival probability than that of female infant patients over a 10 year period. Where as a male patient at age 40 to 50 seems to have higher survival probabilities compared to female patients at the same age. Figure 9,
Figure 8.: Survival probability function surface plot results for Age for male melanoma patients. Plot on left for male patient. Plot on right for female patient.

Figure 9.: Survival probability function surface plot results for tumor thickness. Plot on left is for the male patient diagnosed at age of 20 years old. Plot on right for male diagnosed at age 60 years old.
Figure 10.: Survival probability function surface plot results for tumor thickness. Plot on left is for the female patient diagnosed at age of 20 years old. Plot on right for female diagnosed at age 60 years old displays the surface plot of survival results for male melanoma patients for tumor thickness ranging from 0.01 mm to 9 mm. The surface plot (on left) is for male patient diagnosed at 20 years of age, whereas the plot (on right) for male patient diagnosed at 60 years old. As we can see survival estimates for young men is farther away and lower than that of older men, these findings were found similar to a recent study by D. Fisher and A. Geller in 2013 [55]. Fisher and Geller mentioned that more attention was given to older men over the past years and suggested that more awareness needed to be addressed to young men to help in early detection of melanoma. They also mentioned the difference between young men and young women, which we can figure out by comparing the two left plots of figure 9 and figure 10. The survival probability for young men (diagnosed with tumor thickness larger than 4mm) within two years of diagnosis is too low (almost 0) compared to that of young women. Some of our significant findings were found to be similar to those found in another study by C. Gamba et. al. in 2013 [56]. However, more investigation and statistical data analysis are required to better understand the causes of the differences between young males and females, and to plan new strategies to fight the major pernicious form of skin
3.6 Conclusions

In modeling survival data with artificial neural network, it is more prevalent to utilize DHANN. The prediction accuracy is much better compared to the PLANN. However, the results may change if some how the survival data contains time varying covariates (risk factors). One can attempt to amend the PLANN model by differentiating between the individuals who survived the whole duration time and those who dropped out during the duration time, which opens another area of research in this field. With regard to learning techniques for ANN, P. J. Lisboa [57] have amended the PLANN by adapting the Bayesian learning for neural networks, developed by Mackay in 1995 [44]. It is still an open problem, How the Bayesian learning will affect the performance of DHANN? Whether it’s going to change the comparison results with the PLANN?, among other questions that we need answers and opens more areas of research on this type of problems. Some of the contributions we obtained from the current study is:

1. The use of artificial neural network have priority over conventional statistical methods, which is being able to form a model with several response variables.

2. There exist two strong methods for predicting survival time using artificial neural network. If data used consists of time varying covariates (Use PLANN). If data does not contain time varying covariates (Use DHANN).

3. The current study opened the door to additional research points, one of which, How to use DHANN in the analysis of competing risks?

In chapter 4, we present a solution for using DHANN in the analysis of competing risks, by introducing binary variables for each risk in the matrix of covariates.
Chapter 4
New Bayesian Learning for Artificial Neural Networks with Application on Survival time for Competing Risks

In the present chapter, we introduce a new method of utilizing artificial neural network (ANN) in modeling survival data of competing risks. Additionally, we present and validate a new Bayesian learning technique for neural networks. The new neural network architecture that we propose in this chapter is used to study the risks associated with patients diagnosed with melanoma (skin cancer). Patient’s information diagnosed with melanoma in the United States from the years 2000 to 2010 were gathered from the Surveillance, Epidemiology, and End Results Program (SEER) [58]. We used Harrells c-index to validate the new proposed neural network structure, in addition we used the v-flod cross validation method to check the model on different set of data and to find the best fit number of hidden units.

4.1 Literature Review

In chapter 1 we reviewed several methods used for survival time analysis. Another set of models that arises and uses artificial neural networks (ANN), two of which we discussed in chapter 3. ANN is well known in clinical trials as a predictor classifier, predicting the diagnosis of a specific cancer as in (Khan, et al., 2001), among several others. In survival analysis, ANN was previously used to predict whether a patient will live the current time period or not (Bottaci, et al., 1997) [6], others were proven to be better than the Cox PH model (Taktak, et al., 2007) such as the partial logistic artificial neural network PLANN
In which the authors presented a method by utilizing ANN to model survival time as a function of other variables using the partial logistic regression approach on censored data as we discussed on previous chapter, also an extension of PLANN for competing risks analysis PLANNCR, by Bignazoli, Boracchi, Ambrogi, & Marubini, 2006.[59]. PLANNCR is a neural network, that shares same input layer structure as PLANN where time interval is treated as ordinal variable and activation function use in hidden layers is the logistic function, similar to PLANN. The output layer in PLANNCR consists of a number of output units equal to the number of risks in the study. The activation function for output unit $r$ is the softmax function given by

$$
\hat{h}_{tr} = \frac{Z_{tr}(x)}{\sum_{r=1}^{R+1} \exp[Z_{tr}(x)]}
$$

(4.1)

where $Z_{tr}(x)$ is the function of predictors. Beside the choice of the Kullback-Leibler distance as the error function makes the PLANNCR perform as a generalized linear models. PLANNCR predicts the conditional probability that a subject will be censored in the $t^{th}$ time period for the $R$ risk, and by comparing it to other competing models used for competing risks, the PLANNCR came the best. But still as we clarified in chapter 3 that due to the use of time as one of network inputs makes you repeat subjects information. We have shown that this was one of the reasons why DHANN performed better than PLANN in a single risk case. One of our goals in this chapter is to amend the DHANN to be able to model survival functions for competing risks.

On the other hand, training of ANN should be done with cautions, as over-fitting is a major problem in any ANN system. That is, error of unseen data set (different from the data set ANN was trained on) occurs to be larger than the training error. The use of a penalty term based on Bayesian method was introduced to over come over-fitting issues (Bishop, 1995) [41], and it was used in PLANN and PLANNCR. However, the learning of PLANNCR was further improved by applying the Bayesian regularization (known as the evidence procedure) for training neural networks (MacKay, 1992)[43] by (Lisboa, et al., 41
2009) PLANNCR-ARD [60], that resulted in similar estimates for the cumulative cause-specific hazards if one would use the Nelson-Aalen nonparametric method, with the advantage of determining the effects of covariates on the hazard function using the automatic relevance determination. The idea behind using the PLANN or PLANNCR is based on discrete hazard models, as mentioned in previous chapter. This case the record of one subject in the study will be replicated for the number of time periods the event did not take place or the subject time becomes censored, which results in limitations, especially for large data sets. On the other hand the evidence procedure that was utilized with PLANNCR is not considered as fully Bayesian technique as it does not integrate the posterior distribution but rather searches for optimal parameters, [43] and since it is based on Gaussian approximation to the posterior, the network performance may break down as the number of hidden units increases,[45]. The investigation of the latter point and a solution for it by using the Hybrid Markov Chain Monte Carlo method, (HMC), that was introduced by Radford Neal in 1996, [45].

![Figure 11. PLANNCR structure with logistic activation function for hidden units layer with softmax function (4.1) for output units layer](image_url)
4.2 Melanoma Patient’s Data

Data used in validating our new proposed neural network structure for competing risks, were collected from the Surveillance, Epidemiology, and End Results Program (SEER), [58]. Data consists of patients diagnosed with melanoma in the United States from the years 2000 to 2011, thus the period of study is eleven years. Since we are going to estimate hazard function based on discrete survival method, we chosen to divide the study time to eleven time intervals, each interval consisting of 12 months. We have chosen a total number of patients with complete information of the risk factors used equal to 185,108. As we did in our previous study, we divided our data set according to gender as they do not share same survival time distribution. Figure 12 displays the number of patients divided by gender, then by tumor behavior, then cancer stage. As we can see, 97.5% of patients (Males and Females) are diagnosed with invasive tumor behavior, known as Malignant. Although skin cancer is curable, especially in early stages, it is also known to be deadly especially
with an invasive tumor behavior. So, the need of a model than can predict the survival time of melanoma patients based on such variable (tumor behavior) is very important. The use of such models that describes the behavior of risk factors on survival time is vital, to help patients decide their course of treatment, and to be able to educate people on the importance of checking up and give good care for their skin changes. A full list of selected risk factors from SEER database is

- **Age**: Age of Patient at diagnosis by melanoma.

- **Tu**: Size of Tumor thickness diagnosed.

- **Stage 1**: Localized: An invasive neoplasm confined entirely to the organ of origin.

  *Stage 0: noninvasive is the baseline.*

- **Stage 2**: Regional: A neoplasm that has extended either beyond the limits of the organ of origin or onto regional lymph nodes or combination of both.

- **Stage 3**: Distant: A neoplasm that has spread to parts of the body away from the primary tumor.

- **Stage 4**: Un-staged: Information is not sufficient to assign a stage.

- **Behav**: Identifies the tumor behavior either noninvasive (in situ) or invasive (malignant).

- **Seq**: Sequence number, describes the number and sequence of all reportable malignant, in situ, benign, and borderline primary tumors, which occur over the lifetime of a patient.

For melanoma patients diagnosed in the United States between the years 2000 to 2011, we have identified three possible risks; the first risk *patient deceased by Melanoma*, the second risk *patient deceased by a different Cancer*, and the third risk *patient deceased by Non-Cancer*. All of these risks were identified from the Cause of death variable in the
SEER database. We also used the Vital record variable to identify whether a patient is deceased by the end of the study or still alive. To identify the patients who are censored (specifically patients who lost follow up) we used the survival time variable and added it to time in which patient was diagnosed, along with the vital record status. Those patients with whom vital status was alive and his/her survival time was not complete to end of the study date were marked as *Lost follow up* (right censored).

**Table 5: Distribution of possible risks for melanoma patients**

<table>
<thead>
<tr>
<th>Risks</th>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alive or lost follow up</td>
<td>82873(44.8%)</td>
<td>73891(39.9%)</td>
</tr>
<tr>
<td>Deceased by Melanoma</td>
<td>7607(4.1%)</td>
<td>3427(1.9%)</td>
</tr>
<tr>
<td>Deceased by Other Cancer</td>
<td>3820(2.1%)</td>
<td>1725(0.9%)</td>
</tr>
<tr>
<td>Deceased by Non-Cancer</td>
<td>7929(4.3%)</td>
<td>3836(2.1%)</td>
</tr>
</tbody>
</table>

From Table 5 we see the number of male patients (in blue) for each of our identified risks. Being alive at the end of the study is the baseline, i.e. patients remains in same stage as the starting point (being diagnosed with Melanoma). The percentage of each category (in red) is with respect to the whole sample (count of 185,108). The percentage of lost follow up for Male and Female patients does not exceed 1.5% from the total number of *Alive or lost follow up* category. As we can see the probability of being deceased by melanoma is very small, 4% for male patient & almost 2% for female patient. For fitting a statistical model for survival time with 3 competing risks we kept out 20% of male data and 20% of female data for validating network models and comparing them using unseen data set different from the data set the network models were trained on. Though we have mentioned in chapter 2 that using Bayesian inference in neural network have an advantage of comparing between several networks models, we preferred to use v-fold cross validation technique. We have
chosen to split our data set to 10 approximately equally sized groups. More on the use of
v-fold cross validation will be explained in the Results section of this Chapter.

4.3 Discrete Hazard Artificial Neural Network for Competing Risks

In this section we present a new neural network model for predicting survival time of com-
peting risks. We will use the data discussed in the previous section to test and validate
the new proposed model. As we showed in Chapter 3, that the DHANN performed better
compared to PLANN. So, the new proposed neural network is an update to the DHANN
of single risk. The idea starts from Cox-PH for competing risks or what is known by the
Cox model for the cause-specific hazard [61], in which the conditional hazard for risk \( R \)
is given by:

\[
h_r[t, X(t)] = h_{or}(t) \exp[Z(t) \beta_r] \quad (4.2)
\]

where, \( Z(t) \) is a vector of \( p \) derived covariates, and \( h_{or} \) is the baseline hazards. \( r = 1, 2, .., R \) number of risks. The aim of DHANN is to predict the discrete hazard prob-
ability for all time intervals at the same time by having the number of output units in the
output layer equal to the number of time intervals. It would be difficult to update DHANN
for competing risks, to follow PLANNCR approach which is training network to predict
the conditional probability of being censored for each risk (by having number of output
units in output layer equal to number of risks). So it means that if we consider that we have
3 possible risks, and our time intervals of the study is equal to 10, Then we need 30 output
units in the output layer, which is not practical. Our proposed solution for have DHANN
accommodate for competing risks is to add a number of binary variables equal to the num-
ber of required risks to the vector of derived covariates. In our case we have 3 possible
risks, so our derived vector of covariates \( Z(t) \) is given by

\[
Z(t) = \beta_{01} R_1 + \beta_{02} R_2 + \beta_0 3 R_3 + \beta X \quad (4.3)
\]
where \((\beta_{01}, \beta_{02}, \beta_{03})\) presents the coefficients for the three binary variables \(R_1, R_2,\) and \(R_3\) respectively. The binary variables are defined as:

\[
R_r = \begin{cases} 
0 & \text{for patient deceased of another risk or still alive} \\
1 & \text{for patient deceased of the } r \text{ risk}
\end{cases}
\]

as we discussed previously, we are going to add three binary variables (meaning adding also 3 additional input units in the neural network input layer) to the vector of risk factors (covariates). The coding of these three variables is illustrated in the following table.

A patient that was deceased by melanoma, his/her record/row of information will be starting with three additional values \(1 0 0 \cdots \cdots \). If a patient was deceased by other type of cancer rather than melanoma (and is a melanoma patient), then his/her record will start with \(0 1 0 \cdots \cdots \). If a patient was deceased by a non-cancer cause such as car accident, heart attack, flu, etc. then his/her record starts as \(0 0 1 \cdots \cdots \). The last case would be for patient who is still alive at the end of the study or lost follow and his/her vital status is unknown or not known to be deceased, then his/her record starts as \(0 0 0 \cdots \cdots \).

Table 6: Coding of the three added binary variables representing the possible risks

<table>
<thead>
<tr>
<th>(R_1)</th>
<th>(R_2)</th>
<th>(R_3)</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>Deceased by Melanoma</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
<td>Deceased by Other Cancers</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
<td>Deceased by Non-Cancer</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>Still Alive or lost follow up</td>
</tr>
</tbody>
</table>

As we showed in Chapter 3, using DHANN requires the formation of training vector instead of having one response variable. In case of DHANN the training vector was formed with length \((1 \times T)\), where \(T\) is the number of time intervals on the following form:
Now we are upgrading the DHANN to fit competing risk model based on the addition we did on the input layer. The training vector for the new method DHANN-CR is given by:

\[ h_{ik} = \begin{cases} 
0 & \text{for } 1 \leq k \leq K \\
1 & \text{for } t \leq k \leq K \text{ and event } = 1 \\
\frac{r_k}{n_k} & \text{for } t \leq k \leq K \text{ and event } = 0 
\end{cases} \]

where \( r_k \) is the Kaplan-Meier estimate of hazard probability for risk \( r \). In the following we will take an example of three patients to illustrate the reformation of data prior to applying it to the DHANN-CR. Consider data of 3 patients are given in the following table

<table>
<thead>
<tr>
<th>ID</th>
<th>ST</th>
<th>Age</th>
<th>Tu</th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>Behav</th>
<th>Seq</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>109</td>
<td>51</td>
<td>1.25</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>Melanoma</td>
</tr>
<tr>
<td>2</td>
<td>130</td>
<td>35</td>
<td>0.85</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>Still Alive</td>
</tr>
<tr>
<td>3</td>
<td>96</td>
<td>63</td>
<td>1.01</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>Lost follow up</td>
</tr>
</tbody>
</table>

In Table 7, we have three patient record with the chosen 11 risk factors we identified in the data section. Variables \( S_1, S_2, S_3 \) and \( S_4 \) stands for the the four different cancer stages. Now, first patient was diagnosed by melanoma at Age 51, with tumor thickness of 1.25mm, this patient have survived for 109 months \( ST = 109 \). The status of the first patient mentions that he/she deceased by melanoma. Then this patient record will start with \([1 0 0 51...]\) and his/her training hazard vector corresponding to his risk factors would be
in which for the first 4 time intervals the event did not take place and it took place on the fifth year (i.e 5th time interval). Second patient record with start with three binary values as zeros since patient is still alive at the end of the study and his/her vector of training will be all zeros as neither of the expected 3 events took place (which are deceased by melanoma, or deceased by another cancer, or deceased by a non-cancer cause). In the meantime, the third patient lost follow-up from what appears in his/her record of information. By looking at the third patient survival time ($ST = 96$) we obtain that the patient lost follow-up after year eight, which means the eight’s interval, then the training vector of this patient will start with zeros up to the eight time interval then for the ninth, tenth, and eleventh values will be equal to $d_{rj}/n_j$ and since patient who lost follow-up we repeat his record three times as he/she is in a risk of possible three events. The reformation of the those 3 patients is illustrated in the following table by showing the first 3 values in patient record and his/her corresponding training vector.

Table 8: Example of Re-structuring data for DHANN-CR

<table>
<thead>
<tr>
<th>ID</th>
<th>Covariates</th>
<th>training vector</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$R_1$ $R_2$ $R_3$ $X_1$ ... $X_p$</td>
<td>$h_1$ $h_2$ ... $h_9$ $h_{10}$ $h_{11}$</td>
</tr>
<tr>
<td>1</td>
<td>1 0 0 $x_{11}$ ... $x_{1p}$</td>
<td>0 0 ... 0 1 1</td>
</tr>
<tr>
<td>2</td>
<td>0 0 0 $x_{21}$ ... $x_{2p}$</td>
<td>0 0 ... 0 0 0</td>
</tr>
<tr>
<td>3</td>
<td>1 0 0 $x_{31}$ ... $x_{3p}$</td>
<td>0 0 ... 0.0476 0.0498 0.0292</td>
</tr>
<tr>
<td>3</td>
<td>0 1 0 $x_{31}$ ... $x_{3p}$</td>
<td>0 0 ... 0.5331 0.0564 0.0321</td>
</tr>
<tr>
<td>3</td>
<td>0 0 1 $x_{31}$ ... $x_{3p}$</td>
<td>0 0 ... 0.1115 0.1248 0.0689</td>
</tr>
</tbody>
</table>

Once the data is re-structured the same way we explained in the previous example, we can use the same neural network structure for hidden and output layers as in DHANN. Figure 13, displays the new structure of DHANN-CR with our three identified risks.

4.4 New Bayesian Learning Technique

In Chapter 2, we explained the two Bayesian techniques used in learning of neural network. Most of the application that have been using Bayesian learning were developed using the
Figure 13.: Structure of DHANN-CR with 3 identified risks

evidence procedure. Our aim is to be able to use the fully Bayesian learning technique done through the hybrid Monte Carlo sampling (HMC). We propose a solution for the problems found in the HMC technique making researchers to avoid using it. As we mentioned the HMC technique uses initial constant values for hyperparameters that does not change based on the data. So, if the choice of initial hyperparameters does not capture the actual prior belief, then neural network performance will not be good. A solution for the current problem is use the part of re-estimating the hyperparameters of the evidence procedure as it is done based on the data. Our approach is to reach an approximate values of hyperparameters which capture the true distribution of weights variance. We called this new approach (E-HMC), E: stands for evidence and HMC: stands for hybrid Monte Carlo sampling. Our new proposed method (E-HMC) works as follows:

1. Choose initial values for hyperparameters $\alpha$ and $\beta$, and network weights and bias.

2. Estimate the most probable values of $\alpha$ and $\beta$ that maximize the evidence given by (2.32) considering the initial weight values as the most probable weight values, where these values are estimated by using (2.33) and (2.34).
3. Use the HMC to sample from the posterior distribution of the weights, with hyperparameters value found in step 2.

To validate our proposed method, we trained our neural network models for the chosen data set of melanoma patients using the three Bayesian techniques, the evidence procedure, the HMC sampling method and the E-HMC, the newly proposed method. All neural networks were trained with initial values of hyperparameters equal to 0.01. The automatic relevance determination was also used to determine which of our 11 risk factors contributes more towards the survival time of a melanoma patient. All neural networks models were trained using MATLAB software, by using a set of algorithms done by Ian Nabney [62] called NETLAB. The evidence procedure and the HMC sampling were the two major algorithms we use from NETLAB, although we had to update some of the codes as it was done on older versions of MATLAB.

4.5 Results

The main problem in statistical modeling is to find the model that best fits the data based on two criteria; the first one with respect to the best neural network, mainly the number of hidden nodes in hidden layers, and the second with respect to the learning algorithm. In order to accomplish that we use two model validation techniques, the \textit{v-fold cross validation} and Harrell’s C-index \cite{63}. Harrell’s C-index, discrimination index, measures the predictive accuracy of statistical models, and widely used in survival analysis to compare the effectiveness between models. In 2005, there was an upgrade on the Harrell C-index to get a time-dependent discrimination index for censored survival data by Antolini, Boracchi, and Biganzoli \cite{64}. Harrell’s C-index for proportional hazard models is given by:

\[
c = \frac{\sum_{i=1}^{n-1} \sum_{j=1;j\neq i}^{n} \text{concordance}_{ij}}{\sum_{i=1}^{n-1} \sum_{j=1;j\neq i}^{n} \text{comparable}_{ij}}
\]  

(4.5)
where \( \text{comparable}_{ij} \) is the number of comparable pairs \( \{i, j\} \) from sample of \( n \) observations. And, \( \text{concordance}_{ij} \) is the number of concordant pairs. The two variables represents binary indicators that are defined as:

\[
\text{comparable}_{ij} = I\{ST_i < ST_j \& d_i = 1\} + I\{ST_i = ST_j \& d_i = 1 \& d_j = 0\} \quad (4.6)
\]

and,

\[
\text{concordance}_{ij} = I\{z(x_i) > z(x_j)\} \ast \text{comparable}_{ij} \quad (4.7)
\]

where \( z(x_i) \) is the linear predictor function in (4.2). We have created and implemented a new algorithm in MATLAB to calculate the \( C \)-index for any neural network in the form of PLANN, or DHANN, or DHANN-CR.

### 4.5.1 Algorithm for computing \( C \)-index through MATLAB

To find \( C \)-index we first find the number of comparable pairs then use it to calculate the number of concordant pairs. Recall that we used 10-fold cross validation, so for the same number of hidden units we have a total of 10 different neural networks models (Only in case of using evidence procedure). Thus, the \( C \)-index is calculated for each of these networks and then average it for the same number of hidden units for each time interval (we have a total of 11 time intervals). The code created for counting the comparable pairs is given on next page. As we previously mentioned our data set is divided into 10 groups, in line 4 a cell array consisting of 10 matrices is formed. Each of this matrices has five columns and number of rows equal to the sample size of each of the ten groups. The five columns are as follows, survival time of each patient in that group, plus four binary variables, first binary variable is the status of that patient, indicating whether event of being dead took place or not, other remaining binary variables are for the three risks defined in table 4. Thus, the second column will let us know if the subject experienced the event or not, if event took place the remaining three columns will let us know event occurred due to which risk. The
reason for that is to be able to measure the predictive accuracy for each risk.

In line 7-8 we are retrieving one of ten data group matrix and obtaining its number of rows (sample size of that group). Lines 10-19 finding how many pairs of that group are comparable. For example if we want to find the C-index for being deceased by melanoma, then to obtain the count of comparable pairs in each data group we check for each pair \( \{i, j\} \).

If the survival time of subject \( i \) is less than the subject \( j \) \( (i \neq j) \) and if subject \( i \) experienced the event and it is due to melanoma, then these pairs are comparable. These pairs
are also comparable if their survival times are equal and subject $i$ experienced the event due to melanoma plus subject $j$ also experienced the event. Otherwise the two pairs are not comparable. Now we move to determining the concordant pairs based on the network outcome for each time period.

```matlab
conc = zeros(m,m); yy=prediction{hn,cv}; for i = 1:m-1
    for j = 2:m
        if((FS(i,1)<FS(j,1)) && (FS(i,2)==4&&FS(i,4)==1))
            if(yy(i,pre)>yy(j,pre))
                conc(i,j)=1*comp(i,j);
            else
                conc(i,j)=0;
            end
        end
    end
    sconc = sum(sum(conc));
    scomp = sum(sum(comp));
    c = sconc/scomp;
end
```

The code for counting the number of concordant pairs is shown on the next page. The number of concordant pairs is determined if for each comparable pair \{i,j\} the predicted hazard of $i$ is larger than the predicted hazard of $j$, such that subject $i$ experienced the event while subject $j$ did not, lines 5-19 of code on next page. On the second line we obtain the prediction done with each neural network structure for each of the ten data groups. The whole counting process of comparable pairs and concordant pairs are repeated for the different number of neural networks trained with different number of hidden units. The
same process was applied to all neural networks trained with each of the three learning techniques.

4.5.2 Comparison with C-index

Table 9: C-index for first time interval for female patients

<table>
<thead>
<tr>
<th>TI</th>
<th>LA</th>
<th>HN</th>
<th>Data groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Evid</td>
<td></td>
<td></td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>10</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>11</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>13</td>
</tr>
</tbody>
</table>

| HMC |    |    | 4 | 0.9419 | 0.9410 | 0.9438 | 0.9442 | 0.9385 | 0.9459 | 0.9393 | 0.9489 | 0.9414 | 0.9490 |
|     |    |    | 5 | 0.9487 | 0.9405 | 0.9429 | 0.9449 | 0.9391 | 0.9494 | 0.9397 | 0.9492 | 0.9407 | 0.9489 |
|     |    |    | 6 | 0.9432 | 0.9385 | 0.9416 | 0.9422 | 0.9377 | 0.9452 | 0.9358 | 0.9460 | 0.9414 | 0.9484 |
|     |    |    | 7 | 0.9420 | 0.9399 | 0.9472 | 0.9468 | 0.9387 | 0.9440 | 0.9376 | 0.9514 | 0.9426 | 0.9487 |
|     |    |    | 8 | 0.9412 | 0.9405 | 0.9435 | 0.9462 | 0.9389 | 0.9405 | 0.9383 | 0.9485 | 0.9392 | 0.9496 |
|     |    |    | 9 | 0.9440 | 0.9395 | 0.9444 | 0.9449 | 0.9367 | 0.9467 | 0.9376 | 0.9486 | 0.9397 | 0.9493 |
|     |    |    | 10 | 0.9425 | 0.9309 | 0.9452 | 0.9438 | 0.9378 | 0.9443 | 0.9387 | 0.9460 | 0.9416 | 0.9462 |
|     |    |    | 11 | 0.9424 | 0.9382 | 0.9431 | 0.9421 | 0.9373 | 0.9453 | 0.9360 | 0.9447 | 0.9395 | 0.9492 |
|     |    |    | 12 | 0.9422 | 0.9389 | 0.9429 | 0.9441 | 0.9389 | 0.9467 | 0.9384 | 0.9454 | 0.9393 | 0.9482 |
|     |    |    | 13 | 0.9407 | 0.9405 | 0.9419 | 0.9429 | 0.9366 | 0.9448 | 0.9367 | 0.9471 | 0.9387 | 0.9490 |

| EHMC |    |    | 4 | 0.9447 | 0.9407 | 0.9436 | 0.9448 | 0.9389 | 0.9460 | 0.9397 | 0.9501 | 0.9430 | 0.9495 |
|      |    |    | 5 | 0.9444 | 0.9420 | 0.9453 | 0.9460 | 0.9399 | 0.9473 | 0.9406 | 0.9509 | 0.9427 | 0.9494 |
|      |    |    | 6 | 0.9419 | 0.9423 | 0.9443 | 0.9444 | 0.9416 | 0.4025 | 0.9399 | 0.9481 | 0.9412 | 0.9502 |
|      |    |    | 7 | 0.9423 | 0.9393 | 0.9441 | 0.9429 | 0.9385 | 0.9462 | 0.9380 | 0.9483 | 0.9401 | 0.9485 |
|      |    |    | 8 | 0.9429 | 0.9386 | 0.9432 | 0.9434 | 0.9364 | 0.9462 | 0.9373 | 0.9461 | 0.9397 | 0.9473 |
|      |    |    | 9 | 0.9440 | 0.9395 | 0.9444 | 0.9449 | 0.9367 | 0.9467 | 0.9376 | 0.9486 | 0.9397 | 0.9493 |
|      |    |    | 10 | 0.9425 | 0.9399 | 0.9452 | 0.9438 | 0.9378 | 0.9443 | 0.9387 | 0.9460 | 0.9416 | 0.9462 |
|      |    |    | 11 | 0.9424 | 0.9382 | 0.9431 | 0.9421 | 0.9373 | 0.9453 | 0.9360 | 0.9447 | 0.9395 | 0.9492 |
|      |    |    | 12 | 0.9422 | 0.9389 | 0.9429 | 0.9441 | 0.9389 | 0.9467 | 0.9384 | 0.9454 | 0.9393 | 0.9482 |
|      |    |    | 13 | 0.9407 | 0.9405 | 0.9419 | 0.9429 | 0.9366 | 0.9448 | 0.9367 | 0.9471 | 0.9387 | 0.9490 |

We want to test how good our new neural network performs as to predicting the survival time of competing risks. Additionally, we want to find the best learning algorithm that yields better learning for neural network. We use the C-index to validate our new approach (DHANN-CR) and the new learning algorithms. For each of the ten data groups of males and females, we tried eleven neural network structures starting with four hidden
units in the hidden layer to thirteen hidden units. For each of the neural network structures, we have computed the \textit{C-index} on each of the ten data groups. The \textit{C-index} was also calculated for each time interval and that process was repeated for the three different risks. Results will be displayed in the following table. Table 9 shows \textit{C-index} values obtained from neural network models (DHANN-CR) for prediction done on first risk only (deceased by melanoma). Table 9 shows only results obtained for first time interval and we obtained similar results for all eleven time intervals. As we can see from the table there is no significant difference between \textit{C-index} values among networks trained with different learning algorithms (Evid: Evidence, HMC: Hybrid Monte Carlo, and EHMC: newly proposed method). On the other hand, obtaining \textit{C-index} values greater than 0.9 is an indication that neural network models have high predictive accuracy, which means that the new neural network structure (DHANN-CR) perform very well. In the following table we give a summary of the averaged \textit{C-index} values over the 10 different groups for the eleven time periods for neural networks structure of four hidden nodes. Table 10 also gives 95% confidence interval for \textit{C-index} for each time period.

\textbf{Table 10: \textit{C-index} for the three learning techniques for ANN with 4 hidden nodes for Risk 1}\n
\begin{tabular}{|c|c|c|c|}
\hline
Time Period & Evidence & HMC & E-HMC \\
\hline
T_1 & 0.946 (0.944, 0.949) & 0.943 (0.941, 0.946) & 0.944 (0.942, 0.946) \\
T_2 & 0.949 (0.947, 0.951) & 0.950 (0.948, 0.952) & 0.949 (0.948, 0.951) \\
T_3 & 0.950 (0.948, 0.952) & 0.952 (0.950, 0.954) & 0.951 (0.949, 0.953) \\
\vdots & \vdots & \vdots & \vdots \\
T_{10} & 0.945 (0.943, 0.948) & 0.944 (0.942, 0.947) & 0.946 (0.943, 0.948) \\
T_{11} & 0.946 (0.944, 0.948) & 0.946 (0.944, 0.948) & 0.947 (0.945, 0.949) \\
\hline
\end{tabular}

\textit{C-index} comparison resulted in no difference between the three learning techniques in terms of the survival predictive accuracy. On the other hand, we obtained strong evidence that supports the validity of our proposed neural network approach for modeling survival time of competing risks which is DHANN-CR. Next, we will see the v-fold cross valida-
tion comparison that will compare between the three learning techniques by looking at the errors.

4.5.3 Comparison with \( v \)-fold Cross Validation

In this section, we compare the errors obtained by DHANN-CR trained with three different learning algorithms. We would like to recall that all networks started with the same initial values of hyperparamters which is 0.01. Also we used the ARD, which means that each of our eleven inputs is assumed to be controlled by weights having different distributions.

![Graph showing errors distribution for neural networks trained with 11 different values of hidden nodes with ten different data, using Hybrid Monte Carlo Sampling](image)

Figure 14.: Errors distribution for neural networks trained with 11 different values of hidden nodes with ten different data, using Hybrid Monte Carlo Sampling

We start with neural networks trained with Hybrid Monte Carlo Sampling as it resulted in the highest error (between \(9.6 \times 10^5\) to \(10.4 \times 10^5\)). Figure 14 shows the amount of error \((Y - axis)\) obtained from neural network trained on different groups \((X - axis)\) of data using the same learning algorithm.
Figure 15.: Errors distribution for neural networks trained with 11 different values of hidden nodes with ten different data, using Evidence Procedure

Figure 16.: Errors distribution for neural networks trained with 11 different values of hidden nodes with ten different data, using E-Hybrid Monte Carlo

We wish to mention that this amount of error is for predicting the hazard probability function for 6644 patients, that is, the total number of predictions is $6644 \times 11 = 73084$ for
the female data. The average error using the Hybrid Monte Carlo Sampling is between 13.092 and 14.282, which is not what we expect for an average error to be. In addition, the errors for the same neural network structures are not constant, having variations, rather than having constant behavior on different data sets.

This explains what we mentioned before using HMC requires the knowledge of the correct prior values. One could get better models using HMC by trying different initial values for hyperparameters, but the cost of computational time it takes to run HMC for one model makes it unrealistic to continue. We must mention that Neal [45] had generated 2 million networks to fit a function with 6 data points. In Figure 15, we present the errors distribution using evidence procedure on the same data set as HMC. Note that the errors decreased compared to networks trained with HMC, their variance is with less variability compared to HMC. Figure 16 shows the errors of neural networks prediction, that was trained with our newly proposed method, which is Using the evidence to re-estimate hyperparameters based on the data and then use Hybrid Monte Carlo sampling to obtain samples of the weights distribution from its posterior. As we can see, errors are fewer compared to neural networks trained with Evidence and HMC. Also, errors seems more consistent for different data groups. As the number of hidden nodes increases, error also increase, giving an advantage for using Bayesian learning with neural network no need for large number of hidden units. By looking at the errors of neural networks with five hidden units in the hidden layer (of figure 15) possesses errors with lower variance, have approximately constant variance compared to others. The best fit neural network model for predicting survival time of competing risks is the model trained with E-HMC with five hidden units in hidden layer. All results shown above are for female melanoma patient data, similar results were obtained for male patient data.
4.6 Conclusion

In the current Chapter, we have obtained several important results using artificial neural network for predicting survival time of competing risks. We summarize our findings in the following points:

1. We have presented a new method that utilize artificial neural network in predicting the hazard function of competing risks.

2. We are introducing (proposing) a solution to help use the Hybrid Monte Carlo simulation learning algorithm for neural network in a more efficient way.

3. Using Bayesian inference in the learning of neural networks avoids the need of large number of hidden units in hidden layer.

The DHANN-CR is a new method of utilizing ANN in survival analysis. It’s more useful compared to PLANN, and PLANN-CR especially with the existence of huge data set. The use of Bayesian inference in learning neural network, gives neural network more chance to learn than to memorize, increasing the number of hidden units in neural networks makes it more memorizing tool rather than learning tool. Our future goal is to see if our new proposed learning method can be utilized with artificial neural network in other statistical analysis methods, such as time series analysis, and Categorical data analysis, among others.

In the next Chapter, we introduce a new approach of using Hybrid Monte Carlo Sampling with recurrent neural network for the modeling of time series data.
Chapter 5
Artificial Neural Network for Forecasting Carbon Dioxide Emission in the Atmosphere

In this Chapter, we develop an artificial neural network model utilizing time series approach for forecasting Carbon dioxide, $CO_2$, in the atmosphere. In Chapter three and four we saw the use of Feedforward artificial neural network in survival analysis, however, Feedforward networks can be used in other generalized linear statistical modeling [65–68]. One of the important statistical analysis methods is time series analysis, which deals with forecasting observation in the future for a certain time period that mainly depends on the same reading of that observation on previous time periods. That is, the future predictions depends on previous observations of the same data. Modeling of such situations in ANN requires the use of recurrent neural networks. In this Chapter we will present a new method of using Hybrid Monte Carlo Sampling in the learning of recurrent networks for time series forecasting. We will validate the new proposed model on a popular and vital problem our society is facing, which is carbon dioxide emission in the atmosphere.

Carbon dioxide is strongly connected to climate change, but the impact of carbon dioxide varies according to the source and level of emissions and also according to regional effects,[69]. As we know, the most dominant source for carbon dioxide emission is fossil fuels, making it a major contributing factor in global warming. Other variables that contribute towards the emission of carbon dioxide in the atmosphere are given in Figure 16, [70]. Our future goal is to use artificial neural network to build a model that better understand the contribution of each of the variables in Figure 17, and hence be able to make policies and control the emission of carbon dioxide. United States comes in second place
after China in the ten largest carbon dioxide emitters. United States shares about 14.69% of Global carbon dioxide emission compared to China of 23.43% in year 2014, [71]. Our work starts by fitting a neural network model for predicting the monthly average carbon dioxide emission in United States.

![CARBON DIOXIDE (CO₂) IN THE ATMOSPHERE IN U.S.A.](image)

**Figure 17:** Emission of Carbon Dioxide in the Atmosphere in U.S.A.

### 5.1 Literature Review

There is a number of studies utilizing artificial neural network for the prediction of time series data, making it too hard to track. Models ranging from the use of Feedforward networks to recurrent networks and lately using what is called by wavelet neural network, [72]. One of the most popular series used for forecasting time series is the sunspot series,
a recent study on using neural network with quantum gate was proposed by X. Guan et. al, [73]. The authors of the study presented an improvement for predicting sunspot number series, however no further investigation for testing their model on other time series data. This has been the issue of utilizing neural networks in time series for the last decade. The use of neural networks in time series depends heavily on the data, different models, different architecture of networks were presented some showed better performance compared to the popular ARIMA model others did not, [74]. Fitting ANN model for time series data involves not only finding optimal numbers of hidden units, as it was the case in Chapters 3 & 4, but also the number of input units is variant. The number of input units in forecasting models corresponds to number of previous observations that one would use to predict future outcome. There is no specific method of choosing the number of input units, while researchers tried to find a solution to this problem, one was a claim of choosing the number of hidden units is equal to number of auto-regressive(AR) terms in Box-Jenkins [75]. However, this approach can not be useful as for moving average of order one, MA(1), model there is no autoregressive terms. Another solution was to use Box and Jenkins model identification procedure and then use the number of input terms corresponding to the identified terms. For example, if Box and Jenkins model identification step found AR(2) and MA(1), then the ANN used for such a model would probably have three input nodes, for $X_{t-2}$ and $e_{t-1}$. In a study by Zou et al. [76], compared between performance of ARIMA, ANN and a combined ARIMA, ANN model. In which they used ARIMA to identify AR and MA terms, then use ANN with number of input units equal to number of identified terms from ARIMA model. According to their comparison it was found that the combined ANN and ARIMA model resulted in less mean square error, however, results may vary with different data set. In general there is no specific procedure for the choice of a certain neural network structure for time series analysis, it all depends on the data series. The second part of utilizing ANN in time series involves the choice of learning algorithm. Several learning algorithms were used and proposed in literature, like the use of genetic algorithms as
in, [77–80], while others used regular back-propagation algorithms that were summarized along with results in [74]. On the other hand, some other models used the evidence procedure that was proposed by Mackay [43]. J. Ticknor used evidence procedure along with a three layer Feedforward network to forecast the stock market, [81]. However, Ticknor did not use the evidence, as explained in Chapter 2, to compare between neural network models to choose the optimal number of hidden units.

The aim of the present study is to reduce the issues that need attention for utilizing ANN in time series. We propose the use of recurrent neural network that utilizes Hybrid Monte Carlo sampling learning algorithm by applying it online. In other words, for each input we update the learning of neural networks. The purpose of using HMC is to fix the number of hidden units, as we discussed in Chapter 4 that the networks trained with HMC, does not require large number of hidden units. In the coming sections we highlight the important points of the ARIMA model, then we explain the proposed neural network structure and the developing of a new learning algorithm written for this specific task. The learning algorithm involves the use of HMC algorithm written by Nabney in the NETLAB package for MATLAB software.

5.1.1 Auto-Regressive Integrated Moving Average Models: ARIMA

The auto-regressive integrated moving average (ARIMA) models are the most popular time series methods for modeling stationary time series. ARIMA initially was presented by Box and Jenkins in 1970 [82]. Let \( \{X_t, X_{t-1}, X_{t-2}, ..., X_1\} \) be a time series data, the general mathematical expression of the ARIMA model is given by:

\[
\Phi_p(B)(1 - B)^d X_t = a + \Theta_q(B)\varepsilon_t
\]

where \( \Phi_p(B) \) is the autoregressive operator with order \( p \) defined as

\[
\Phi_p(B) = 1 - \phi_1 B - \phi_2 B^2 - \cdots - \phi_p B^p
\]
where $\Theta_q(B)$ is the moving average operator with order $q$ defined as

$$\Theta_p(B) = 1 - \theta_1 B - \theta_2 B^2 - \cdots - \theta_q B^q$$

where $B$ is the back shift operator, $\varepsilon_t$ is the error term that is normally distributed with mean zero and constant variance. If the time series data shows a seasonal trend, then a generalization of the ARIMA model is to fit seasonal data is called Seasonal autoregressive integrated moving average model (SARIMA) given by:

$$\Phi_p(B) \Phi_p(B^s)(1 - B)^D (1 - B)^d X_t = a + \Theta_q(B) \Theta_Q(B^s) \varepsilon_t$$  \hspace{1cm} (5.2)

where $\Phi_p(B^s)$ and $\Theta_Q(B^s)$ are the seasonal autoregressive and moving average, respectively. The procedure that we follow to fit the ARIMA or SARIMA models know as the Box and Jenkins procedure can be summarized into the following steps.

1. **Model Identification**: In which the order of AR($p$) and MA($q$) are identified. By computing and plotting the autocorrelation function (ACF) and the partial autocorrelation function (PACF). If the series is non-stationary then it may requires filtering (either by non-seasonal or seasonal difference), and if series possesses a non constant variance over time then a transformation may take place.

2. **Model Estimation**: After step 1 is done, and the series is made sure to be stationary (As required by ARIMA procedure), then it follows estimation of the parameters $\phi_1, \phi_2, \ldots, \phi_p$ and/or $\theta_1, \theta_2, \ldots, \theta_q$ using maximum likelihood estimation to minimize the mean square error function.

3. **Model Validation**: Includes tests for white noise errors (making sure errors follows the assumed distribution) by checking errors autocorrelation and partial autocorrelation functions. Additionally, the use of Akaike information criterion, [83](AIC), to compare among several models.
4. **Forecasting**: After model validation, the selected (best) model is used for forecasting and forecasting errors are calculated.

After these four steps, one can use the fitted model to perform forecasting. We will discuss those steps practically on our data set of \( CO_2 \) in the atmosphere. And explain how we picked the number of terms for AR and MA, along with non-seasonal or seasonal difference that took place.

5.2 **The Data**

The monthly average carbon dioxide emission that we use was collected by the National Oceanic & Atmospheric Administration (NOAA). The data contains monthly average carbon dioxide emission from March 1958 to February 2015. The data from March 1958 through April 1974 have been obtained by C. David Keeling of the Scripps Institution of Oceanography as mentioned by NOAA in the data description file, while NOAA started their own measurements through Mauna Loa Observatory station starting in 1974. The carbon dioxide emissions is expressed as parts per million (ppm) which is the number of molecules of \( CO_2 \) in every one million molecules of dried air [84].

![Atmospheric CO2 at Mauna Loa Observatory](image)

Figure 18.: Monthly Average Emission of Carbon Dioxide in the Atmosphere in U.S.A.
Figure 18 displays the monthly average emissions of CO$_2$ in the atmosphere in the U.S.A, obtained from NOAA website [84]. In the current analysis we use information from January 1960 to December 2008 to train the ANN model, and left the remaining to test the forecasting accuracy of our models. Data entering neural network is required to be on small scale to smooth and faster conversion. We did standarized carbon dioxide data using the following formula:

$$\hat{x}_i = \frac{x_i - min\{x_t, x_{t-1}, \ldots, x_1\}}{max\{x_t, x_{t-1}, \ldots, x_1\} - min\{x_t, x_{t-1}, \ldots, x_1\}}$$ (5.3)

5.3 Time Series Data Modeling

In this study we present a model by combining both the ARIMA and ANN. We are going to use the first step in the ARIMA procedure to identify the number of input units in the input layer. Then, we will train a recurrent neural network using Hybrid Monte Carlo Sampling using online learning. First we start with model identification.

5.3.1 Model Identification

Based on Box and Jenkins methodology, a time series data should be stationary, so the first step in model identification is to make sure that the series is stationary, and this is done by looking at the time series plot of the data.

By looking at Figure 19, we notice that as time increases the series also increase suggesting a possible first nonseasonal difference(filtering) is needed as the average of the series is not constant over fixed time periods. So we take the first difference and then plot the series again. As we can see in Figure 20, the series now shows constant mean making it stationary with respect to mean. It also appear that there exist a seasonal trend, in which its order will be determined by looking at the autocorrelation function and partial autocorrelation function.
Figure 19.: Time Series plot for Average CO$_2$ Emission

Figure 20.: CO$_2$ Series after first nonseasonal difference (filtering)

Figure 21, displays the autocorrelation function for the series after taking the first difference. This result indicates the need of seasonal difference as the series express non-stationary in seasonal lags (12, 24, ...). Figure 22, after taking additional seasonal difference gives an indication of AR in nonseasonal term and either the presence of seasonal AR of MA for lags (12, 24, 36, ...). The graph for partial autocorrelation function (Figure 23) con-
firms that we have AR of order one in the non-seasonal part and a MA of order one in seasonal part. Figure 22, plots the autocorrelation function for series after taking 1 non-seasonal difference and one seasonal difference. Figure 23 plots the partial autocorrelation function for the same series. The ACF behavior for AR model and MA model is the same, while the difference will be obtained from the partial autocorrelation function. Since, in partial autocorrelation function we have one term above zero, then it is an indication of AR(1) model. The seasonal lags in Figure 22 and Figure 23 gives the indication of a seasonal term in SMA(1). Overall, model identification suggests the best model of the following form

\[ \text{SARIMA}(1,1,0)\times(0,1,1)_{12}, \]  

that is,

\[ \Phi_1(B)(1-B)^1(1-B)^1X_t = a + \Theta_Q(B)\varepsilon_t \quad (5.4) \]

with 1 nonseasonal difference and one seasonal difference, and with one nonseasonal AR(1) with one seasonal MA (SMA(1)).

So for fitting the neural network model to \( CO_2 \) data, we are going to use three input units in the input layer. Since we have identified one nonseasonal difference we will take
into consideration two previous values $X_{t-1}$ and $X_{t-2}$ to predict $X_t$ along with one seasonal moving average $\varepsilon_{t-12}$. The moving average term will be represented by the recursive property of neural network. We have fitted the model in (5.3) using Minitab Statistical Software and obtained the results in Table 11.

### 5.3.2 Recurrent Neural Network for $CO_2$ Data

To fit the $CO_2$ data we decided to choose three input units in the input layer, two for two successive data points in the past to predict the next outcome. The third one is fed from the error observed by the neural network outcome as it is shown in Figure 24. We will discuss how the data is restructured to be trained correctly using the recurrent neural network in
Figure 23.: Partial autocorrelation function after taking one seasonal difference

Figure 24. In which, the third input for the first eleven patterns will be zero, since our Moving average starts from lag 12.

To train neural network in Figure 24, to fit the time series model, the data need to be restructured in forms of patterns. The patterns should be formed such that the column vector of time series data is changed to a matrix with three columns with each row representing a
For example, the first row will consist of
\[ X_1, X_2, \text{ and } (\varepsilon_{-9} = 0) \] to predict \( X_3 \),
next row will consist of
\[ X_2, X_3, \text{ and } (\varepsilon_{-8} = 0) \] to predict \( X_4 \),
and so on till we reach the row that consists of

\[ X_{10}, X_{11}, \text{and } (\varepsilon_1 = \hat{X}_3 - X_3) \] to predict \( X_{12}, \)

the following row consist of the pattern:

\[ X_{11}, X_{12}, \text{and } (\varepsilon_2 = \hat{X}_4 - X_4) \] to predict \( X_{13}, \)

and so on till you reach the end of the data. In our case we created 600 pattern for training proposes and 58 pattern to check forecasting. The linear function was used as activation function for output unit in Figure 24. We have created a MATLAB function that can be used for general number of input units.

The code on previous page titled: Restructuring of Time Series data for Neural Networks, is for a function that will form the patterns of data to be modeled with neural network. Function \textit{restructure} takes a number of input units (\( \text{nin} \)), number of output units (\( \text{nout} \)), training data column vector of size \( n \), number of units (\( \text{nloop} \)) in input layer that are fed from output units (recursive structure), and finally the number of lags that present the AR terms identified from first step of Box and Jenkins procedure. In lines 6 and 7 the function is obtained the sample size and the number (\( t \)) of previous observation that the predicted value depends on (as in case we have two AR terms 1 and 2). Although the code currently works for successive terms only but can easily be generalized to non-successive terms. In lines 9 and 10 we are initiating the matrix of patterns that acts as our predictor variables, and the target outcome vector, acting as our response,, and both are based on the number of input units and the number of output units, in our case we have \( \text{nin} = 3 \) and \( \text{nout} = 1 \). The rest of the code loops over the vector of data forming the patterns as explained earlier and the function returns the matrix of observations and its corresponding targets. The matrix of observations will have its last column all zeros, the last column is reserved for error obtained from prediction of the neural network for each single pattern, representing the SMA(1) with \( S = 12 \). Neural networks usually takes its input data from a matrix, where each column corresponds to one of the input units and each row represents a pattern. To preform learning online after each pattern, then we need to store each row of \( Xtrain \) in a
separate row vector and this is done by storing $X_{train}$ in a cell array as it is shown in the following code, same is also made for the target vector.

```matlab
for i = 1:n
    Xi{i,1}=Xtrain(i,1:nin);
    Ti{i,1}=YTrain(i,1);
end
```

Once we have the cell array ready, next initialize the network and start training it using Hybrid Monte Carlo Sampling as follows:

```matlab
%initialize neural network
nin= 3;
nout=1;
nhidden=10;
%Assigning values for hyperparameters
aw1 = [0.01, 0.01, 0.5]; %3 different distributions for input units
ab1 = 0.01; %hyperparameter for bias of first layer
aw2 = 0.01; %hyperparameter for weights fanning from hidden to ...
            %output layer
ab2 = 0.01; %hyperparameter for bias of hidden layer
beta = 50; %hyperparameter for data error
prior = mlpprior(nin, nhidden, nout, aw1, ab1, aw2, ab2);
net = mlp(nin, nhidden, nout, 'linear',prior,beta);
% Start loop to preform Online training
for j =1:600
    seed = 10; % Seed for random weight ...
    rng(seed,'v5uniform');
    rng(seed,'v5normal');
    % Set up vector of options for hybrid Monte Carlo.
```
nsamples = 500; % Number of retained samples.
options = foptions; % Default options vector.
options(1) = 1; % Switch on diagnostics.
options(5) = 1; % Use persistence
options(7) = 25; % Number of steps in trajectory.
options(14) = nsamples; % Number of Monte Carlo samples returned.
options(15) = 200; % Number of samples omitted at start of chain.
options(17) = 0.75; % Alpha value in persistence
options(18) = 0.002; % Step size.

w = mlppak(net);
% Initialise HMC
hmc('state', 42);
% Calling HMC after feeding network with each pattern (Online ... learning)
[samples, energies] = hmc('neterr', w, options, 'netgrad', net, ... 
    Xi{j}, Ti{j});
pred = zeros(size(Ti{j}));
Ed = 0;

for k = 1:nsamples
    w2 = samples(k,:);
    net2 = mlpunpak(net, w2);
    network11{1,k}=net2;
    Y = mlpfwd(net2, Xi{j});
    [Err, edata, eprior] = mlperr(net2, Xi{j}, Ti{j});
    % Average sample predictions as Monte Carlo estimate of true ...
    integral
    pred = pred + Y;
    Ed = Ed + edata;
end
pred = pred./nsamples;
Ed=Ed./nsamples;
Lines 2-12: the network is being structured and the prior object is built up. We used the automatic relevance determination to assign different distribution of weights for each of our three input units, line 6, error hyperparameter value is 0.5. Line 12: the neural network is set with the choice of input units, output units, linear function as activation function for output unit and prior structure. Then online learning is applied by calling the Hybrid Monte Carlo sampling for each record of the cell array $X_i$ with corresponding targets $T_i$. Errors are computed after each pattern and input cell array is updated, lines 46-50. In the current analysis the error obtained from the first pattern of $X_i$ is returned and stored in the 12th pattern of $X_i$ completing the third input information, corresponding to SMA(1). Lines 15-44 are taken from Nabney algorithm for HMC implementation, more details can be found in his book [62]. We sampled 500 samples for each pattern using HMC.

5.4 Results

As we mentioned in Chapter 2 the use of Hybrid Monte Carlo Sampling requires several trails to obtain initial values for hyperparameters. We will present two graphs to illustrate the difference between the initial values of hyperparameters. After training the neural network with the previously mentioned procedure we obtained a mean square error of $0.000616$, by comparing it with the one obtained from using SARIMA model in Table 9 ($\text{MSE} = 0.0864$) we can tell that the recurrent neural network model outperformed the SARIMA model. After we compared SARIMA and the recurrent neural network trained with HMC, we also compare their forecasting accuracy. As we mentioned in the data section that we left five years for determining the forecasting accuracy.
Figure 25.: Comparison between forecasting accuracy of RANN with (0.01 hyperparameters) and SARIMA

Figure 26.: Comparison between forecasting accuracy of RANN and SARIMA
In Figure 25, Black dots are for observed $CO_2$ average monthly emission, while red dots from the SARIMA model. The green series is the forecasting from (RANN) trained with initial values of hyperparameters of the input units all equal to 0.01. We have realized that the third input unit that corresponds to the error needs to have higher value of hyperparameter, meaning errors have less weight compared to $X_1$ and $X_2$ in forecasting $X_3$. The final neural network we developed is the one trained with initial values of hyperparameters of 0.01 for two data inputs and 0.5 for the error input.

Figure 26, displays the final outcome of the best fit neural network. The red series here represents the forecasting of the RANN, it is either similar to SARIMA model in some cases and in others it outperform it.

5.5 Conclusion

Our main objective in this Chapter is to introduce a general framework for utilizing neural network in time series analysis. Step 1 in achieving our objective, we proposed a new methodology for utilizing ANN in fitting stationary time series data. The proposed method resulted in better performance compared to the popular ARIMA model. Step 2 is to model non-stationary time series data using same method (Taking into account the high capability of neural network in fitting non-linear patterns). We would like to mention that we first tried using the evidence HMC method (The new proposed Bayesian learning presented in chapter 4) in training the recurrent neural network in Figure 24, but we could not observe any accepted samples from the HMC after re-estimating the hyperparameters using the evidence. Thus, the evidence procedure do not perform well with online learning.

The secondary objective of the current study, is to better understand the behavior of $CO_2$ in the atmosphere. Additionally form a model that predicts carbon dioxide emission from the various source in Figure 17. Also, use the automatic relevance determination to rank these sources based on their percentage of contribution to carbon dioxide emission.
5.5.1 Contributions

The development of ANN models for time series analysis revealed some very important findings that can be summarized as follows:

1. The use of ARMIA model identification step is important in determining the number of input units in neural networks used to model time series data.

2. Best neural network structure to model time series data is recurrent (recursive) neural networks.

3. The Hybrid Monte Carlo Sampling proposed by Neal, train perfectly recurrent neural networks.

4. Carbon dioxide emission in the atmosphere can be better explained through a recurrent neural network model trained with Hybrid Monte Carlo Sampling compared to popular ARIMA model.
Chapter 6
Future Work

Our future work consist of two main goals, first is to introduce Bayesian techniques to other artificial intelligence methods such as genetic algorithm, among others. The second is to use artificial neural network with Bayesian techniques to find a solution for other vital problems our society is facing.

6.1 Neural Network models in Cancer research

Our proposed model can be used with other cancer diseases, such as lung cancer, breast cancer, brain cancer, among others. For example, in survival studies, treatment options can be added to risk factors, and by the use of automatic relevance determination, doctors can suggest patients their suitable course of treatment that would make them survive longer. Of course these kind of models need to be trained on more specialized data than the one with used in chapters three and four. Additional information needed to be able to come up with decisions on individual bases. Same idea can be extended to all cancer types.

In breast cancer especially (and for other types too), studying the factors that causes cancer has great attention among medical doctors now a days. An artificial neural network model can be built to study the effect of several factors on the incident of cancer. The use of ARD will help tremendously in finding the most contributing factors to the incident of cancer.
6.2 Neural Network for Time Series Forecasting

The area of building neural network models for time series forecasting is still open. The challenges that exist need more research to find a suitable answer for it. Our future work in this area is to build a framework to use artificial neural network in time series forecasting. Special attention will be given to non-stationary time series, due to the wide existence in real life of such time series. We will be working on a time series model that predicts carbon dioxide emission in the atmosphere conditioning on the source of emission. The use of automatic relevance determination will help in ranking the sources of emission based on their contributions, allowing policy makers and government agencies to better control these sources to reduce or control carbon dioxide emission in the atmosphere. We can extend this study by examining the regional effect and use one of neural network advantages in the capability of predicting several variables at once. A similar neural network architecture to DHANN-CR can be used to predict the carbon dioxide emission of several regions. The input layer will present the different sources of emission, and the output layer represents the different regions.
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