NEURAL NETWORK OPTIMIZATION USING SHUFFLEDFROG ALGORITHM FOR SOFTWARE DEFECT PREDICTION

1REDDI. KIRAN KUMAR, 2S.V.ACHUTA RAO

1Department of Computer Science, Krishna University, Machilipatnam, India- 521001  
2Research Scholar, Krishna University, Machilipatnam, India- 521001  
E-mail: 1kirankreddi@gmail.com, 2achutaraosv@gmail.com

ABSTRACT

Software Defect Prediction (SDP) focuses on the detection of system modules such as files, methods, classes, components and so on which could potentially consist of a great amount of errors. SDP models refer to those that attempt to anticipate possible defects through test data. A relation is present among software metrics and the error disposition of the software. To resolve issues of classification, for the past many years, Neural Networks (NN) have been in use. The efficacy of such networks rely on the pattern of hidden layers as well as in the computation of the weights which link various nodes. Structural optimization is performed in order to increase the quality of the network frameworks, in two separate cases: The first is the typically utilized approximation error for the present data, and the second is the capacity of the network to absorb various issues of a general class of issues in a rapid manner along with excellent precision. The notion of Back Propagation (BP) is quite elementary; the result of neural networks is tested against the desired outcome. Genetic algorithms (GA) are a type of search algorithms built, based on the idea of natural evolution. A neural network using Shuffled Frog Algorithm for improving SDP is proposed. Keywords: Software Defect Prediction (SDP), Neural Network (NN), Back Propagation (BP), Genetic Algorithm (GA) and Shuffled Frog Leaping Algorithm (SFLA)

1. INTRODUCTION

This Software Defect Prediction (SDP) holds an important place in the domain of software quality and dependability. Software faults may be defined as errors, mistakes or defects in computer programs or systems which lead to false or unusual outcomes, or leading the software to perform in unexpected ways. When a software module consists of too many flaws that gravely inhibit execution, it is said to be fault-prone. The procedure of identifying such faulty modules in a particular software is SDP. The conventional methods of discovering faults in a software are code review, unit testing, integration testing and system testing. But, when a particular software’s dimensions increase by way of the number of coding statements and complexity, the process of discovering as well as repairing errors becomes harder as well as costly with regard to computation; through the usage of advanced testing and evaluatory processes. Furthermore, Boehm gleaned that the discovery and repair of issues once a particular software has been delivered, is costlier, money and effort-wise, when compared to repairing the fault in the initial phases of the life cycle of the software. Identification of fault-prone software modules allows professionals to focus their effort and capabilities on the problematic regions of the software under construction [1 & 2].

The NN framework focuses on the prediction of the quality of object oriented software through estimation of the amount of flaws as well as the amount of statements altered in each class. Object-oriented metrics along with conventional complexity metrics are both used here. Object-oriented metrics includes inheritance related measures, cohesion measures, coupling measures, memory allocation measures as well as the usage of Ward Neural Network and General Regression Neural Network (GRNN) to increase the accuracy of predictions regarding the quality of software [3].

Neural network consists of back propagation network with several activation functions. To achieve more accurate predictions, they are used on hidden layer slabs so as to identify dissimilar characteristics in a design analysed using a network. A Gaussian function is employed in a
hidden slab for the identification of features in the mid-range of the information and a Gaussian complement is employed in a different hidden slab for the identification of characteristics for the upper and lower limits of the data. Hence, the outcome layer will receive dissimilar “views of the data”. Compounding the two characteristics sets in the outcome layer points to more accurate predictions. The other structure picked is the GRNN, which is a memory-based network that gives an estimation of continuous variables and comes together at the base (linear or non-linear) regression surface. As it is a one-pass learning algorithm possessing a very parallel architecture, the barest minimum of information in a multidimensional measurement environment; the algorithm allows for seamless movement from one ascertained value to the next.

Discovering the appropriate algorithm for the modeling of software elements to various stages of error severity is a crucial task [4]. A model grounded in the Biological Neural Network, is the Artificial Neural Network (ANN) and is typically referred to as merely Neural Network. Artificial neurons, described as nodes are interlinked for the purpose of constructing ANNs. The structure of neural networks is crucial for the performance of a specific operation. Certain neurons are placed specifically for obtaining input from the external environment. Because they are not linked to one another, the structure of the neurons is as a layer, referred to as an input layer. The neurons present in the input layer provide certain output which functions as the input for the adjacent layer [5].

Two types of NN structures that may be combined are feed-forward networks and recurrent/recursive networks. Networks that possess completely linked layers, like the multi-layer perceptron, and networks with convolution and pooling layers are feed-forward networks. Every single network acts as a classifier, however each has a distinct strength of its own. Completely linked feed-forward NNs are non-linear learners which may, at the most, be utilized as an alternate to linear learners.

Because the network is non-linear and has the capacity to incorporate previously trained word embeddings, it results in exceptional classification precision. A set of studies achieved better syntactic parsing outcomes through the mere replacement of the linear model of the parser with a completely linked feed-forward neural network [6]. The networks are initially directed to simulate the behaviour of active and passive circuits. These networks, now trained are called as neural network models. These may now be utilized in high level simulation and designs, and are shown to produce the fastest and most precise responses to the task because of the information they acquired during the process of training.

When compared to traditional approaches; such as numerical modeling techniques, that are exceptionally costly with regard to computation, or analytical techniques that could be hard to attain for newly obtained gadgets, or empirical modeling techniques which possess a large range and poor precision, neural networks prove to be more efficient and may be employed for a great number of applications as well as modeling techniques [7].

A three-layer feed-forward NN is proven to be capable of approximating a non-linear continuous function to a random precision. They are employed in plenty of fields like prediction, system modeling and control. Because of the specific structure, NNs are excellent in absorbing learning algorithms like Genetic Algorithm (GA) as well as Back Propagation. Typically, there are two learning stages for an NN: Firstly, a network architecture is determined with a specific set of inputs, hidden notes as well as outputs. Secondly, an algorithm is picked to execute the training procedure.

But, it is not a must for a stable architecture to give its best results during the learning stage. A small network might be incapable of performing well due to the restrictive processing potential. A big network, though, might possess repetitive links and the execution cost remains heavy. Both constructive and destructive algorithms may be utilized in order to attain the network architecture. Constructive algorithms begin with the small networks while the destructive algorithm begins with the big networks.

Hidden layers, nodes and links are later eliminated to compress the network in a dynamic manner. The pattern of a network architecture may be modeled as a search issue. Gas are used to achieve the results. Pattern-classification approaches may be seen designing networks [8]. Fitness testing consumes too much time in various real-world applications of evolutionary calculation. A way to decrease the computation time is through the replacement of the fitness function with an approximate framework with lesser computational cost. These frameworks are called meta-models or substituted in optimization. A model for evolutionary optimization uses approximate models for applications in order to achieve optimization.

In this model, the approximate model is compounded with the initial fitness function to
manipulate the evolutionary procedure; that is, to
determine how frequently the approximate model
ought to be utilized as opposed to the initial fitness
function, to make sure of the merging of the
evolutionary algorithm to the right optimum of the
initial issue as well as to decrease calculation cost
to the highest degree. The better the quality of the
framework, it should replace the initial fitness
function more frequently. The learning capacity of
the NNs is especially crucial for the time online
learning is required to be executed at the time of
optimization.

Optimization of the structure of NNs is
performed before they are used for the purpose of
fitness testing in evolutionary pattern optimizations
[9]. GAs are based on the evolution of people. In a
certain environment, people who are more suited to
it, are fit to survive and pass down their genes to
future generations; and less adaptable humans die.
The goal of GAs is to utilize basic representations
to code complicated architectures and basic
computations to better those architectures. GAs are
hence, defined by their representations and
operators.

In a GA, a single chromosome is
correlated to a binary string. The bits in each string
are named as genes and their changing values are alleles. A set of single chromosomes are a
population. Fundamental genetic operators
incorporate reproduction, crossover and mutation
[10]. GAs and artificial NNs are methods for
learning and optimization, which have been based
on biological mechanisms. NNs utilize inductive
learning and typically need samples, whereas GAs
adopt deductive learning and need objective
evaluation function.

This paper suggests neural network
structure optimization using Shuffled Frog
Algorithm. Section 2 deals with literature related to
this work, section 3 reveals the methods used in the
work, section 4 deals with results and discusses
obtained results and finally section 5 concludes the
work.

2. LITERATURE REVIEW

Yang et al., [11] researched this issue and
proposed a network pruning algorithm grounded in
sparse representation, called SRP. The suggested
method begins with a sizeable network, extracted
crucial hidden neurons from the initial architecture
utilizing a forward selection criterion which lessens
the residual output error. Additionally, the proposed
algorithm possesses no restrictions of network kind.
The efficacy of the suggested technique was tested
on various benchmark data sets. The outcomes
reveal that SRP is more accurate as opposed to
other methods.

Fan and Wen [12] proposed a novel mind
evolutionary algorithm (MEA) grounded in the
basic MEA model in order to optimize the NN’s
architecture and weights, where like taxis and
different operators of architecture optimization are
planned. Using such taxis operators, the local
optimum is discovered and then overshooting the
constriction of local range by the usage of
dissimilation operators, the global optimum is
attained in the global solution environment. The
outcomes of the simulations revealed that the
approach was efficient and accurate.

Mohseni and Tan [13] suggested a novel
mixed training algorithm comprising error back-
propagation (EBP) as well as variable structure
systems (VSS) for the optimization of parameter
updating of NNs. In order to optimize the number
of neurons in the hidden layer, a new term
dependent on the outcome of the hidden layer is
combined to the cost function as a penalty term so
as to optimally utilize hidden blocks connected to
weights correlating to every single block in the
hidden layer. Additionally, apart from analysing the
imposed dynamics of the EBP method, the global
reliability of the mixed training methodology and
restrictions on the design variables were observed.
The suggested method has plenty of benefits
including assured convergence, increased strength
as well as decreased sensitivity to preliminary
weights of the network.

Shahriari-kahkeshi and Askari [14]
proposed the plan of a recurrent neural network
(RNN) trained Shuffled Frog Leaping Algorithm
(SFLA) for the purpose of identifying and
following control of non-linear continuous stirred
tank reactors (CSTR). RNNs were applied to
approximate unknown dynamic systems and SFLAs
were employed for the training and optimization of
connection weights of the RNNs. The suggested
control design used neural control system synthesis
was employed in the closed-loop control system for
accurate control inputs. The outcomes revealed that
the RNN-SFLA controllers have extraordinary
dynamic responses and also that they adapted
excellently to changes.

Feng et al., [15] suggested a wavelet
neural network (WNN) sound source model
grounded in the SFLA. The SFLA is employed for
the optimization of weights and thresholds of
WNN, attain preliminary weights and thresholds
having particular periodicity and later train
WNNs. This overpowers certain failings of NNs
that have slow searching speed. Outcomes revealed
that the novel WNN algorithm has improved convergence rate, positioning accuracy, favourable application prospects and future research value.

Yu et al., [16] utilized SFLA as a basis for NNs in speech emotion detection. SFLA is utilized to train arbitrary preliminary information, optimize link weights and thresholds of NNS with rapid network convergence speech. Outcomes revealed that the SFLA network shows excellent detection rate in the field.

Ye et al., [17] used the SFLA with pseudo code and flow charts to ease its execution. The researches utilized SFLA to optimize weight and threshold value of BP networks. Experimental results revealed that SFLA outperformed GA in optimizing BP networks’ weight and threshold values that are utilized in non-linear function fitting.

Che et al., [18] proposed a brilliant technique of swarm neural networks (SNN) for dealing with equalities-constrained non-convex optimization issues. The suggested technique dealt with the issue in two sections that compound local searching capacity of one-layer RNN and global searching capacities of SFLA. Firstly, an RNN framework grounded in typical non-convex optimization was issued. Also, dependent on the SFLA model, NNs are considered as frogs that ought to be split into many memeplexes and develop individual differential equations in order to find an accurate local solution. And lastly, numerical samples with outcomes are provided to display the efficacy and excellent features of the suggested method resolving non-convex optimization issues.

3. METHODOLOGY
In this section, we discuss neural network utilizing BP, neural network using GA and proposed neural network using SFLA.

3.1. Neural Network using Back Propagation
A BPNN learns by utilizing the generalized delta rule in a two phase propagate-adapt cycle. The first randomly drawn weights are applied to the NN in order to create a starting point in which to initiate the search. During the starting point, approximates are compared with the expected outcome, and an error is calculated for each of the observations. The error for the first observation is computed by decreasing an amount equal to the estimated value from the true value. Total of the squared errors is typically utilized as objective function [19].

One of the most commonly used NN algorithms is BP and it may divided into four major stages. After choosing the weights of the network arbitrarily, the BP algorithm is utilized to calculate the required rectifications. The algorithm may be divided into the below mentioned four steps [20]:

i) Feed-forward computation
ii) Back propagation to the output layer
iii) Back propagation to the hidden layer
iv) Weight updates

The best technique for carrying out supervised learning is the BP algorithm. BP has been employed in many learning tasks and has come out as the standard algorithm for the training of multi-layer perceptron networks. BP is grounded in gradient descent as it is q conjugate gradient. It typically utilizes a least-square optimality criterion, determining a technique for computing the gradient of the error specifically for the weights for a certain input, through the propagation of the error backwards through the network. Error BP is essentially a search procedure that tries to decrease a whole network error function like the sum E of the squared error of the network output over an ensemble of training input/output pairs:

\[ E = \frac{1}{2} \sum_{j=1}^{m} (t_j - o_j)^2 \]

where \( t_j \) is the target and \( o_j \) is the actual \( j^{th} \) output of the network.

The back propagation algorithm determines two sweeps of the network: firstly, a forward sweep from the input layer to the output layer, and next, a backward sweep from the output to the input layer. In the first stage, input vectors are propagated through the network to give outputs of the network. The backward step is alike to the forward one, except that error values calculated, are propagated back through the network, so as to evaluate how the weights ought to be altered during the training phase. The fundamental process for the BP algorithm is reported in Figure 1.

The training is monitored through a target pattern corresponding with an input pattern. A pattern is displayed to the network and an error vector is computed to analyze how the weights ought to alter; the procedure is repeated for every pattern. An epoch is a complete cycle for each pattern. The patterns are continually displayed to the network, epoch after epoch, and training continues until the alterations in the absolute value of the the averaged squared error falls to within some tolerance between one epoch and the next [21].
Initialize network weights randomly

while not termination condition do
    Assign as net input to each unit in the input layer its corresponding element in the input vector.
The output for each unit is its net input.
    Calculate network output by forwarding input signals in the network.
    Calculate the error of each output neuron
    for all hidden neurons do
        calculate weights updates
        propagate the error back through the network.
    end for
    Update weights of the network.
end while

Figure 1: Pseudo code of the Back Propagation algorithm

3.2. Neural Network Using Genetic Algorithm

Genetic Algorithms is a way of approaching machine learning by appropriating behaviour of the human gene as well as Darwin’s theory of natural selection. GA is a part of Evolutionary Algorithms which provide solutions depending on the methods more generally discovered in nature like mutation, selection, crossover etc.

Genetic Algorithms are applied starting with an individual population that is typically represented in the form of trees. A probable solution is represented by each tree or say chromosome in this case. Nodes on the tree denote certain traits that relate to the issue for which the solution is being searched. Collectively, the set of possible solutions to the problem is (represented by the chromosomes) as known as the population [22].

GA’s appear in the picture when it is required to solve issues which can have various solutions. Here, genetic algorithms are utilized to cluster the classes delineated according to object oriented metrics into subsystems or typically called components of software. As described earlier, GA utilizes an approach similar to Darwin’s “Survival of the Fittest” theory or natural selection. Why this approach is being deliberated upon is due to the large solutions set that provides a many possible solutions to an issue.

When employing a GA in a problem, a couple of implications are made. They are mentioned below

- A fitness function must be present for evaluating if a solution is a possible or not.
- When a solution is found, a representation of it must be made by a chromosome.
- Which genetic operators will be employed must be decided.

Furthermore the definition of a solution in this case would be one which would be both complete as well as valid. In terms of a representation, it may be assumed that the possible solutions have been encoded in the solutions space.

The GA is a search technique grounded in the idea of evolution, and particularly with the notion of the survival of the fittest. The usage of GA on NNs make a hybrid neural network where the weights of the neural network are computed through GA method. From all the search spaces of the probable weights, the GA will create new points of the probable solution.

The first stage to compute its values, is to delineate the solution field with a genetic representation (problem encoding) as well as a fitness function to define the better solutions. These two components of a GA are the only problem dependent on the GA method. When initial populations of elements (genotype or chromosomes) have been generated, the methods utilized by these algorithms in order to reach a solution to the issue are connected to evolutionary theories:

i. Selection → few chromosomes of the current population are chosen for the breeding of a new generation. A small set is chosen arbitrarily and the remaining are chosen based on how they suit a fitness function.

ii. Genetic operations → when the initial chromosomes have been delineated by those that fit more appropriately with the fitness function, the remaining population will be built through genetic operations. Fatnesses of the new chromosomes will additionally, be verified and compared to the worst chromosomes of the previous generation to determine who remains in the population.

Mutation: Altering one or more gens of few chromosomes of the entire set & Crossover: Crossing two or more chromosomes in the set amongst themselves.

A pseudo-code for Genetic algorithm is [23]:

1. Creation of the initial population
2. while (solution)
   (a) Evaluate the fitness of all the chromosomes of the population
   (b) The best chromosomes will be selected to reproduce, using mutation and crossover.
   (c) Substitute the worsts chromosomes of the previous generation by the new produced chromosomes.

Figure 2: Simple Genetic Algorithm
Genetic algorithms are algorithms for optimizing and machine learning grounded loosely in many features of biological evolution. They require five components [24]:

i. A method of coding solutions to the problem on chromosomes.

ii. An evaluation function which returns a rating for each chromosome given to it.

iii. A method to initializing the population of chromosomes.

iv. Operators that may be applied to parents when they reproduce to alter their genetic composition. Standard operators are mutation and crossover.

v. Parameter settings for algorithms, operators etc.

3.3. Proposed Neural Network Using Shuffled Frog Algorithm

ANN technology, inspired by neurobiological (the brain’s structure) theories of massive interconnection and parallelism, is perfect for undertakings like pattern recognition and discrimination. One of the first researchers to extend the application of neural networks into optimization was Hopfield (1984), and it has been successfully applied to a variety of combinatorial optimization problems. The ANN learns to solve a problem by developing a memory capable of associating a large number of input parameters with a resulting set of outputs or effects. The ANN develops a solution system by training using examples given to it [25].

SFLA is a meta heuristic for solving optimization problems. SFLA is a cooperative search metaphor that is population based and is occasioned by natural memetics. Memetic evolution by way of influencing ideas from one individual to the next in a local search is employed by the algorithm. Theoretically, this is alike to Particle Swarm Optimization (PSO). A shuffling technique allows for the exchange of data amongst local searches, thereby attaining global optimum [26].

In the SFLA, the population comprises of frogs of like structures. Every frog denoted one solution. The total population is split into various subgroups. Different subgroups may be considered as different frog memes. Every subgroup attempts a local search. They can correspond with one another and better their memes among local units. Once a pre-determined number of memetic evolution stages are done, data is transferred between memeplexes in a shuffling manner. This shuffling makes sure that the cultural evolution in the direction of any specific interest is completely unbiased. The local search and the shuffling procedures alternate till the stopping criteria is satisfied. [27].

The SFLA is based on observing, imitating, and modeling the behavior of a number of frogs during the process of searching for the location that possesses the most amount of available food. The primary benefit of SFLA is its fast convergence speed. The SFL Emerges the benefits of both the genetic-based memetic algorithm (MA) and the social behavior-based PSO algorithm.

SFLA is a population based arbitrary search algorithm occasioned from nature meme tics. In the SFLA, a set of possible solutions determined by a set of frogs that is divided into several communities known as memplexes. Every frog in the memeplexes performs a local search. Inside of each memplex, the single frog’s behavior may be influenced by other frogs and evolution through a procedure of memetic evolution will occur. After a particular number of memetic evolution stages, the memeplexes are pressured to join together and novel memeplexes are developed through a shuffling process.

The flowchart of Shuffled frog leaping algorithm is illustrated in Fig. 1. The various steps are as follows:

1. The SFLA includes a population ‘P’ of possible solutions, determined by a set of virtual frogs(n).
2. Frogs are arranged in descending order as per their fitness and then divided into subsets called as memeplexes (m).
3. Frogs $i$ is expressed as $X_i = (X_{i1}, X_{i2}, ..., X_{iS})$ where S represents number of variables.
4. Within each memplex, the frog with worst and best fitness are identified as $X_w$ and $X_b$.
5. Frog with globe best fitness is defined as $X^g$.
6. The frog with worst fitness is improved according to the following equation.

$$X_{new} = X_{old} + \Delta_i (-D_{max} \leq \Delta_i \leq D_{max})$$

where rand is an arbitrary number in the range of $[0,1]$; $\Delta_i$ is the frog leaping step size of the i-th frog and $D_{max}$ is the maximum step allowed.
is better than the current one, $X_s$ will be accepted. If it isn’t improved, then the calculated $X_s$ will be replaced by $X_s$. If there is no possibility of improvement in the case, a new $X_s$ will be generated arbitrarily. Reiterate the update operation for a certain number of iterations [28]. The flowchart of SFLA is given below in Figure 3.

4. RESULTS & DISCUSSION

In this section, the classification accuracy, precision, recall and F measure are evaluated from table 1 to 4 and figure 4 to 7.

Table 1: Classification Accuracy

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Classification Accuracy %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neural Network - BPP</td>
<td>92.41</td>
</tr>
<tr>
<td>Neural Network - GA</td>
<td>94.48</td>
</tr>
<tr>
<td>Neural Network - proposed Shuffled frog</td>
<td>95.86</td>
</tr>
</tbody>
</table>

Figure 4: Classification Accuracy

From table 1 and figure 4, it is observed that the Neural Network method improved classification accuracy by 2.22%, 1.45% and 3.66% when compared with the BPP, GA and proposed Shuffled frog methods.

Table 2: Precision

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Precision</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neural Network - BPP</td>
<td>0.8677</td>
</tr>
<tr>
<td>Neural Network - GA</td>
<td>0.8979</td>
</tr>
<tr>
<td>Neural Network - proposed Shuffled frog</td>
<td>0.9297</td>
</tr>
</tbody>
</table>
From the table 2 and figure 5, it can be observed that the Neural Network method improved precision by 3.42%, 3.48% and 6.9% when compared with the BPP, GA and proposed Shuffled frog methods.

**Table 3: Recall**

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Recall</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neural Network - BPP</td>
<td>0.8787</td>
</tr>
<tr>
<td>Neural Network - GA</td>
<td>0.9213</td>
</tr>
<tr>
<td>Neural Network- proposed Shuffled frog</td>
<td>0.9297</td>
</tr>
</tbody>
</table>

**Figure 6: Recall**

From table 3 and figure 6, the observation is made that the Neural Network method improved recall by 4.73%, 0.91% and 5.64% when compared with the BPP, GA and proposed Shuffled frog methods.

**Table 4: F Measure**

<table>
<thead>
<tr>
<th>Methodology</th>
<th>F Measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neural Network - BPP</td>
<td>0.8731</td>
</tr>
<tr>
<td>Neural Network - GA</td>
<td>0.909</td>
</tr>
<tr>
<td>Neural Network- proposed Shuffled frog</td>
<td>0.9297</td>
</tr>
</tbody>
</table>

**Figure 7: F Measure**

From the table 4 and figure 7, it can be observed that the Neural Network method improved F measure by 4.03%, 2.25% and 6.28% when compared with the BPP, GA and proposed Shuffled frog methods.

4. **CONCLUSION**

NN are widely used for prediction. The weights in various layers of the network are optimized using a BP or GA. To enhance the performance of the NN, a novel method based on SFLA for optimizing the NN for improving the SDP is proposed in this paper. The shuffled frog-leaping algorithm with local search can guarantee the feasibility of the updated solution. The proposed method was evaluated for classification accuracy, precision, recall and f measure for classifiers. It is seen that the NN method improved classification accuracy by 2.22%, 1.45% and 3.66% when compared with the BPP, GA and proposed Shuffled frog methods.
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