A DEEP LEARNING-BASED PROCEDURE FOR ESTIMATION OF ULTIMATE LOAD CARRYING CAPACITY OF STEEL TRUSSES USING ADVANCED ANALYSIS

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Abstract
In the present study, Deep Learning (DL) algorithm or Deep Neural Networks (DNN), one of the most powerful techniques in Machine Learning (ML), is employed for estimation of ultimate load factor of nonlinear inelastic steel truss. Datasets consisting of training and test data are created based on advanced analysis. In datasets, input data are the member cross-sections of the truss members and output data is the ultimate load factor of the whole structure. An example of a planar 39-bar steel truss is studied to demonstrate the efficiency and accuracy of the DL method. Five optimizers such as Adadelta, Adam, Nadam, RMSprop and SGD and five activation functions such as ELU, LeakyReLU, Sigmoid, Softplus, and Tanh are considered. Based on analysis results, it is proven that DL algorithm shows very high accuracy in the regression of the ultimate load factor of the planar 39-bar nonlinear inelastic steel truss. The number of layers can be selected with a small value such as 1, 2 or 3 layers and the number of neurons in each layer can be chosen in the range \([N_i, 3N_i]\) with \(N_i\) is the number of input variables of the model. The activation functions ELU and LeakyReLU have better convergence speed of the training process compared to Sigmoid, Softplus and Tanh. The optimizer Adam works well with all activation functions considered and produces better MSE values regarding both training and test data.

Keywords: deep learning; artificial neural networks; nonlinear inelastic analysis; steel truss; machine learning.

1. Introduction
Classical methods for design of steel structures are based on two main steps, where an elastic analysis is used first to calculate the forces in each structural member and then the safety of each member is checked using strength equations, that are inelastic analyses to account for nonlinear effects, by assuming each member as an isolated member. Obviously, these methods do not consider directly structural nonlinear behaviors and their member separate check cannot make sure the compatibility between the members and whole structure. Therefore, although these methods yield acceptable solutions for design of structure and save lots of computational efforts, they have been gradually being replaced by advanced analysis methods \([1–4]\) which can account for geometric and material nonlinearities directly and model complex contact conditions. Advanced analysis methods can also predict

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the load-carrying capacity of whole structure that allows elimination of the tedious individual member check approach used in the classical methods. However, advanced analysis methods are excessive computing times to solve the design problems which require lots of structural analyses such as optimization or reliability analysis of the structure [5–8]. In such cases, using metamodels based on machine learning (ML) techniques are considered as an efficient solution.

Metamodel is an approximate mathematical representation used to perform the complicated relationship between input and output data. In light of this, nonlinear inelastic responses of the structure are predicted without performing advanced analysis. Some popular ML methods are Support Vector Machine (SVM) [9], Kriging [10], Random Forest (RF) [11], Gradient Tree Boosting (GTB) [12], Decision Tree (DT) [13], and so on. The applications of ML methods into structural design are quite diverse but focused primarily on damage detection [14, 15] and health monitoring [16, 17]. Besides, researchers have been applying ML methods for structural optimization [18], reliability analysis [19], prediction of structural ultimate strength [20], etc.

The performance of traditional ML methods largely depends on the data representation choice of the users since these methods cannot automatically detect the representations or features needed for classification or detection from the raw input data. The pattern-recognition often requires complex techniques with high expertise. Therefore, using ML methods is complicated. On the contrary, modern ML methods are called representation learning methods because the data presentations can be automatically discovered. This not only improves the efficiency and accuracy of ML methods but also makes the use of these methods simpler. A review of the representation learning methods is provided by Bengio et al. [21].

Deep learning (DL) in Artificial Neural Network (ANN), one of the best branch of the ML methods, has been commonly used in various structural design and analysis problems such as damage detection [22], health monitoring [23], etc. Several studies also listed by LeCun et al. [24] to demonstrate the efficiency of DL with other ML methods such as image and speed recognitions, natural language understanding, regression, classification, etc. Recently, by solving a well-known ten-bar truss problem, Lee et al. [25] showed the efficiency and accuracy of the DL comparing to the conventional neural networks in structural analysis. It is noted that most DL models are based on an ANN that consists of multiple levels of representation by utilizing simple but nonlinear interconnected layers with many neurons per layer. In DL models, the presentation at the following layers have higher abstraction levels than the previous one. Important information is amplified whilst non-critical information is gradually decreased and excluded through the layers. With such a complex and flexible organization system, as a result, DL can handle complicated and high-dimensional data. Additionally, developing and using of a DL model do not need a high expertise of the users. For this reason, these methods can be effectively applied in many fields of technology, medical, business, and science.

This paper presents a DL-based procedure for estimating the ultimate load-carrying capacity of nonlinear inelastic steel truss. Firstly, advanced analysis is presented to capture the structure nonlinear inelastic behaviors. Then, data consisting of inputs and outputs are collected from advanced analyses. The inputs are the cross-section of members and the output is the ultimate load factor of the truss structure. In order to demonstrate the efficiency and accuracy of DL algorithm, an example of a planar 39-bar steel truss is taken into consideration. In addition, sensitivity analyses are performed to examine the influences of $N_h$, the $N_n$, activation functions, and optimizers on the accuracy of DL method for the regression of the ultimate load factor of this structure.
2. Advanced analysis for calculating ultimate load factor of steel trusses

The stress-strain curve proposed by Blandford [26] presented in Fig. 1 is employed to perform the constitutive model since it includes most important behaviors of material such as: elastic, elastic and inelastic post-buckling, unloading and reloading. Compressive stress is positive in this figure. The equations of the parts of the stress-strain curve in Fig. 1 as follows:

- Part (a):

\[ \sigma = E\epsilon, \quad \epsilon < \epsilon_k \]  

- Part (b):

\[ \sigma = \sigma_k, \quad \epsilon_k \leq \epsilon < \epsilon_{ib} \]  

- Part (c):

\[ \sigma = \sigma_1 + (\sigma_k - \sigma_1) e^{-(X_1 + X_2 \sqrt{\epsilon - \epsilon_{ib}})/(\epsilon - \epsilon_{ib})}, \quad \epsilon \geq \epsilon_{ib} \]  

- Part (d):

\[ \sigma = \sigma_1 + \frac{\epsilon - \epsilon_1}{0.5E} + \left( \frac{1}{\sigma_2 - \sigma_1} - \frac{1}{0.5E(\epsilon_2 - \epsilon_1)} \right) (\epsilon - \epsilon_1) \]  

- Part (e):

\[ \sigma = E\epsilon \text{ for } |\epsilon| < |\epsilon_y| \]  

- Part (f):

\[ |\sigma| = \sigma_y \text{ for } |\epsilon| \geq |\epsilon_y| \]  

- Part (g):

\[ |\sigma| = E \left( |\epsilon| - (|\epsilon_3| - |\epsilon_3|) \right) \]
The notations in the above equations are defined as follows: $\sigma$ is axial stress; $\varepsilon$ is axial strain; $E$ is elastic modulus; $\varepsilon_{ib}$ is axial strain at the beginning point of inelastic post-buckling; $\sigma_y$ is yield stress; $\varepsilon_y$ is yield strain; $\sigma_{cr}$ is Euler critical buckling stress, $\sigma_{cr} = \frac{\pi^2 EI}{AL^2}$; $\varepsilon_{cr}$ is Euler critical buckling strain, $\varepsilon_{cr} = \frac{\sigma_{cr}}{E}$; $A$ is cross-section; $I$ is inertia moment of weak axis; $L$ is length of the member; $\sigma_k = \sigma_{cr}$ if yielding occurs and $\sigma_k = \sigma_y$ if the elastic buckling occurs; $\varepsilon_k = \varepsilon_y$ if yielding occurs and $\varepsilon_k$ if the elastic buckling occurs; $\sigma_l$ is asymptotic lower stress limit; $X_1$ and $X_2$ are parameters based on $(L/r)$ of compressive member.

The incremental form of equilibrium equation for a truss element is expressed as [27]

$$\begin{bmatrix} [k_E] + [k_G] + [s_1] + [s_2] + [s_3] \end{bmatrix} \{d\} + \{1 \} f = \{2 \} f$$

in which $\{1 \} f$ and $\{2 \} f$ are the initial nodal element forces at previous and current configurations, respectively; $[k_E]$ and $[k_G]$ are the elastic and geometric stiffness matrices, respectively; and, $[s_1]$, $[s_2]$ and $[s_3]$ are the higher-order stiffness matrices. The detail of these matrices can be found in [4].

3. Deep learning in artificial neural network

As mentioned above, DL algorithms or deep neural networks (DNN) are effectively used in handling both regression and classification problems. Due to the training data features, DNN model can be classified as (1) supervised learning, (2) unsupervised learning, and (3) semi-supervised learning. For the estimation problems of steel trusses considered in this study, the supervised learning is employed since the input and output data are determined. In the supervised learning algorithm, feedforward neural networks (FNN) using backpropagation (BP) algorithm are commonly used. The FNN basic issues using BP algorithm are indicated in the following sections.

3.1. Supervised learning

In supervised learning form, the training data is given in an expression as follows:

$$T = \{(X_i, Y_i)\}_{i=1}^N$$

where $X_i$ is the input vector of the data $i^{th}$ or the feature vector of the data $i^{th}$; $Y_i$ is the output vector or the label of the data $i^{th}$; $N$ is the number of data samples. In the supervised learning algorithm, the correct answers are given; therefore, the training process results can be controlled by minimizing the error between the predicted and exact values. As an example, with the input $X_i$, the predicted result $Y'_i$ is given by the training model. To enhance the training model, the optimization process is used to minimize the mean-square error function which is given as follows:

$$E_{MSE} = \frac{1}{N} \sum_{i=1}^{N} (Y_i - Y'_i)^2$$

3.2. Feedforward neural network

In the FNN, there are many layers and each layer has many units. Each layer includes connections to the next layer. The first layer plays the role of the input layer where each unit in this layer is a data feature. The last layer plays the role of the output layer, in which each unit is a data label. Remaining layers are the hidden layers and they play the role of transferring the important features from the input
layer to the output layer. An example of a fully connected FNN with 2 hidden layers is described in Fig. 2.

The input of an unit is computed based on the weight function, which can be mathematically stated as follows:

$$u_i^{(j)} = \sum_{l=1}^{N_u^{(j-1)}} w_{li}^{(j)}$$  \hspace{1cm} (11)

where $u_i^{(j)}$ is the input of the $i^{th}$ unit of the $j^{th}$ layer; $w_{li}^{(j)}$ is the weight at the connection between the unit $l^{th}$ of the $(j - 1)^{th}$ layer and the $i^{th}$ unit of the $j^{th}$ layer; $N_u^{(j-1)}$ is the number of units of the $(j - 1)^{th}$ layer. In each layer, to transform the input signal to the output signal in the unit, an activation function $f()$ is employed as below:

$$v_i^{(j)} = f(u_i^{(j)})$$  \hspace{1cm} (12)

where $v_i^{(j)}$ is the output of the $i^{th}$ unit in the $j^{th}$ layer. It is noted that the users should choose the suitable activation functions to obtain the best result for their problems.

In the supervised learning algorithm, to evaluate the accuracy of the training model, loss functions are computed based on the predicted output and the exact output of the model, are employed. Several common loss functions are mean_squared_error, mean_absolute_error, mean_absolute_percentage_error, etc. In the present work, we use mean_squared_error loss function which is expressed in Eq. (10). The objective of a training process of an DNN model is to minimize the loss function.

### 3.3. Backpropagation algorithm

In the BP algorithm, the objective function is loss function, while design variables are the weight parameters. The purpose of an DNN model training process is to find the optimal weight parameters in which the loss function is minimum. In order to optimize the weight parameters, the BP algorithm is commonly utilized based on the gradient descent method. In multilayer FNN, an BP algorithm is employed to compute the effect of each weight corresponding to loss function. The weight parameters can be revised in the BP algorithm based on the gradient descent method given as follows:

$$W^{(j+1)} = W^{(j)} - \varepsilon \partial L^{(j)} + \mu \Delta W^{(j-1)}$$  \hspace{1cm} (13)
4. Deep learning model for estimating ultimate load factor of steel trusses

The main steps of the DNN model for estimating the ultimate load factor of steel trusses as follows:

- **Step 1**: Definition of problem and generation of data
  
  Cross-sectional areas of structural members are considered as the inputs, while the ultimate load factor of the structure is the output. Generation of data starts by creating \(m\) input samples, \((X_1, X_2, \ldots, X_m)\), where \(X = (x_1, x_2, \ldots, x_n)\) is the vector of structural cross-sectional areas. \(x_i\) is the cross-section area of the structural member group \(i^{th}\) which is generated randomly in the predefined range \([x_i^{lowb}, x_i^{upb}]\). Corresponding to each input vector \(X\), the ultimate load factor of the structure, \(lf_i\), is calculated by using advanced analysis.

- **Step 2**: Data scale
  
  To improve the performance of the DNN training process, all the data is scaled down in the range \((0,1]\) by using the “MinMaxScaler” method as follows:

  \[
  x_i^{scale} = \frac{x_i - x_i^{lowb}}{x_i^{upb} - x_i^{lowb}} \quad (14a)
  \]
  
  \[
  l f_i^{scale} = \frac{lf_i - \min(lf_i)}{\max(lf_i) - \min(lf_i)} \quad (14b)
  \]

  All scaled data is now divided into two groups of training data set, \((X_{train}, Y_{train})\), and test data set, \((X_{test}, Y_{test})\). The training data and test data are used to develop and check the DNN model, respectively.

- **Step 3**: Definition of a DNN model structure
  
  The DNN model is developed using sequential model where the layers are defined from the input layer, through the hidden layers, to the output layer. In each layer, the number of units and the type of activation function are chosen. Notes that, the number of units in the input layer can be equal or different to the number of input variables, but the number in the output layer must be equal to the number of output variables. The activation function of each layer can be chosen as the same or different. The connections between the units in two adjacent layers are fully connected or using dropout to prevent overfitting in training the model. The number of hidden layers and units in each layer affect the computation time and accuracy of training model. Therefore, several values of number of hidden layers and units in each layer should be tried to find the acceptable DNN structure.

- **Step 4**: Compile and train the DNN model
  
  To compile the DNN model, the loss function and optimizer are defined first. Some popular regression loss functions are mean_squared_error, mean_absolute_error, mean_absolute_percentage_error, etc. And, some optimizers are stochastic gradient descent (SGD), Adagrad, RMSprop, Adam, etc. Different loss functions and optimizers need to be tried in order to find the acceptable ones for the studied problem.

  To train the DNN model, the ‘fit()’ function is used with the main parameters as: training data, test data, mini-batch, the number of epochs. The training and test data are presented in Step 2. Mini-batch method has been proposed as a way to speed-up the convergence of the model training by dividing the
training data into several smaller batches and then training the model through these batches in turn. Epoch is a time that all training data is passed forward and backward through the neural network only once. The more epochs are used, the more the model is improved, up to a certain point where the model is converged and stops improving during each epoch.

After this step, the DNN model is obtained. It can be used for predicting the ultimate load factor of structure if a new input data is given.

5. Numerical examples

In this section, the planar 39-bar steel truss presented in Fig. 3 is studied. The cross-sectional areas of 39 structural members are divided into 39 groups that have the same design range [645.16, 11290.3] (mm²). Steel material has the yield strength of 172.375 (MPa) and the elastic modulus of 68.95 (GPa). The horizontal applied loads according to the X-axis at all nodes are equal to 136 (KN), and the gravity loads at all nodes equal 170 (KN). For developing the DNN model, the programming language Python and the open-source software libraries Tensorflow and Keras are employed, while PAAP program [4] is used for performing advanced analysis. Fig. 4 shows the histogram of ultimate load-factor of the truss with the number of samples of 50,000.

In the current section, parameters consisting of \( N_h, N_n \), activation functions, optimizers are taken into consideration to investigate their influence on the accuracy of the DL model for estimation of the truss structure. It is noted that the accuracy of the DL model is evaluated by using average MSE in all training samples over 30 independent runs. In addition, the number of training and test data is chosen to be 5,000 and 10,000 respectively, and the number of epochs is fixed at 10,000 for all cases. In addition, it is noteworthy that training and test data are generated from advanced analyses of PAAP software with a similar computational time of approximately 10s for each sample.
5.1. Effect of the number of hidden layers and neurons

In this section, the effect of \( N_h \) on the accuracy of the developed DL model is investigated by considering various \( N_h \), while the activation function is chosen to be LeakyReLU and the optimizer is selected to be Adam. \( N_h \) is varied from 1 to 5, while \( N_n \) is considered to be \( N_i/2 \) (20), \( 2N_i \), \( 3N_i \), \( 4N_i \) (\( N_i \) is the number of input variables, \( N_i = 39 \)). Tables 1 and 2 present the average MSE for training and test data after 10,000 epochs, respectively. It appears from Table 1 that the accuracy of the training model significantly increases when \( N_h \) increases from 1 to 5 with respect to the increase of \( N_n \) (i.e., with \( N_n \) increases from 20 to 156, the accuracy increases from 1.4 to 26.74 times, respectively). It is apparent that the DL model with \( N_h \) equal to 5 always has the best accuracy in regression of the ultimate load factor regardless of \( N_n \). This means that when the number of hidden layers and nodes increase, the DNN model more well recognize the features of the data. It is observed from Table 2 that the accuracy of the DL model for test data is similar in all cases. This implies that the model is not overfitting, furthermore the number of layers can be chosen small such as 1, 2 or 3 layers and the number of nodes is in the range \([N_i, 3N_i]\) in the light of the accuracy of the model for test data.

<table>
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<tr>
<th>Hidden nodes in each layer</th>
<th>( N_i/2 = 20 )</th>
<th>( N_i = 39 )</th>
<th>( 2N_i = 78 )</th>
<th>( 3N_i = 117 )</th>
<th>( 4N_i = 156 )</th>
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<td>Number of hidden layers</td>
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<td>Time (s)</td>
<td>MSE</td>
<td>Time (s)</td>
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5.2. Effect of the activation function

In order to investigate the effect of activation functions on the performance of a developed DL model, five activation functions consisting of ELU, LeakyReLU, Sigmoid, Softplus, and Tanh are taken into consideration, where ELU and LeakyReLU are advanced activation functions. It is noted that \( N_h \) is fixed at 3, while \( N_n \) is 117 in all analyses since this case gives the best accuracy reported in Section 5.1. Figs. 5 and 6 display the comparison among various activation functions (in combination with the Adam optimizer) based on convergence history of MSE of the ultimate load factor of the truss corresponding to training and test data. As can see in Fig. 5, for training data, the model using Tanh has the best convergence rate, while the model using Sigmoid has the least convergence rate compared with other models. At the final iteration, the average MSE of the model using Tanh is smallest \((1.55 \times 10^{-5})\), while the average MSEs of the models using ELU, LeakyReLU, and Softplus are close each other which is much lower than that of the model using Sigmoid. For test data, using
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Sigmoid has the smallest MSE among other activation functions as can be observed from Fig. 6. Using Tanh activation function provides the highest MSE value compared to using other ones. It is noteworthy from Figs. 5 and 6 that the use of Sigmoid gives the worst results for training data but it provides the best accuracy for test data.

5.3. Effect of the optimizer

In order to examine the influence of optimizers on the efficiency and accuracy of a developed DL model, five optimizers consisting of Adadelta, Adam, Nadam, RMSprop, and SGD are considered, while $N_h$ and $N_n$ are 3 and 117, respectively. Table 3 illustrates the comparison of various optimizers (in combination with the various activation functions) based on average MSEs of the ultimate load factor of the truss structure. It can be seen from this table that the combinations between activation functions and optimizers provide a high accuracy of the model for training and test data in most cases. It appears that AdaDelta, Adam and Nadam can work well with all activation functions considered. However, Nadam produces higher MSE for test data than AdaDelta and Adam, while AdaDelta has greater MSE for training data than Adam and Nadam in all cases. SGD does not work well with Sigmoid and SoftPlus since they yield a high MSE for both training and test data. Similarly, RMSprop does not work well with Tanh. Adam optimizer has the lowest value of average MSE for training data regardless of the activation function used. This was also found by Lee et al. [25] for the linear 10-bar...
truss problem. In addition, the combination of Adam optimizer and Tanh activation function gives the best accuracy for training data. The combination of SGD optimizer and Tanh activation function provides the best accurate results for test data. On the other hand, it is also found that the computational times for training the DNN model for various activation functions and optimizers are almost similar to each other (approximately 400 seconds).

6. Conclusions

In the current study, an efficient method is proposed for estimation of nonlinear inelastic steel truss using DL algorithm, one of the most powerful branch of ML methods. Datasets include training and test data, which are collected from advanced analyses of steel truss. In this data, inputs are cross-sections of members and the outputs are the ultimate load-factor of the truss structure. The number of training data is chosen to be 5,000, while the number of test data is 10,000 in datasets. A planar 39-bar steel truss is taken into account for demonstration of the performance of the DL algorithm. Based on the analysis results, it is demonstrated that DL has a very high accuracy in the regression of the ultimate load-factor of the nonlinear inelastic steel truss. Additionally, sensitivity analyses are carried out to investigate the influences of the number of hidden layers, the number of neurons in each layer, activation functions, and optimizers on the accuracy of DL algorithm for the regression of the ultimate load factor of this structure. It is concluded that most of combination of optimizers and activation functions give a high accuracy of the model. The number of layers can be selected with a small value such as 1, 2 or 3 layers and the number of nodes can be chosen in the range \([N_i, 3N_i]\) to reduce the running time but still have a high accuracy of the model. The activation functions ELU and LeakyReLU improve the convergence speed of the training process compared to Sigmoid, Softplus and Tanh. The optimizer Adam is recommended to use since it works well with all activation functions considered and produces the better MSE values with both training and test data.

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