Application of Neural Networks Technique for Predicting of Abrasiveness Characteristics of Thermal Coal

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Abstract—In this work, Neural Networks (NN) approach was proposed to deal with abrasiveness behavior of thermal coal. Back-propagation neural network (BPNN) and Generalized regression neural network (GRNN) techniques were employed to assess the Abrasive index (AI) of coal to mineral. The multivariate statistical results revealed that the BPNN and GRNN models were successfully developed to model the abrasiveness characteristics of thermal coal with the coefficient of determination, $R^2 = 0.9003$ for BPNN and $R^2 = 0.937$ for GRNN. These good results indicated that the NN techniques were capable of accurately modeling the abrasiveness characteristics of coal.

Keywords: Abrasive index, back-propagation neural network, coal, generalized regression neural network.

I. INTRODUCTION

Coal is a combustible, organic rock, which is composed mainly of carbon, hydrogen and oxygen. Coal is primarily used as a solid fuel to produce electricity and heat through combustion. When coal is used for electricity generation, it is usually grinded to an efficient burnable size in a mill and then burned in a furnace with a boiler. During grinding, frictions occur, cause abrasive wear or erosion on the critical components and thereby affect the performance of power plant. It is therefore important to assess the relative abrasion characteristics of thermal coal by selecting the right type of materials for grinding and burning of the coal [1]. To study the abrasion of coals, an index namely abrasion index (AI) which has been used to assess the abrasive nature of the thermal coal was established first by Yancey, Geer and Price (YGP) in 1951 using YGP test rig. Over the years there have been some modifications to this method by both mining houses and coal users which have resulted in inconsistent and conflicting results [2]. AI was then measured using different methods, which include the AI tester pot using four iron blades as cutting elements [3]. Reference [4] revealed that two and three-body abrasive wear is not only concerned with hard material, as material of less hardness than the concerned metal blades can still cause material wear. Reference [3] revealed that the effects of particles that are less hard than the cutting blades are inconsistent in comparison to harder minerals. To improve the prediction of abrasiveness of thermal coal, it is important to understand the nature and properties of the mineral matters in a coal that would contribute to abrasive wear. Most of the empirical equations available in literature [5]-[9] for predicting AI of coal, are based on linear assumptions which may lead to erroneous estimations and do not take into consideration most of the relevant factors. To achieve this, Neural Network (NN) based predictive techniques was suggested to understand the nonlinear relationships and thereby achieving ability to predict accurately. Reference [10] used non-linear multivariable regression and NN to find the correlation between Hardgrove grindability index and the proximate analysis of chemise coals. Reference [11] also used NN to studies the relationship between petrography and grindability for Kentucky coals. To our knowledge, this is the first time that NN has been used to predict the abrasive index of the coal. The objective of this study is to investigate the possibility for the prediction of abrasiveness characteristics of thermal coal abrasive index using neural network.

II. MATERIALS AND METHODS

A. Experimental Data

The coal samples used in this study was sourced from different colliers in South Africa. These samples were analyzed for both their chemical and physical characteristics based on the premise that they can contribute information that can make it possible for this study to adequately reveal coal constituents that cause abrasion during grinding. The abrasion index tester pot was used to determine AI of the coal samples. A Perkin-Elmer simultaneous thermogravimetric analyzer (STA 6000) equipped with Pyris manager software was used to determine the proximate analyses that gives information about moisture and ash percentage, and petrographically determined minerals in the coal samples.

A multivariate statistical analysis was conducted, and from this analysis it was determined that four variables (Ash, Quartz, Pyrite and Moisture) were significant contributors to AI models.

B. Back-Propagation Algorithm (BPNN)

In this study, the first step was done to scale the inputs and targets within the range 0 and 1 in case the higher values would drive the training process and mask the
contribution of lower valued inputs, as well as to perform a principal component analysis to eliminate redundancies of the data set. In the second steps, the data set was divided into training, validation and testing subsets. The training set was

used for updating the network weights and computing the gradient. The validation set was used for improving generalization. The testing set was used for validating the network performance. The data in each subset were selected randomly, and then a network was created. A total of ten training algorithms were conducted to simulate the test data. The performances of the network in each training process and the best network with the highest prediction performances were recorded.

C. Generalized Regression Neural Network (GRNN)

The GRNN falls into the category of probabilistic NN that can solve any function approximation problem if sufficient data are available [12]. The additional knowledge needed to get the fit in a satisfying way is relatively small and can be done without additional input by the user. This makes GRNN a very useful tool to perform predictions of system performance in practice [12]. The expected value of the output \( y \) given the input vector \( x \) is given by:

\[
E\left( \frac{y}{x} \right) = \frac{\int y f(x,y)dy}{\int f(x,y)dy}
\]

(1)

when the density \( f(x,y) \) is not known, it must usually be estimated from a sample of observation of \( x \) and \( y \). The probability estimator \( g(x,y) \) in (2) is based on sample values \( x' \) and \( y' \) of the random variables \( x \) and \( y \):

\[
g(x,y) = \frac{1}{2\pi\sigma_1\sigma_2\sigma_3\sigma_4} \exp \left\{ -\frac{1}{2\sigma_1\sigma_2} (x-x')^2 - \frac{1}{2\sigma_3\sigma_4} (y-y')^2 \right\}
\]

(2)

Where \( n \) is the number of sample observations, and \( p \) is the dimension of the vector variable \( x \). A physical interpretation of the probability estimate \( g(x,y) \) is that it assigns sample probability of width \( \sigma \) (smoothing factor) for each sample \( x' \) and \( y' \), and the probability estimate is the sum of those sample probabilities [12]. The squared distance \( D \) between the input vector \( x \) and the training vector \( x' \) is defined as:

\[
D^2 = (x - x')^T (x - x')
\]

(3)

And the final output is determined by performing the integrations in (4). This result is directly applicable to problems involving numerical data.

\[
g = \frac{\sum_{i=1}^{n} y' \exp\left( -\frac{D^2_{i}}{2\sigma^2} \right)}{\sum_{i=1}^{n} \exp\left( -\frac{D^2_{i}}{2\sigma^2} \right)}
\]

(4)

The smoothness parameter \( \sigma \), considered as the size of the neuron’s region, is a very important parameter of GRNN [12]. When \( \sigma \) is large, the estimated density is forced to be smooth and in the limit becomes a multivariate Gaussian with covariance \( \sigma^2 I \) (\( I \) = unit matrix), whereas a smaller value of \( \sigma \) allows the estimated density to assume non-Gaussian shapes, but with the hazard that wild points may have a great effect on the estimate [12]. Therefore, a range of smoothing factors should be tested empirically to determine the optimum smoothing factors for the GRNN models [12]. In this study, NN toolbox 5.1 in Matlab 7.4 (R2007a) was used to develop the NN models.

D. Performance Indicators

The root mean square error (RMSE), mean absolute error (MAE), and coefficient of determination (R\(^2\)) between the modelled output and measures of the training and testing data set are the most common indicators to provide a numerical description of the goodness of the model estimates. They are calculated and defined according to (5), (6) and (7), respectively [13]:

\[
RMSE = \left[ \frac{1}{N} \sum_{i=1}^{N} (O_i - P_i)^2 \right]^{1/2}
\]

(5)

\[
MAE = \frac{1}{N} \sum_{i=1}^{N} |O_i - P_i|
\]

(6)

\[
R^2 = 1 - \frac{\sum_{i=1}^{N} (P_i - A)^2}{\sum_{i=1}^{N} (O_i - A)^2}
\]

(7)

where \( N \) = number of observations, \( O_i \) = observed value, \( P_i \) = predicted value, \( A \) = average value of the explained variable on \( N \) observations.

RMSE and MAE indicate the residual errors, which give a global idea of the difference between the observed and predicted values. \( R^2 \) is the proportion of variability in a data set that is accounted for by a model. When the RMSE and MAE are at the minimum and \( R^2 \) is high (\( R^2 \geq 0.80 \)), a model can be judged as very good [14].

III. RESULTS AND DISCUSSION

A. BPNN Model Development

The development of a good BPNN model depends on parameters determined using error methods (RMSE, MAE and \( R^2 \)). Three important aspects (number of layers, neurons in the hidden layer and the type of activation functions for the layers) must be decided on the BPNN structure. A three-layer BPNN was constructed to determine if its prediction performance was superior to a two-layer network. Unfortunately, the results were almost the same. It is worth noting that the bigger network structure would need more computation and could cause overfitting of the data [15]. To optimize the network, two neurons were used as an initial guess in the hidden layer. By increasing the number of neurons, the network gave several local minimum values and different error values were obtained for the training set.

Table I gives the dependence between the neurons numbers in hidden layer and RMSE for the Levenberg-Marquardt Algorithm selected as the best BP algorithm.

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In Table I a gradual decrease was observed in the RMSE. With 6 neurons in hidden layer, the MSE reached its minimum value of 0.000257. Hence, the neural network containing 6 neurons in hidden layer was chosen as the best case. When the number of neurons exceeded 6 in hidden layer, the MSE showed an increase at 7 and 8 neurons. The networks are sensitive to the number of neurons in their hidden layers. The optimum number of neurons required is problem dependent, being related to the complexity of the input and output mapping, the amount of noise in the data, and the amount of training data available.

Sigmoid transfer functions are usually preferable to threshold activation functions because with sigmoid units, a small change in the weights produces a change in the output, which makes it possible to tell whether that change in the weights was good or bad. The following sigmoid transfer functions are often used for BPNN: Hyperbolic tangent sigmoid (tansig), log-sigmoid (logsig) and linear (purelin) transfer function. The tansig transfer function, which can produce both positive and negative values, tended to yield faster training that the logsig transfer function, which can produce only positive values. Table II summarizes the BPNN performance using different transfer functions, 6 neurons were used in the hidden layer for all transfer functions.

All of the transfer function combination tested, the logsig transfer function at hidden layer and tansig transfer functions at output layer with the highest $R^2$ (Table 2) were used in this work.

Once the BPNN was constructed, the weights were initialized and the network was ready for training. Ten different BP training algorithms were compared to select the best suited algorithm. For all BP training algorithms, a three-layer NN with a lo-sigmoid transfer function at hidden layer and a hyperbolic tangent sigmoid transfer function at output layer were used. For all BP training algorithms, 6 neurons were used in the hidden layer. Their characteristics deduced from the experiments are shown in Table III.

The benchmark comparison showed that the traingd (gradient descent BP) algorithm had the lowest training speed compared to all other algorithms, follow by trainda (gradient descent BP with adaptive learning rate) with $R^2 = 0.7383$, traindx (gradient descent BP with momentum and adaptive learning rate), trainoss (one step secant BP), traincfg (conjugate gradient BP with Fletcher-Reeves), trainbfg (BFGS quasi-Newton), trainscg (scaled conjugate gradient BP), traincgp (conjugate gradient BP with Polak-Ribiere), trainrp (resilient BP). The trainlm (Levenberg-Marquardt) with the highest $R^2$ was suitable as BP training algorithm.

The BPNN algorithm trainlm was developed using a three layer approach with a logsig transfer function at hidden layer and tansig transfer functions at output layer. This BPNN have six input neurons (moisture, ash, SiO$_2$ and pyrite) and one output neuron (AI) as shown in Fig. 1.

The error rates of proposed network were 2.12, 3.72 and 1.85 for training, testing and validation data, respectively. The coefficient of determination of proposed network was 0.84, 0.59 and 0.94 for training, testing and validation data respectively. All input and output data in the training phase were scaled so that they changed in the range of 0 and 1. The network was trained to predict the AI using a total of 40 training dataset, 26 testing datasets and 25 validation dataset. Identifying of the optimal NN structure was studies

### Table I

<table>
<thead>
<tr>
<th>Number of Neurons</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0.134843</td>
</tr>
<tr>
<td>3</td>
<td>0.093658</td>
</tr>
<tr>
<td>4</td>
<td>0.054025</td>
</tr>
<tr>
<td>5</td>
<td>0.039396</td>
</tr>
<tr>
<td>6</td>
<td>0.000257</td>
</tr>
<tr>
<td>7</td>
<td>0.012368</td>
</tr>
<tr>
<td>8</td>
<td>0.013655</td>
</tr>
</tbody>
</table>

### Table II

<table>
<thead>
<tr>
<th>Transfer function Versus $R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hidden layer</strong></td>
</tr>
<tr>
<td>logsig</td>
</tr>
<tr>
<td>logsig</td>
</tr>
<tr>
<td>logsig</td>
</tr>
<tr>
<td>tansig</td>
</tr>
<tr>
<td>tansig</td>
</tr>
</tbody>
</table>

### Table III

<table>
<thead>
<tr>
<th>BP Training algorithm</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>traingd</td>
<td>0.6393</td>
</tr>
<tr>
<td>traindx</td>
<td>0.7477</td>
</tr>
<tr>
<td>trainda</td>
<td>0.7383</td>
</tr>
<tr>
<td>trainrp</td>
<td>0.8317</td>
</tr>
<tr>
<td>trainlm</td>
<td>0.9167</td>
</tr>
<tr>
<td>trainbfg</td>
<td>0.8031</td>
</tr>
<tr>
<td>traincg</td>
<td>0.8139</td>
</tr>
<tr>
<td>trainoss</td>
<td>0.7765</td>
</tr>
<tr>
<td>traincfg</td>
<td>0.7807</td>
</tr>
<tr>
<td>traincgp</td>
<td>0.8256</td>
</tr>
</tbody>
</table>

Fig. 1. BPNN structure uses in this study
by changing the number of layers, nodes, transfer function, and iteration number. The prediction of BPNN versus actual measure for all (training, testing and validation) is shown in Fig. 2.

**B. GRNN Model Development**

The smoothness factor $\sigma$ is the only parameter which affects the GRNN performance by taking several aspects into account depending on the application the predicted output is used for [12]. For a prediction that is close to one of the training sample and a sufficiently small smoothness factor the influence of the neighboring training samples is minor. The contribution to the prediction is a lot smaller than the contribution of the training samples that are further away from the point of prediction can be neglected [12]. Due to the normalization the prediction therefore yields the value of the training sample in the vicinity of the each training sample. For a bigger $\sigma$ the influence of the neighboring training samples cannot be neglected. The prediction then is influenced by more point and the prediction is getting smoother. With even larger $\sigma$ the predicted curve will get flatter more smooth as well. In some cases this is desirable. For example when the available data include a lot of noise, then the prediction has to interpolate the data whereas if the data are correct [12]. GRNN has to fit the data more precisely and has to follow each little trend the data makes. If the $\sigma$ approaches infinity the predicted value is simply the average of all the sample points. Due to the fact that data are not generally without measurement errors and that the circumstances change from application to application, there cannot be a right or wrong way to choose $\sigma$. Table IV summarizes the results for GRNN model using different $\sigma$ values. The $\sigma$ value 0.05 can fit data very closely, with higher $R^2$ (0.951) value than when using the larger $\sigma$, but the larger $\sigma$ can make the function approximation smoother.

**Fig. 2.** AI predicted by BPNN in testing process versus actual measured

![Graph](image)

**Table IV**

<table>
<thead>
<tr>
<th>$\sigma$</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td>0.951</td>
</tr>
<tr>
<td>0.10</td>
<td>0.908</td>
</tr>
<tr>
<td>0.25</td>
<td>0.725</td>
</tr>
<tr>
<td>0.50</td>
<td>0.660</td>
</tr>
<tr>
<td>1.00</td>
<td>0.523</td>
</tr>
</tbody>
</table>

The proposed GRNN has three layers. To avoid overtraining, the spread constant was changed to allow testing data error became close the training data error. If a smaller smoothness constant is selected, the output of network would completely fit on training data, but the generalization ability of network might be decreased. The larger the smoothness factor, the smoother is the function approximation. Smoothness factor is used to fit data very closely than the typical distance between input vectors. Larger smoothness factor is used to fit data more smoothly. Fig. 3 represents the prediction of data using proposed GRNN versus actual data. The coefficient of determination ($R^2$) was 0.937 for all (training, testing and validation).

**C. Statistical performance of predicted models**

The statistical performance of the developed predicted models using Statistics toolbox 6.1 in Matlab 7.4 (R2007a) are given in Table V. The results showed that all the methods were quite stable. The values of each performance indicator ($R^2$, MAE and RMSE) were within 2% change in every case.

**Table V**

<table>
<thead>
<tr>
<th>Number of data points</th>
<th>GRNN $R^2$</th>
<th>MAE</th>
<th>RMSE</th>
<th>BPNN $R^2$</th>
<th>MAE</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>91</td>
<td>0.995</td>
<td>1.92</td>
<td>3.10</td>
<td>0.900</td>
<td>2.60</td>
<td>3.58</td>
</tr>
</tbody>
</table>

**D. Parametric Studies**

Abrasiveness refers to the capacity of coal to wear away or erode the surface with which it comes into contact. The parameters affecting wear rate abrasion rate are not well understood. It has been suggested that a high content of mineral matter and the presence of hard minerals like quartz and pyrite contribute to the abrasive qualities of coal [16]. The proposed NN was performed by a set of input parameters to study the influence of different parameters on the amount of AI. This set of parameters was as follows: $H_2O$, ash, $SiO_2$ and pyrite. The network prediction was obtained by varying a single parameter each time while keeping all other parameters constant.

Quartz and pyrite are the main components in the coal
responsible for the wear and abrasion [17]. Quartz is twice more abrasive than pyrite on a weight percent basis in coal [17]. This factor was attributed to quartz which is generally found as large “excluded” particles, whereas the pyrite is often “included” in the soft clays and the coal matrix [5].

Fig. 4 illustrates the effects of Quartz on AI. Quartz is the hardest common mineral associated with coal [9]. The relationship between abrasive index (AI) and quartz in the coal is represented in Fig. 6 with $R^2 = 0.725$.

The relationship between abrasive index (AI) and pyrite in the coal is represented in Fig. 5 with $R^2 = 0.776$.

The correlation between abrasive index and ash percentage has been reported in literature [9], [18], [19]. A linear relationship was found between the two variables, with $R^2 = 0.805$ as presented in Fig. 6. The ash yield essentially reflects the non-combustible residues of the different minerals associated with the coal.

High moisture content constituents of coals influenced AI of coal as illustrate in Fig. 7.

![Fig. 4. Effects of Quartz on abrasion index](image)

![Fig. 5. Effects of pyrite on abrasion index](image)

![Fig. 6. Effects of ash on abrasion index](image)

**IV. CONCLUSION**

BPNN and GRNN techniques were employed to explore the nonlinear relationships between AI and four variables (Ash, Quartz, pyrite and moisture) on the abrasiveness characteristics of thermal coal. It was found that the obtained results of BPNN and GRNN predictions were in good agreement with the actual measurements, with coefficient of determination, $R^2 = 0.9003$ for BPNN and $R^2 = 0.937$ for GRNN. The good results indicated the NN techniques were capable of accurately modeling AI from thermal coal.

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**REFERENCES**


