**Significance of Artificial Neural Network Analytical Models in Materials Performance Prediction**

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**Abstract**

In materials science, performance prediction of materials plays an important role in improving the quality of materials as well as preventing serious damage to environment and threat to public safety. Traditional regression analysis models in materials science are still not perfect, unable to handle large amount of data, limited to capture nonlinearities of data and time-consuming for prediction. It makes a strong demand on data analysis of materials data with the help of computer science method. In recent years, artificial neural network (ANN) is increasingly performing as a strong bridge between materials science and computer science. In this paper, various ANN applications for predicting various properties of different materials in various materials related areas were reviewed. It highlights the significance of ANN in materials related problems and the trend of ANN in context of materials science with some limitations.

Keyword: materials performance prediction, artificial neural network, multilayer perceptron, backpropagation, feedforward, radial basis function

# Introduction

Since material service safety is important to be maintained, many researchers devote to experimental testing and data analyzing to discover the different factors’ influencing on materials performance. Traditional mathematical analysis models used in materials science are still not perfect, unable to handle large amount of data, limited to capture complex relationship and time-consuming for prediction. In recent years, an emerging area of materials performance evaluation and prediction by artificial neural network (ANN) was evolved as an efficient way. ANN analysis can produce satisfactory prediction results of materials properties, compared to traditional data analysis [[1](#_ENREF_1)] [[2](#_ENREF_2)] [[3](#_ENREF_3)] [[4](#_ENREF_4)].

One hot research target in materials science is to determine the unknown relations among materials properties. ANN can predict the output by exploring arbitrary interactions among the variables and enabling quantitative expression and understanding of very complicated problems [[5](#_ENREF_5)]. Because of its parallel structure and fast learning capacity, it can be used for linear and non-linear data analyses in applications such as pattern recognition, prediction, system identification and control. The main benefit of neural network is its generalization ability, noise tolerance and prediction ability of output of unseen test data with savings in cost and time [[6](#_ENREF_6)].

This paper aims to give a hand to materials and computer researchers with a comprehensive view of ANN models implemented in various prediction applications in material science researches. This review will highlight the significance of ANN in materials related areas and the facts that arouse continuous ANN implementation in future materials related problems.

The rest of the contents are arranged as follows. The basic introduction to ANN followed by the principal idea of ANN, the simplest ANN model, various activation functions used in ANNs and different classifications of ANN models is firstly described in Section 2. Section 3 gives the significance of ANN in materials related researches and analyzes their advantages and shortcomings. The future trend of ANN application in materials science is pointed out in Section 4.

# Artificial Neural Network

## The Principal Idea of Neural Network

Artificial neural network is one kind of network model established in mathematical way that is based upon the working principle of biological neurons connected to each other in biological neural network. In ANN, a neuron is an element that receives multiple inputs and generates a single output and is characterized by its weight, bias and activation function. Those neurons are firstly grouped into layers and then those layers are connected to each other, thus forming a network model. The information from the input layer is moved as the input to the hidden layer that will do processing to produce the final output and sent the output to output layer. That kind of movement from the input layer to the output layer is known as feed forward propagation. When there is error between the actual output and the expected output at the output layer, the weights and biases of neurons will be updated to reduce the error in backpropagation process. Learning process, called as training, can learn from previous data and iteratively adjust the weight values to reduce error between actual output and desired output. When the desired output is achieved at a satisfactory level, training is stopped and the weighted links between processing units are saved. The final weights and biases after training are used as analytical tools for prediction of unseen input data. Details of the neural network method itself can be found in [[7](#_ENREF_7)] [[8](#_ENREF_8)] [[9](#_ENREF_9)] [[10](#_ENREF_10)].

## The Simplest ANN

Single layer neural network, also known as perceptron, is the first and simplest neural network model [[7](#_ENREF_7)]. It consists of neurons in a single layer. Fig 1 demonstrates a single layer neural network, in which the number of input (m), the number of neuron (n) and input signals (xi) are connected to all neurons in only one layer.

As all neurons in a single layer perceptron network produce their corresponding output, the number of output is the same as the number of neuron (n). The input signals (x1, x2, …, xm) are connected to the neurons of the single layer via their corresponding weights (w11, w12, …, wnm) so that every neuron has the corresponding weight value for each input signal respectively. It can be seen in Fig 1that the first neuron, neuron-1 has weight data (w11, w12, …, w1m) for input signal (x1, x2, …, xm) as well as neuron-2 also has its corresponding weight data (w21, w22, …, w2m) for the same input signal (x1, x2, …, xm). It goes in the same way for the nth neuron that its weight data will be (wn1, wn2, …, wnm). The input signal X can be expressed in vector form as

**(1)**

and weight data as

**(2)**

where n is the number of neuron and m is the number of input. Any connection between neurons and input of the network can be calculated based on the number of n and m. For example, Fig 1 shows the connection path between the first input (x1) and the second neuron (n2) can be easily seen as w21.



**Fig 1**. The structure of perceptron (single layer) neural network



**Fig 2**. The working principle of a neuron

The working principle of an artificial neuron is shown in Fig 2. As it is mainly constituted with two components: sum and activation function, it can be said that an artificial neuron works in two steps. In the first step, all input data in a neuron are multiplied with their respective weight data and those multiplications are summed to pass through the activation function. The second step accepts the sum as the input to the activation function that will produce the output. The netinput to the activation function in first step can be calculated in vector form as follows:

***netinput= WX =******w1\*x1 + w2\*x2 + w3\*x3+……+ wn\*xm* (3)**

The weighted data (weight data \* input data) are summed and passed as the input to the activation function of the neuron in the second step. The activation function accepts the input and process it to produce the output. The output of the neuron can be calculated as

***Output*n= (4)**

The output of the entire network can also be calculated by passing the summation of multiplication of the input vector and weight vector to the activation function as follows:

***y= (Netout = F(netinput) =F (WX))* (5)**

To improve the performance of neural network, some neurons includes weight data as well as bias data. By adding bias values b, the performance of the neurons is improved by shifting decision boundary of classifiers to left or right. The value of the bias is generally set to 1 and the mathematical model of the neuron with bias value can be expressed as

***y=Net out = ƒ(netinput) = ƒ(WX+b)* (6)**

## Activation Function

Since every neuron uses an activation function to generate output from the weighted data and the accuracy of the output is dependent on activation function, it is important to choose the correct activation function for neurons. The widely used activation functions in neural network architecture are shown in Table 1.

**Table 1**. Activation functions mostly used in ANN

|  |  |  |
| --- | --- | --- |
| **Name** | **Formula** | **Range** |
| **Linear** |  |  |
| **Semi-linear** |  |  |
| **Logistic (sigmoidal)** |  |  |
| **Hyperbolic tangent (sigmoidal)** |  |  |
| **Exponential** |  |  |
| **Sinusoidal (sine)** |  |  |
| **Rational (sigmoidal)** |  |  |
| **Step** |  |  |
| **Hard limit (threshold)** |  |  |
| **Modular** |  |  |
| **Signed (signature)** |  |  |
| **Quadratic** |  |  |
| **ReLu** |  |  |
| **Leaky ReLu** |  |  |
| **Gaussian** |  |  |
| **Softmax** | for i=1,…, J |  |

Among them, the three most widely used activation functions are linear, sigmoid and hard-limit function. Linear function operates no changes in incoming input signals to generate the original output values. Sigmoid non-linear function, also known as logistic function, can be defined as . It can generate the output between 0 and 1for input value range of -∞ and ∞. Hard-limit function gives the output of neuron as 0 if the input is less than 0; otherwise, it gives the output as 1.

## Classification of Neural Network

Different network models can be basically classified due to the architecture of neural networks: network topology, layer number, activation function and feedback connection.

### From network topology

The classification of neural network defined by network topology includes fully connected and unfully connected. Fully connected network means that every neuron of each layer is connected to all the neurons of another neighbor layer. In such kind of network, neurons of one layer transmit their output signal as input to the neurons of next layer. One of well-known fully connected neural network is hopfield network[[11](#_ENREF_11)]. In unfully connected neural network, neurons of one layer do not connect to all neurons of another layer, just connecting to some of corresponding neurons from another layer.

### From layer number

The simplest neural network has only one layer. The input and output are connected directly with the neurons on the single layer. Neural networks composed of at least two layers are known as multi-layer neural networks. In multi-layer neural networks, number of neurons of each layer can be same or different. Layers are connected to each other from left to right order. Network architecture can be described as (number of input layer neurons-number of hidden layer neurons-number of output layer neurons); for example, (5-8-1) structure means 5 neurons in input layer, 8 hidden neurons in hidden layer (hereby (5-8-8-1) means two hidden layers with 8 hidden neurons respectively) and 1 neuron in output layer.

### From activation function type

Activation function is important for the performance of ANN. From Table 1, we can see that there are many different kinds of activation function. ANN can be classified by the types of neurons of the network that use activation function: homogeneous network in which activation functions of the neurons in the network are same and heterogeneous network in which the neurons in the network use different activation functions. Weingaertner et al. presented a framework to develop the automatic design of heterogeneous neural networks by using different activation functions in hidden layer neurons of feed forward neural networks [[12](#_ENREF_12)]. Lee et al. compared their proposed heterogeneous neural network with homogeneous neural network composed of the conventional neuron model, achieving improvement in error rate and training speed, and also extends to memory capacity [[13](#_ENREF_13)].



**Fig 3**. Types of Artificial Neural Network Classified by Feeding Direction

### From feedback direction

ANN can also be classified if it is feed forward (static) or feedback (dynamic) network according to its feeding direction between layers. ANN layers are usually connected in one direction from left to right order, known as feed forward. Layers of neural networks known as static networks are connected in one direction from one layer to another in feed forward way without feedback connection.

But some networks are both feed forward and feedback connected. Feedback networks introduce signal loops in network in both directions, which are powerful and have more complicated structure. One of popular network models with feedback connection are known as recurrent networks (dynamic networks). Fig 3 illustrates a classification of current popular neural network models based on those two classes of feed forward neural network and feedback neural network [[14](#_ENREF_14)].

Besides neural network classifications mentioned above, ANN can also combine different types of neural networks together for special purpose. It combines two or more different neural networks above, known as hybrid neural network. Such kind of neural networks are multilayer neural networks composed of multiple layers with different separate topologies and a special learning algorithm is needed for hybrid neural networks.

# Significance of Neural Network Applications in Materials Researches

The generic nature of ANN was widely applied in materials related fields, covering mechanical [[15](#_ENREF_15)], wear [[16](#_ENREF_16)], hydrological [[17](#_ENREF_17)], atmospheric science [[18](#_ENREF_18)] [[19](#_ENREF_19)], civil engineering [[20](#_ENREF_20)]and etc., for several application areas in which materials failures can occur, including pressurized water reactors [[21](#_ENREF_21)], quenching operation [[22](#_ENREF_22)] , wrought application [[23](#_ENREF_23)], distribution and transmission electric energy lines [[24](#_ENREF_24)], aqueous chloride solution [[25](#_ENREF_25)] , nuclear waste disposal [[26](#_ENREF_26)], aerospace and aviation sectors [[27](#_ENREF_27)], steel production lines[[28](#_ENREF_28)] and etc. ANN was established for predicting corrosion in different environments such as atmosphere[[29](#_ENREF_29)], water[[30](#_ENREF_30)], sea[[24](#_ENREF_24)], high temperature[[31](#_ENREF_31)] and so on as well as forecasting materials performance as failure forms of fatigue [[32](#_ENREF_32)], crack [[21](#_ENREF_21)], pitting [[4](#_ENREF_4)], creep [[33](#_ENREF_33)], flow stress[[34](#_ENREF_34)], wear [[16](#_ENREF_16)], elasticity[[35](#_ENREF_35)], toughness and hardness [[36](#_ENREF_36)], hot deformation behavior [[37](#_ENREF_37)] [[3](#_ENREF_3)], yield strength [[38](#_ENREF_38)], ultimate strength [[39](#_ENREF_39)], elongation to fracture [[23](#_ENREF_23)], etc.

Materials on researches were varied due to the intention materials researchers focused on. Several of them were steel[[40](#_ENREF_40)] [[25](#_ENREF_25)], zinc[[41](#_ENREF_41)], aluminum[[42](#_ENREF_42)], copper[[24](#_ENREF_24)], iron and titanium [[43](#_ENREF_43)], magnesium[[44](#_ENREF_44)], other metal alloys [[21](#_ENREF_21)] [[22](#_ENREF_22)] [[26](#_ENREF_26)] [[27](#_ENREF_27)] [[34](#_ENREF_34)] [[45](#_ENREF_45)] [[46](#_ENREF_46)]and so on. The majority of materials related applications are focused on implementing feed forward backpropagation neural networks. Recent works on other powerful, complicated and efficient neural networks models, such as radial basic function neural network (RBFNN), convolutional neural network (CNN) and generalized regression neural network (GRNN) can also give satisfactory prediction. In materials science researches, it is important to know the role of variables affecting on material properties and determine which input variable is the most influencing factor on desired output variable. Some input ranking methods, such as change of MSE (COM), fuzzy curves and sensitivity analysis, have been applied to extract knowledge from trained ANN [[47](#_ENREF_47)]. Previous recent works have shown that integration of neural networks with other computing paradigms such as Bayesian framework [[48](#_ENREF_48)], genetic algorithm [[49](#_ENREF_49)] [[50](#_ENREF_50)] [[51](#_ENREF_51)], sensitivity analysis [[52](#_ENREF_52)] [[41](#_ENREF_41)] [[25](#_ENREF_25)] and fuzzy logic[[44](#_ENREF_44)] can effectively be used to make the performance of neural network models more efficient. The development of more than one ANN types, such as linear model (LM) (simplest form of ANN), multilayer perceptron (MLP), radial basis function (RBF) and so on, and their comparison results were also discussed in some researches in literature [[53](#_ENREF_53)] [[54](#_ENREF_54)].

Bhadeshia [[55](#_ENREF_55)] reviewed on some weakness of traditional linear regression analysis and why neural network architecture came up to replace linear regression analysis with specific neural network applications in context of materials. We summarize the majority of neural network prediction models exploited in materials related problems for predicting different properties of different materials in general, shown in Table 2. Most of those ANN prediction results were agreed with experimental results.

**Table 2**. Neural network predictive models used in materials related problems

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Materials | Applied Area/ Environment | Input | Activation Function | Output | Predicted/ relevant Property | Neural Network Model |
| steel and zinc[[41](#_ENREF_41)] | industrial | temperature, time of wetness TOW, exposure time, Sulphur dioxide concentration and chloride concentration | sigmoid | corrosion depth | chemical | multilayer perceptron |
| 304 Stainless Steel[[25](#_ENREF_25)] | aqueous chloride solution | temperature, Chloride ions concentrations and applied stress | sigmoid | time of failure as a result of stress corrosion cracking | mechanical | multilayer perceptron |
| Al, low carbon steel, copper[[24](#_ENREF_24)] | distribution and transmission electric energy lines | environmental parameters | linear and sigmoidal tangent, logarithmic | laboratory evaluated atmospheric corrosion rates | chemical | multilayer perceptron |
| Iron alloy, carbon and alloy steel, grade-2 titanium[[43](#_ENREF_43)] | industrial | Mo content x, Ni content (14 x), (x + 1), HCl, H2SO4, potential (E), dE,  compositions (B, C, Cr, Cu, Fe, Mn, Mo, Ni, P, S, Si, Ti, and V), duration of exposure, log (duration of exposure),  incubation period and the duration of experiments | hyperbolic tangent | log(i), corrosion potential Ecorr, corrosion current icorr, and polarization resistance Rp (general corrosion)  corrosion rate and log (corrosion rate) (general corrosion)  maximum pit depth Dmax, sample weight change W, charge amount Q, ratio Q/W, O2 consumed, log (Dmax), log (W), log (Q), log (Q/W), and log (O2consumed) (localized corrosion) | chemical | multilayer perceptron |
| nickel-based alloy 22[[26](#_ENREF_26)] | industrial (nuclear waste disposal) | temperature values, sample treatment types: welded or annealed, and the exposure time,  sample treatments: welded or annealed, temperature, pH, Cl-, aging time, types of corrosion  frequency (w) and log(w) | hyperbolic tangent | corrosion weight loss in mg (general corrosion),  repassivation potential (localized corrosion),  Zr and Zi i.e. real and imaginary impedances, respectively, abs(Z), and Phase (localized corrosion) | chemical | multilayer perceptron |
| A357 alloy[[22](#_ENREF_22)] | quenching operation | deformation temperature, strain rate and strain | tan sigmoid and log sigmoid, linear | thermal flow stress | thermal | backpropagation Neural Network |
| Alloy 600[[21](#_ENREF_21)] | pressurized water reactors (PWR) | temperature, KI, pH, Conductivity, ECP, Yield strength, B(OH)3, LiOH | hyperbolic tan (tanh), linear | crack growth rate (CGR) | mechanical | backpropagation Neural Network |
| Sn-Ag alloy[[45](#_ENREF_45)] | electronics industry | the contents of In. Bi, Sb, RE, Sn, Ag, Cu, | sigmoid, purelin | tensile strength, shear strength and solidification temperature | mechanical | backpropagation Neural Network |
| Ti-5Al-2Sn-2Zr-4Cr-4Mo alloy[[46](#_ENREF_46)] | elevated temperature | deformation temperature, equivalent strain rate, and equivalent strain, | sigmoid, linear | flow stress | mechanical | backpropagation Neural Network |
| steel[[40](#_ENREF_40)] | concrete structures and buildings | time intervals | sigmoid | electrical potential as a factor for steel corrosion | chemical | backpropagation Neural Network |
| dual-phase (DP) steels[[32](#_ENREF_32)] | corrosive environment | stress intensity factor (DK), and volume % of martensite content (%M) | gaussian | fatigue crack growth rate (da/dN) | mechanical | backpropagation Neural Network |
| Al7075-T6, Al2024-T315, D16 aluminum alloy,  350 WT steel[[27](#_ENREF_27)] | aerospace and aviation sectors | SIF (stress intensity factor), stress ratio | gaussian function | crack growth rate | mechanical | radial basis function neural network |
| polyphenylene sulfide (PPS) matrix composites [[2](#_ENREF_2)] | tribological application | material compositions (PPS matrix, Short carbon fiber, TiO2, PTFE, Graphite), Testing conditions (Sliding speed, Applied pressure) | tansig, purelin | tribological characteristic (Coefficient of friction, Specific wear rate) | tribological | backpropagation Neural Network |
| microalloyed pipeline steels[[56](#_ENREF_56)] | petroleum industry | chemical compositions (carbon, magnesium, niobium, titanium, nitrogen, molybdenum, nickel, aluminum, copper, chromium, vanadium and carbon equivalent) and corrosion cell characteristics (reference electrode, scan rate, temperature, relative pressure of oxygen, pressure of purged CO2, chloride ion, bicarbonate concentration) | hyperbolic sigmoid, linear | corrosion current density and corrosion potential | chemical | backpropagation Neural Network |
| microalloyed API X70 line pipe steels[[36](#_ENREF_36)] | oil and natural gas industries | chemical compositions and tensile properties | hyperbolic sigmoid, linear | toughness and hardness | mechanical | backpropagation Neural Network |
| Ni–SiC composite coatings[[57](#_ENREF_57)] | various applications which need enhanced microhardness, better wear and corrosion resistance | ultrasonic power, SiC particle concentration, and current density | not clearly defined | weight loss | chemical | multilayer perceptron |
| prestressed, posttensioned concrete (PC)[[58](#_ENREF_58)] | civil engineering and structural engineering applications | stress level of concrete, water-cement ratio, cement-fine aggregate, cement-coarse aggregate ratio and testing age | not clearly defined for BPNN, radial basis function for RBFNN | carbonation depth | chemical | backpropagation neural network and radial basis function neural network |
| steel reinforced concrete[[20](#_ENREF_20)] | civil engineering applications | air temperature, AC resistivity over the steel bar, AC resistivity remote from the steel bar, DC resistivity over the steel bar | not clearly defined | corrosion current density | chemical | multilayer perceptron |
| mild steel[[59](#_ENREF_59)] | turning process | feed rate, cutting speed, depth of cut and acceleration of radial vibration | gaussian function | surface roughness | mechanical | radial basis function network |
| aluminum alloys[[60](#_ENREF_60)] | aerospace and other fields | altitude, annual average temperature, average humidity, annual average sunshine and annual rainfall | not clearly defined | atmospheric corrosion | chemical | backpropagation neural network |
| structural carbon steel[[61](#_ENREF_61)] | atmospheric environment | local temperature, relative humidity, amount of precipitation, pH of rainfall, air pollution by Sulphur dioxide and exposition time | not clearly defined | atmospheric corrosion weight loss | chemical | multilayer perceptron |
| AA3004 aluminum alloy[[42](#_ENREF_42)] | can making | the ratio of initial to final thicknesses, reduction, preheating time and temperature, finish rolling temperature and the final annealing temperature | logistic sigmoid | yield strength, elongation, ultimate tension strength, , | mechanical | backpropagation neural network |
| steel strip[[62](#_ENREF_62)] | hot strip mills | spray pressure, water flowrate, spray angle and vertical spray height | purelin, tansig | spray impact. | mechanical | backpropagation neural network |
| aluminium-based, particulate-reinforced metal matrix composites (MMCs) [[63](#_ENREF_63)] | structural applications | different SiC (µm) particle size ranges | sigmoid | density and tensile strengths | mechanical | multilayer Perceptron |
| IN625 alloy [[34](#_ENREF_34)] | aeronautic, aerospace, marine, petrochemical industries and gas turbine engine components | temperature, strain rate and strain | sigmoid (tansig and logsig) | flow stress | mechanical | multilayer perceptron |
| mild steel [[64](#_ENREF_64)] | chemicals and industries | Hammett constants, dipole moment, HOMO energy, LUMO energy, energy gap, molecular area and volume | sigmoid | corrosion inhibitor efficiency | chemical | multilayer perceptron |
| sandcrete materials [[35](#_ENREF_35)] | civil engineering projects | water-to-binder ratio (W/B), metakaolin addition (MK), binder (B), superplasticizer (SP), and ultrasonic velocity (UV). | logistic sigmoid, hyperbolic tangent | compressive strength, Modulus of Elasticity | mechanical | backpropagation neural network |
| structural carbon steel[[29](#_ENREF_29)] | atmospheric environment | local temperature, relative humidity, amount of precipitation, pH of rainfall, air pollution by sulphur dioxide and exposition time | not clearly defined | corrosion weight loss | chemical | multilayer perceptron |
| Ti–6Al–4V alloy[[65](#_ENREF_65)] | turning process | feed rate, depth of cut, acceleration amplitude of vibration in radial and tangential direction | tansig, Purelin | surface roughness | mechanical | multilayer perceptron |
| carbon steel manganese [[66](#_ENREF_66)] | oil industry | time, temperature and inhibitor concentration of inhibitor | gaussian function | real and imaginary parts of impedance | chemical | radial basis function neural network |

### ANN for Materials Performance Prediction

To predict materials properties is a challenge in materials related fields because it relies on a large number of variables that are in complex nonlinear relation with each other. ANN are empirical models that can elucidate any unknown correlation in physical models and satisfy complex problems in materials [[67](#_ENREF_67)]. Therefore, ANN was given an attention of materials researchers for predicting properties related with mechanical [[28](#_ENREF_28)] , tribological [[68](#_ENREF_68)] and wear [[69](#_ENREF_69)]. There were also many published satisfactory ANN predictions dealing with corrosion related problems [[70](#_ENREF_70), [71](#_ENREF_71)]. Some experimental techniques such as corrosion test, tensile test, theoretical deterministic methods, metallographic corrosion characterization, mechanical testing and AC impedance spectroscopy, etc. were used to acquire the data for ANN training and result verifying.

Regarding to space requirement, it is not possible to describe all ANN applications in details so that some of them will be grouped and reviewed into different sections.

1. **Corrosion**

Electrochemical reaction on materials surface by nature is known as corrosion. The atmosphere can cause a big deal in corrosion degradation process of materials. Statistical regression analysis models mostly used for prediction cannot give superb performance because complexity and non-linearity of meterological variables on atmospheric corrosion is apparently high. To observe local atmospheric corrositivity, in the region of São Luis, five Atmospheric Corrosion Stations (ACS) were installed to monitor the local atmospheric corrosivity and three representative metallic specimens of low-carbon steel, copper and aluminum were exposed to the atmosphere for almost two years. According to corrosivity degree measurement of ACS test sites, the corrosion rates and thickness losses of low-carbon steel, copper and aluminum were respectively categorized into three corrosion metrics: generalized corrosion, alveolar corrosion and localized corrosion. A mathematical model based multilayer perceptron ANN was established by Kenny, Paredes et al. (2009) [[24](#_ENREF_24)], with linear and sigmoidal tangent transfer function at the first hidden layer and logarithmic transfer functions at the second intermediate layer by using Levenberg–Marquardt backpropagation training algorithm, to analyze corrosion rates of three different matereials. Meterological data of ten specific environmental variables (time of exposure, temperature, relative humidity, precipitation, time of wetness, wind velocity, solar radiation, chloride ions, sulfur compounds and dustfall) collected were used as the input to train ANN and laboratory evaluated corrosion rates of specimens were used as the output to analyze the atmospheric corrosion rates. The good correlations obtained in the analysis demonstrate ANN performed greatly in estimating the corrosion rate. It supports one major benefit of ANN that ANN performs well in estimating high complexity and nonlinear correlation between input and output parameters. Long-term corrosion prediction could be calculated by Pourbaix equation[[72](#_ENREF_72)] evaluated with parameters derived from ANN analysis for two year period.

To deal with different corrosion metrics of different materials, Kamrunnahar and Urquidi-Macdonald (2009) [[43](#_ENREF_43)] developed more than one multilayer perceptron backpropagation ANN model to predict different kinds of corrosion behavior. Three ANNs were modelled for two general corrosion data on iron-alloy, and carbon and alloy steel, and one localized corrosion data on Ti-alloy. The ANN model developed for grade-2 titanium showed excellent agreement with experimental data and then used for prediction of future maximum pit depth and other variables under similar operating conditions, giving very good agreement results with experiments. In 2010, Kamrunnahar and Urquidi-Macdonald have also done an extended work [[26](#_ENREF_26)] of their previous work [[43](#_ENREF_43)]. But in that work, they emphasized on different corrosion metrics of one material. They established three different neural networks for localized and general corrosion of alloy 22, with the best network structures of 8-5-5-1 for localized cervie corrosion, of 2-100-100-10-4 for localized corrosion, AC impedance data and of 4-5-5-1 for general corrosion study. The aim of both works was to predict corrosion behavior of metal alloys such as metallic glasses, carbon steel, and grade-2 titanium as well as alloy 22. Data from two different sources (publicly available publications, etc. and personal communication) were together brought into ANN models. Both works have shown that ANN prediction results were good agreement with experimental data.

As a note came up from above materials corrosion predictive researches, more than one ANN models can differently and separately be developed for different aspects of the problem. Kenny, Paredes et al. trained an ANN model to analyze corrosion rates of three different matereials. Kamrunnahar and Urquidi-Macdonald constructed three separate ANN models to observe different corrosion behaviors of three materials in one research and three different neural networks to determine different corrosion metrics of one material in their another work. No matter what the number of ANN models are, the critical point to take account is that more data points would be needed to establish a more robust and reliable ANN model with higher confidence and more representative results.

1. **Thermal Property**

Multivariate nonlinear regression method have been used to represent complicated interaction of flow stress with strain, strain rate and temperature. But the complicated correlation of flow stress with their affecting factors (microstructure and processing parameters) is difficult to be evaluated by any simple constitutive model. To overcome the repeated calculation of regression parameters for new experimental data in simple constitutive model, Yang, Zhu et al. (2012) [[22](#_ENREF_22)] developed an accurate thermal stress model to solve the problems of quench distortion of parts by ANN with a back propagation learning algorithm. In this model, the author used batch training for two network training types: sequential training and batch training. Batch training of backpropagation neural network (BPNN) with the best network structure (3-6-8-1) including two hidden layers provided about 3% error rate between the predicted output and the actual output. According to a strong positive linear correlation relationship between predicted data and experimental data by correlation coefficient calculation, the model was chosen to achieve good network performance for predicting for thermal stress of quenched A357 alloy with a higher precision. In this work, the authors were concerned with training types of ANN so that stability was found in performance of network.

1. **Structural Property**

Sasikumar, Rajendraboopathy et al.(2008) [[39](#_ENREF_39)] presented a BP neural network, that had the best training result with 66-45-1 network architecture, and AE (Acoustic Emission) amplitude frequency data collected up to 50% of failure loads were used as input to BP neural network for failure load (ultimate strength of unidirectional T-300/914 tensile specimens) prediction. Levenberg-Marquardt algorithm and the linear transfer function were used for establishing ANN. In this work, it can be found that the network could not predict the ultimate loads outside the training range in trained network. At this point, it made incapability of the network because it is necessary to subsequently add the AE data of specimens with extreme error rate that lies beyond the training range into the training set, thereby decreasing the acceptable maximum error rate from 8.37% to 1.22%. Other key parameters such as duration, energy, events and counts were also used along with BP Neural Network for material characterization and structural integrity evaluation for further analysis. From analysis, it was found that amplitude frequency is the most affecting factor in predicting failure load of composite hardware.

The author have done another similar work in [[73](#_ENREF_73)] in which three different BP neural networks were built for predicting failure strength of composite tensile coupons using acoustic emission technique. But up to 30%, 40% and 50% of failure loads were used as input to three neural networks. Three optimal network structures are 66-37-1 constructed with 30% of AE data, 66-22-1 constructed with 40% of AE data and 66-45-1 constructed with 50% of AE data, respectively. All of them applied the same network parameters like learning rate of 0.01, momentum of 0.9, hyperbolic tangent transfer function, and backpropagation training algorithm. However, the comparison of three networks showed that the network trained with more data could provide better performance.

The purpose of both works was to to reduce the structural degradation of composites. Reducing the proof test load is the only way for structural degradation. By ANN analytical models, the prediction results of tensile strength in both works indicated that proof test can be done on more sophisticated composites at lower loads so that the actual failure load of composite hardware was predicted. But there was same finding from both works that inadequate training data did not produce the optimal prediction.

1. **Electrochemical Property**

Bassam, Ortega-Toledo et al. (2008) [[74](#_ENREF_74)] and Colorado-Garrido, Ortega-Toledo et al. (2008) [[75](#_ENREF_75)] implemented prediction ability of multilayer perceptron neural networks in the application of electrochemical impedance techniques. Bassam, Ortega-Toledo et al. trained a multilayer perceptron neural network model with 4-5-1 network architecture, Levenberg–Marquardt learning algorithm, the hyperbolic tangent sigmoid transfer function and the linear transfer function for the determination of different types of corrosion by using electrochemical impedance spectroscopy curves from different inhibitor concentrations. Correlation coefficient of R>0.9905 described good agreement between simulations and theoretical data test were in good agreement. It was expected that the model may also enable the implementation of smart sensors for online quality determination (corrosion type) in pipeline steel. Colorado-Garrido, Ortega-Toledo et al. described the application of multilayer perceptron neural network to electrochemical testing by predicting impedance spectroscopy values, with the same network parameters used in the work of Bassam, Ortega-Toledo et al. The best network architecture was obtained with 2-5-1 for 5 ppm of inhibitor and 2-2-1 for 25 ppm of inhibitor, and correlation coefficients R value of 0.984 for 5 ppm and 0.994 for 25 ppm described good agreement between simulations and theoretical data test were in good agreement. In this case, the model’s aim was to predict the imaginary impedance based on the real part of the impedance as a function of time to determine the corrosion resistance of the tested material in the solution. By using the same network parameters, the performance results of those two works gave good agreement between actual and predicted data. Hence, it can be concluded that a good ANN model can progressively be used for future prediction in context of the problem with same nature.

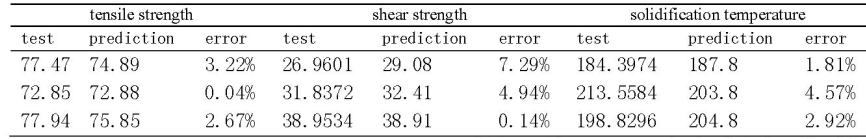
1. **Tribological Property**

Predicting tribological properties of polymer composites to determine performance potential and is a challenge in materials science because the available experimental dataset is relatively small, noisy, complex and highly non-linear. ANN can make the problem-independent solution due to its advantage of nonlinearity, adaptive learning, and generalization. Gyurova and Friedrich (2011) [[2](#_ENREF_2)] established two multilayer perceptron neural networks for predicting tribological properties (coefficient of friction and specific wear rates) of polyphenylene sulfide (PPS) matrix composites. A new measurement series was performed with thermoplastic matrix composites and preexisting five experimental training datasets collected or expanded throughout a period of three years were used for training and testing the developed network models. Material compositions and testing conditions were used as input parameter to predict the output parameters the tribological properties. The tansig function and purelin function were used as transfer functions for the hidden layer and the output layer respectively. The best network structures achieved with coefficient of friction and specific wear rate were 7-[3-1]2-1 and 7-[9-3]2-1, respectively. A user-friendly graphical interface was developed for performing network training and prediction and can easily predict even for unseen input. The network performance was evaluated by a comparison of the MRE (mean relative error) generated after training of the ANN with the different training datasets, thereby improving significantly with the enlarged datasets. The satisfactory ANN predicted results had good correlation with the measured values. The optimized ANN models was also used to analyze the impact of SCF (short carbon fiber), sub-micro TiO2, on the sliding friction and wear properties of PPS composites. It gave the excellent generalization of ANN without performing exhaustive experimental work. When those ANN models were also applied for studying the effect of internal lubricants (Gr and PTFE) in hybrid polymer systems, good correlation existed between actual and predicted output. This work highlighted the advantages of ANN, especially generalization ability, by implementing the existing ANN models for other problems with similar nature.

### ANN for Materials Design

The relation among composition, process and properties of new materials design are influenced by many complicated factors. In the context of materials science, ANN performs a great job for prediction and function approximation with respect to materials’ performance, but there are not many ANN applications enough for forecasting materials properties for materials design. Li, Qi et al. (2007) [[45](#_ENREF_45)] studied a new route for the designing of lead-free solder that can find out the influence of adding input variables into tensile strength, shear strength and solidification temperature of Sn-Ag alloy by using multilayer BP Neural Network. The best network structure 7-40-40-40-1 with three hidden layers was chosen. Data from one reference came into the ANN for training and testing. The analysis result in Table 3 showed that the error rates between expected output and predicted output was within 5%, but only one of them gave 7% error rate because of less amount of training data, unexact measurement of shear strength and poor-proportionally distributed data. This work also indicated that more accurate prediction results can be obtained with big amount of training data. However, ANN still gave good agreement with given experimental conditions and it was verified for making the feasible estimates of the property of the lead-free solder.

**Table 3**. Comparison of test data and predict data [[45](#_ENREF_45)]



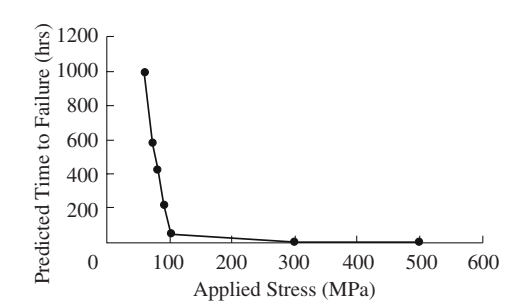
Xie, T. and J.C. Grossman (2018) [[1](#_ENREF_1)] developed a highly accurate prediction model of density functional theory (DFT) calculated properties from Materials Project [[76](#_ENREF_76)], with a database of a diverse set of inorganic crystals ranging from simple metals to complex minerals (including 46 744 materials covering 87 elements, 7 lattice systems, and 216 space groups). It was a crystal graph convolutional neural networks (CGCNN) framework, an innovated universal and interpretable representation of crystalline materials, to directly learn material properties from the connection of atoms in the crystal and discover empirical rules of materials design, that were consistent with the common knowledge for discovering more stable materials. Thus, the search space for high-throughput screening could significantly be reduced by additional information provided by CGCNN for material design. Traditional methods used to represent the arbitrary size of crystal systems as a fixed length vector was well replaced by CGCNN by grabbing the benefit of deep learning. The authors also designed a new convolution function to make learning deeper networks easier, that improved generalization ability of CGCNN as well as prediction performance of different materials properties. Compared to previous statistical learning (SL) framework presented by Jong et al. [[77](#_ENREF_77)] that implemented multivariate local regression to predict elastic properties on the same dataset, CGCNN provided similar results, but it had advantage of predicting properties by extracting features only from the crystal structure. And when CGCNN model was used for 1585 new crystals that were recently added into Materials Project database, it could also provide a good generalization to materials from potentially different crystal groups. It was a kind of evidence that ANN has good generalization ability. CGCNN could also predict discrete properties as well as the classifications of metal and semiconductor with the same framework by using a softmax activation function for the output layer and a cross entropy cost function.

Most current ANN applications highlighted the advantage of feed forward neural networks and ANN performance results were pointed out being superior or similar in making comparison with experimental results. The authors in [[1](#_ENREF_1)] just brought a great contribution into materials science by implementing the concept of feedback neural network and deep learning to mine valuable knowledge from a large dataset to accelerate the design of crystalline materials. It can be found that the use of dynamic neural networks in materials related problems could also gave satisfactory results as great as static neural networks.

### ANN with Other Computing Paradigms

Integration of neural networks with other computing paradigms, such as Bayesian framework, fuzzy logic, sensitivity analysis and genetic algorithm, can also enhance the performance of ANN. Cai, Cottis et al. (1999) [[41](#_ENREF_41)] used multilayer perceptron ANN analysis integrated with sensitivity analysis to analyze phenomenon of atmospheric corrosion behavior of steel and zinc. This work investigated a system with environmental variables including temperature (T), time of wetness (TOW), exposure time and concentrations of Sulphur dioxide (SO2) and chloride (Cl-), and found out the complex interactions between those variables and atmospheric corrosion. The database was big enough to be used for this work, including information from corrosion tests, climatological tables and rainfall data from thirty-three countries around the world. But those data were measured by different methods in different sources so that corrosion data might have inherently scattered. According to sensitivity analysis, almost linear relationship between TOW, SO2 (the most affecting input variables on atmospheric corrosion in industrial environment) and atmospheric corrosion of steel and zinc were founded at a specified condition. The network structure (5-8-1) with one hidden layer and sigmoid transfer function showed that the ANN results accounted for approximately 70% of the variance seen in the real data sets. Although this work did not consider about other possibly more important parameters on atmospheric corrosion (such as effect of microclimate) for prediction model, it can be seen that ANN could still give more satisfactory results by its fault-tolerance ability than previous linear regression analysis.

To find out the solution to the challenge to predict the time to failure of 304 stainless steel as a result of SCC in aqueous chloride solution, Lajevardi, Shahrabi et al. (2009) [[25](#_ENREF_25)] have implemented a similar approach adopted by Cai, Cottis et al, in which a multilayer perceptron ANN with backpropagation algorithm was constructed and evaluated using the collection of experimental data associated with the SCC (stress corrosion cracking) of AISI 304. The data used to train the network were extracted from 24 references. The best network structure was (3-4-4-1) with two hidden layers and sigmoid transfer function. In this work, Montecarlo optimization have been used to evaluate the initial set of weight values to reduce the local optima phenomenon. Both of those works have conducted sensitivity analysis to improve the network efficiency by demonstrating the effects of the important variables and showed the possible effects of the other affecting variables. Sensitivity analysis showed in Fig 4 that among three input parameters (applied stress, temperature and chloride ions concentration), applied stress was regarded as the most influencing parameter on time to failure. Those ANN models can account for about 70% and 74 % of the variance of the experimental data, but these two works have some limitations in unseen possible affecting variables, unexplained variances in prediction and data accuracy.



**Fig 4.** Effect of applied stress on time to failure (Cl– = 2100 ppm and a = 130 MPa) [[25](#_ENREF_25)]

Xia, Nie et al. (2016) [[44](#_ENREF_44)] used another effective input ranking method called fuzzy curve to rank the importance of input parameters that have great impact on the output. The authors built two BP neural networks for investigating non-linear correlation and predicting hardness and corrosion rate of magnesium alloy as a function of compositions. ANN prediction results demonstrated that there was an excellent correlation for hardness and although correlation for corrosion rates was lower than hardness, it was still considerably good result because of the inherently chemical activity of Mg and complex interactions between alloy additions. Fuzzy curve analysis was also used in integration with ANN to analyze the impact of each alloying element on the performance of alloy. The sequence of the alloying elements in terms of their ability to develop Mg alloys was ranked with relatively low alloying level for high hardness as Ca ≥ Zr  Sr Zn Gd whereas fuzzy curves gave complex results for corrosion rate, but the impact of each alloy element on corrosion rate were respectively described. It can be seen that non-linear correlations have been existing between hardness and corrosion rate. However, fuzzy curve analysis could improve the performance of ANN with satisfactory accuracy and the model can be used for corrosion prediction of other existing and future Mg alloys.

ANN is a technology that can learn by examples and then make prediction for new situations. ANN model needs sufficient amount of adequately extensive and representative data to constitute a good training set to solve the problem. Many unnecessary input parameters that do not have a strong interaction with desired output in training data makes an exhausted training and degrades the network performance. In materials science researches, it is critical to know the importance of each input variables affecting on material properties and identify which input variable is the most influencing factor on materials’ properties. By using input ranking methods like sensitivity analysis and fuzzy curve, unimportant input parameters will easily be found out to eliminate from the problem and then model will be made simpler and easier to define future possible mechanisms, and the solution will be more fruitful and exact. And modelling ability of ANN can be improved by considering all other possible variables that have stronger impact on output at a given condition and by constructing a more enriched dataset.

The benefit of other computing technologies, such as genetic algorithm and Bayesian framework, can save the defects of ANN model, like overfitting occurred while training the network and uncertainty and instability of ANN, and then the performance of network is significantly improved. That fact is supported by the work of Huang, Jia et al. (2018) [[50](#_ENREF_50)] in which the authors have made a comparison of three different approaches to predict flow behavior and find out the optimum hot working processing parameters of 5754 aluminum alloy. Three forecasting models: strain-compensation Arrhenius (SA), back propagation (BP) artificial neural network (ANN) and an optimized BPANN model based on genetic algorithm (ANN-GA) were constructed. Their comparison showed that four-layered BPANN model provided accurate prediction than SA model. But it was found that BPANN model was weak to optimize globally so that local optimum occurred in learning algorithm. This weakness made instability in network performance. BPANN was optimized by genetic algorithm (GA) to form a new model with network structure (3-10-8-1) that produced high accuracy and stability (Correlation Coefficient R=0.9999 and Average Absolute Relative Error ARRE=0.0232–1.0485) within the whole deformation range by calculating the fitness values of each individual in GA by the predictive error norm of the neural network. Therefore, ANN-GA model was found as the most efficient and accurate model among three models because genetic algorithm (GA) can remove the uncertainty of ANN, improve its stability and accuracy by searching for the optimal weight and threshold for ANN at the same time.

In [[62](#_ENREF_62)], Kermanpur, A., A. Ebnonnasir, and M. Hedayati (2007) used data set from the hot strip mills of Mobarakeh Steel Complex to construct a novel analytical predictive framework, named analytical–artificial neural network (AANN) model to estimate the spray impact from spray angle, spray pressure, vertical spray height and water flowrate. The predictive model was coupled with Bayesian regularization that can reduce the overfitting problem of Levenberg–Marquardt ANN learning algorithm and improve the network prediction ability for unseen data. The combined network can be used as a proper tool to evaluate the efficiency of a high pressure (HP) hydraulic descaling operation using flat spray nozzles in terms of achieving the highest spray impact under any process condition. With an optimal network structure of one input layer with four input variables, two hidden layers and one output layer with one output variable, a maximum amount of output (spray impact) was obtained. The prediction result showed good agreement with correlation coefficient R2 of 0.992. In this work, sensitivity analysis was also carried out in which input parameter spray angle resulted in a higher impact is followed by the spray height. The authors were also progressively using their current AANN model to predict the effect of all input parameters on the spray overlap.

In recent works, we can see that there are many successful ANN applications in materials related problems. Herein, the hybrid system, the joint implementation of other computing methodologies with ANN, can make satisfactory improvement in ANN performance compared to simple ANN model alone.

### Other Powerful ANN Models in Materials Applications

The other powerful and efficient neural network models rather than multilayer BPNN have also brought successful prediction results in recent works to determine the tensile mechanical properties of materials. Yilmaz and Ertunc (2007) [[78](#_ENREF_78)] introduced a neural network model, named generalized regression neural network (GRNN), to predict tensile strength of steel wires by using all the production parameters (the amount of carbon, ferrite and inclusions) and the diameter of steel wires. That approach was constructed with four layers: input layer, a layer of radial basis centers, a linear layer of regression unit and the output layer. A very good statistical performance with a 0.96 correlation coefficient between the target data and the predicted output have demonstrated the result of good agreement to determin the tensile strength of steel with high accuracy and reliability.

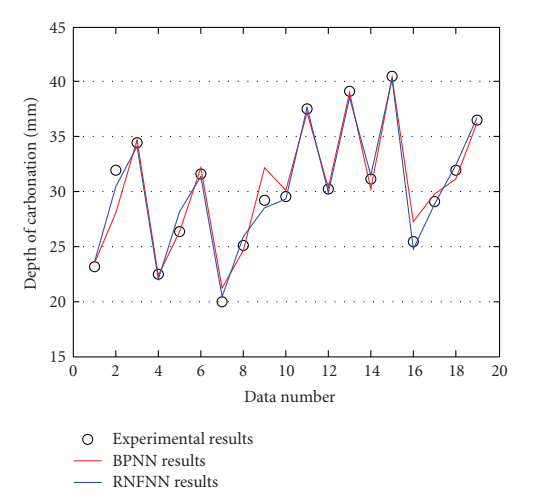
Kappatos, Chamos et al. (2010) [[23](#_ENREF_23)] adopted a radial basis function (RBF) neural network, one of the mostly used feed forward network models after multilayer network, to assess the effect of existing corrosion damage on the tensile behaviour of the wrought magnesium alloy AZ31 by predicting the time dependency of the tensile mechanical properties degradation. The data used to train the network was derived from extensive experimental investigation. The authors used only two input parameters (pit depth and pitting density) that had stronger discriminant capability on output (elongation of fracture and ultimate tensile strength). Although there was no need to determine the number of hidden layers for RBF, extensive trials were carried out to determine the optimum number of hidden neurons to achieve the best prediction accuracy. Conventional linear modelling techniques fully optimized the simple linear transformation at the output layer of RBF network. Greater RBF networks with 40 and 20 hidden neurons gave higher prediction accuracy from 99.61% to 99.83% for elongation of fracture and from 99.27% to 99.68% for ultimate tensile strength, respectively. ANN results made a comparison between two input parameters such that more accurate output tensile strength was achieved with pitting density while the pit depth had stronger effect on the decrease of the mechanical properties than pitting density. RBF neural network prediction results were very good agreement with experimental results.

Most of successful materials-related ANN researches have been done with multilayer backpropagation neural networks. Here, it can be found that other ANNs are as powerful and efficient as multilayer backpropagation neural networks in prediction performance. Regarding to physical background of the problem, input parameters should be selected as important features that have a strong relation with output parameters so that computational cost and size of training data can be reduced to improve the classification performance.

### ANN Models in Comparison

In some literatures, ANN prediction result was compared to not only experimental results but also other feed forward ANN models’ results. In turning process, to control the desired surface roughness by manufacturing makes high expense. To establish a functional relationship between process parameters of turning process and surface roughness, multilayer perceptron (MLP) neural networks have been mostly implemented for a decade. But MLP needs to determine the number of hidden layers and hidden neurons in each layer with a costly trial-and-error approach. Unlike MLP, as radial basis function (RBF) neutral network has the advantage of having only one hidden layer and the number of hidden neurons can automatically be adjusted in learning algorithm, less computational time and simpler training can give faster and more accurate results. Sonar, Dixit and Ojha (2005) [[59](#_ENREF_59)] used the advantage of RBF neural network to construct a predictive model for the surface roughness in a turning process and its performance was compared with the reported performance of a multilayer perceptron (MLP) neural network by Kohli and Dixit [[79](#_ENREF_79)] as well as experimental results. The results showed that experimental values exist between the range of the lower and upper estimate obtained by fuzzy linear regression model and the most likely estimate in most cases is close to experimental value. Also, compared to MLP, the root mean squares errors of RBF are nearly the same as MLP. But RBF made slightly inferior performance against MLP and with a few testing data, the errors were more than 20%. The authors discussed about increasing the number of training and testing data to increase the network accuracy.

Another similar comparison of different ANN models was found in [[58](#_ENREF_58)]. Lu and Liu (2009) analyzed the carbonation depth of prestressed concrete under different stress states by adopting two feed forward ANNs, BPNN and Radial Basis Function neural network (RBFNN), together with comparison in their performance and comparison with experimental results as well. Training data and test data for two ANNs were produced by an accelerated carbonation practical experiment of prestressed concrete. The 5-10-1 structure of BPNN is the optimal network structure for prediction. RBFNN was also established with the same number of input and output parameters. RBFNN results have the largest absolute percentage error of 8.46% whereas the largest absolute percentage error of BPNN is 10.88%. In this work, although RBFNN gave better precision than BPNN, the precision level of BPNN can be improved by adding the number of hidden layers appropriately that will lead to a costly trial-and-error approach. Because the error rates of both models were acceptable and their estimation results were good agreement with experimental results shown in Fig. 4, ANN has a high generalization ability to analyze and predict carbonation depth in concrete. The authors suggested that RBFNN is a better model for predicting carbonation depth in concrete and solving other civil engineering related problems as well. However, they reported that enough training and testing data was still needed to construct a perfect ANN for providing a new practical approach to concrete carbonation analysis.



**Fig 5.** Relationship between the two ANNs’ training and testing results and the experiment results in [[58](#_ENREF_58)]

However, there is no defined universal ANN model that always outperforms on other ANN models. The right choice of ANN model should be made in accordance with the problem to be solved at hand. The network performance relies on the level of complexity and nonlinearity of input-output mapping as well as the amount of training and testing data. Moreover, to make a result comparison with experimental results for optimal result, ANN models must be trained and tested with the same data set as experimental data.

### Summary

Materials data are becoming so abundant that their relationship is quite complicated to handle well. Traditional linear regression analysis is mostly not able to handle a large number of variables and nonlinear relationship among them. The main benefit of ANN over conventional regression analysis is that ANN can model problems in which there is no clear relationships between inputs and outputs and construct the solutions that can not be easily formulated within short time without specifying the form of interactions between variables. From previous works, it can be said that ANN is a good estimator for materials related problems against statistical linear regression analysis due to its advantages of fault-tolerance, noise-tolerance and generalization ability. It can solve both nonlinear and linear problems with high accuracy. As a disadvantage, although ANN has benefit of parallel structure, it sometimes spends long training and operating time and occurs instability when applied to larger problems. Important features that have a strong relation with output parameters should carefully be selected as input into ANN training such that unnecessary computational overhead and increased size of training data can be reduced to attain high classification performance.

But it is critical to be noted that misuse of over complicated networks cannot give the optimal results. ANN should be correctly implemented according to types of problems to be solved to avoid the misuse of ANN benefit. If the number of hidden neurons, compared to the number of data pairs in evaluating the fitting parameters, are not enough as in [[63](#_ENREF_63)] [[80](#_ENREF_80)] or more redundant as in [[42](#_ENREF_42)], it will lead to the misuse of ANN. Moreover, it does not mean the more complicated the networks, the better the results. The number of neurons in hidden layers of neural networks should also be increased within a limit depending on the problem, otherwise; mathematically undefined situation, known as overfitting, in which determining more fitting parameters than the available data points is difficult, will make poor network generalization ability on unseen data. The joint implementation of ANN and other computing paradigms should further be considered due to their good cooperation results.

# Trends of the ANN application in materials area

At present, researchers are trying to implement ANN models to predict the performance, stability and structure of materials, that is also a great challenge in materials science. According to [[81](#_ENREF_81)], in 2020, materials industries, that supply steel, petrochemical products, cement and etc., will be the key technology of automotive industry to develop new structural materials. ANN can accelerate the design of new materials by predicting material properties with high accuracy. To combine materials science with computer science by ANN prediction for materials properties is a great solution for researchers not only at present but also in future.

Previous works described their network parameters: network structures, training algorithms, transfer functions and data used to train and test ANN, in related researches so that those works can be expanded and enhanced by new researchers. Since their prediction results gave satisfactory and acceptable outcomes against experimental results, ANN prediction models can significantly reduce the quantitative experimental works. But most recent works showed large amount of data are still needed to train ANN. ANN performance can be extended when there is data big enough for constructing an optimized neural network model and then the prediction results will be more exact for future use.

Most neural network models in context of materials science are feed forward types and other popular, powerful and efficient feedback neural networks (Recurrent Neural Network, Deep Convolutional Neural Network, etc.), that are currently widely implemented in image and video recognition, speech recognition, automated robotics and machine translation applications in computer science researches, would be future research trends in materials science too. Although there are currently few materials researches implementing feedback ANN, it is still necessary to conduct more researches related with those networks to give strong recommended results to new researchers. Hence, the implementation of ANN in the prediction of material properties may achieve higher score and find more application prospects in the future.

The purpose of this paper is to reveal the significance of ANN in many aspects of materials related problems and it can be seen that ANN prediction accuracy was apparently improved in recent researches compared to traditional consecutive equations. It is expected that this paper can give a hand to materials researchers with a comprehensive review of some applications of neural network in the context of materials science.

# Acknowledgements

Acknowledge the support from the project of Prof. Shi.

And I would suggest adding also the following projects:

We also thank the financial support from the Ministry of Science and Technology, China for the national key research and development plan (2017YFB0701904). And the authors are also grateful for support from the 111 Project (Grant No. B12012) in China for promoting international exchange.

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