On the Optimal Node Ratio between Hidden Layers: A Probabilistic Study

Alan J. Thomas, Simon D. Walters, Saeed Malekshahi Gheytassi, Robert E. Morgan, and Miltos Petridis

Abstract—Two-hidden layer feedforward neural networks (TLFNs) have been shown to outperform single-hidden-layer neural networks (SLFNs) for function approximation in many cases. However, their added complexity makes them more difficult to find. Given a constant number of hidden nodes \( n_h \), this paper investigates how their allocation between the first and second hidden layers \( (n_{h1} + n_{h2}) \) affects the likelihood of finding the best generaliser. The experiments were carried out over a total of ten public domain datasets with \( n_{h1} = 8 \) and 16. The findings were that the heuristic \( n_{h1} = 0.5n_h + 1 \) has an average probability of at least 0.85 of finding a network with a generalisation error within 0.18\% of the best generaliser. Furthermore, the worst case over all data sets was within 0.23\% for \( n_{h1} = 8 \), and within 0.15\% for \( n_{h1} = 16 \). These findings could be used to reduce the complexity of the search for TLFNs from quadratic to linear, or alternatively for ‘topology mapping’ between TLFNs and SLFNs, given the same number of hidden nodes, to compare their performance.

Index Terms—ANN, optimal node ratio, topology mapping, two-hidden-layer feedforward, function approximation.

I. INTRODUCTION

Since the introduction of fully interconnected feedforward neural networks into the automotive industry in 1993 [1], they have enjoyed increasing popularity and are still widely used for function approximation to date, as in for example, [2]. The most popular training algorithm by far is the ‘trainlm’ algorithm available in the Matlab neural network toolbox. This is an implementation of the Levenberg-Marquardt algorithm [3], which generally yields the best generalisation and fastest convergence for function approximation problems [4]. It is widely used for training both SLFNs [5]-[8] and TLFNs [9], [10], though the former is more common in the literature. Unfortunately it is very memory hungry, and the training time rises exponentially with the number of weights. This makes it unsuitable for networks with more than a few hundred weights [4], and this is particularly problematic for networks with two hidden layers. Because of this, the complexity of an exhaustive search through two hidden layers is actually greater than \( O(n^2) \). By way of illustration, whereas an exhaustive search through a single hidden layer with 1 to 32 neurons takes less than an hour, a search through two hidden layers takes approximately 48 hours. This for 30 networks of each topology on a Quad-core i7 4710HQ processor running at 2.5GHz.

This study investigates whether there might be a faster route to take in the search for a TLFN rather than the conventional raster scan. For example only; taking the ‘short cut’ route shown by the solid line in Fig. 1 would reduce the complexity of the search from quadratic to linear since only 8 topologies are tested instead of 64. However, in order for this method to be useful there must be a high probability that it will find networks with only a small penalty in generalisation error compared to the longer raster route. Furthermore, this should still be the case regardless of the dataset.

In the experiments, ten public domain datasets suitable for function approximation tasks are tested to find a ‘short cut’ which works well for all. By suitable, it is meant that they have a single output. Although many examples in the literature have multiple outputs (e.g. [9] [10]), this should be avoided if possible since the validation error is calculated across all outputs. This means the training might stop prematurely due to the poor generalisation capability of a single output, perhaps resulting in sub-optimal generalisation of the other outputs. There is also mathematical evidence that reducing the number of outputs increases the storage capacity of the network [11]. If multiple outputs are required, it is better to split the network into several sub-networks each with a single output. These could run in parallel on hardware or multi-threaded software, and are likely to require fewer nodes each, especially for those where the functions are simpler, and/or for those which require fewer inputs.

II. RELATED WORK

Two-hidden-layer feedforward neural networks have gained popularity ever since it was shown that they can outperform those with a single hidden layer [12]. Related literature concerning the allocation of nodes between hidden layers can be classified as mathematical proofs, empirical studies and anecdotal. These are discussed in the following...
sections:

A. Mathematical Proofs

These are concerned with proving an upper bound for the total number of hidden neurons needed to exactly reproduce the training data, or to do so with negligible error. Using Akaike’s information criteria, Tamura and Tateshi [13] proved that including the output neuron, the upper bound for SLFNs is \( N_o \), and that for TLFNs this is \( N_o / 2 + 3 \) where \( N_o \) is the total number of training samples. This not only proves that the upper bound for TLFNs is lower than that for SLFNs for a given dataset, but suggests that this would be achieved using \( N_o / 2 \) neurons in the first hidden layer and two neurons in the second hidden layer.

Inspired by the work in [13], Huang constructed a proof using a network consisting of different neural subnetworks each playing different roles [11]. He rigorously proved that the upper bound on the number of hidden nodes \( N_h \) for TLFNs with sigmoid activation function is given by equation (1), where \( N_o \) is the number of outputs. These can learn at any \( N_i \) distinct samples with any degree of precision. Conversely, the storage capacity \( N_i \) samples of a TLFN with \( N_h \) hidden neurons is at least that defined in equation (2).

\[
N_h = 2\sqrt{N_o + 2}N_i \quad (1)
\]

\[
N_i = \frac{N_h^2}{4(N_o + 2)} \quad (2)
\]

This proves that reducing the number of outputs increases the storage capacity of a TLFN and that for the best storage capacity, function approximators should be constructed with a single output. Huang further specifies that the total number of hidden neurons \( N_h \) should be allocated between the first and second hidden layers (\( N_1 \) and \( N_2 \) respectively) according to equations (3) and (4).

\[
N_1 = \sqrt{(N_o + 2)N_i} + 2\sqrt{N_i/(N_o + 2)} \quad (3)
\]

\[
N_2 = N_o\sqrt{N_i/(N_o + 2)} \quad (4)
\]

For a single output, i.e. \( N_o = 1 \), equations (1), (3) and (4) can be rewritten as (5), (6) and (7) respectively.

\[
N_h = 2\sqrt{3N_i} \quad (5)
\]

\[
N_1 = \sqrt{3N_i} + 2\sqrt{N_i/3} \quad (6)
\]

\[
N_2 = \sqrt{N_i/3} = N_o - N_i \quad (7)
\]

Substituting \( N_h - N_1 \) from (7) into (6), and gathering terms leads to equation (8).

\[
3N_1 = \sqrt{3N_i} + 2N_h \quad (8)
\]

Finally, by substituting \( 3N_1 \) for \( N_i / 2 \) from equation (5) and rearranging, the ratio of nodes between the first hidden layer and the total number of hidden nodes is given by equation (9). So for a single output Huang’s proof suggests that a TLFN should have a wide first hidden layer and a narrow second hidden layer with a ratio of 5:1 between them.

\[
N_i / N_h = 5/6 \quad (9)
\]

However, this ratio relates to the upper bound on the number of nodes, and is merely a consequence of the way that Huang constructed the network for his proof. Another point to consider is that his work is concerned with exactly reproducing the training data (including any noise). So at this upper bound, any network would definitely be overfitting unless a suitable regularisation scheme such as weight decay or early stopping were deployed. The same can also be said for the work of Tamura and Tateshi [13]. This current work differs from both in that it is an empirical study and arrives at a different conclusion for the ten public domain datasets used.

B. Empirical Studies

In a novel study, Thomas et al. [14] compared every possible combination of hidden nodes in the first and second hidden layers for a given constant number of hidden nodes. Mathematically speaking, this is equivalent to all possible \( n_1 + n_2 = n_h \), where \( n_1 \) and \( n_2 \) are the number of hidden nodes in the first and second hidden layers respectively and the total number of hidden nodes \( n_h \) is constant. This equation describes any of the backward diagonals in Fig. 1. This was repeated for all the different values of constant \( n_h \) which is given by the set \( n_h = \{34, 20, 16, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3\} \). When the average generalisation errors were plotted centred on the forward diagonal \( n_1 = n_2 \), (alternatively \( n_1 = 0.5n_h \)) the result is an ‘isonode’ map as shown in Fig. 2. Each solid contour line is referred to as the \( i^{th} \) isonode, where \( i = n_h \) is the number of hidden nodes. Thus the first contour line in the set, and leftmost on the map is the 34\(^{th}\) isonode.

Over two different datasets, it was concluded that the relationship which gives the lowest average generalisation error is given by:

\[
n_i = \text{int}(0.5n_h + 1), n_2 = n_h - n_i \quad (10)
\]

Although the current study can be viewed as an extension of this work in as much as it shares the concept and basic method of data gathering, it is a new study which uses a completely different (probabilistic) approach.

![Isonode map](image)

Unlike the previous study, it uses relative generalisation errors which permits direct comparison between node combinations across the entirety of the datasets. This in turn allows probabilistic methods to be used by way of a frequency analysis of the data. Another major difference is five times as many datasets were used in the current study.

C. Anecdotal

In the absence of any other related work, it was decided to
examine what actual node combinations other researchers in the automotive field had selected on for their TLFN designs. This indirect evidence can only be considered anecdotal because in many cases there is no information at all about how these were designed, such as [15] [16]. Others have carried out some form of search but this is unspecified [17], whilst some use trial and error methods [18], [19]. Of those which actually perform a search, some are quadratic [9], [10], whilst other use a linear search with equal number of nodes in the first and second hidden layers [20], [21]. Nonetheless it may be that the authors are basing their searches from past experience. It is impossible to tell.

These choices are summarised in Table I, and the same data represented graphically in Fig. 3. Whatever the underlying reason, or reasoning, it is still interesting that the trend is for an equal number of nodes in the first and second hidden layers, with a slight bias towards more nodes in the first hidden layer.

![Graphical representation of node selections](image)

**Fig. 3.** Graphical representation of node selections.

**TABLE I. TLFN NODE SELECTIONS IN THE LITERATURE**

<table>
<thead>
<tr>
<th>Authors</th>
<th>( n_1 )</th>
<th>( n_2 )</th>
<th>( n_\text{choice} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akciyol and Cinar 2005 [22]</td>
<td>40</td>
<td>40</td>
<td>0.5( n_2 ) + 0</td>
</tr>
<tr>
<td>Hunicz et al. 2002 [16]</td>
<td>13</td>
<td>11</td>
<td>0.5( n_2 ) + 1</td>
</tr>
<tr>
<td>Asik et al. 1997 [15]</td>
<td>8</td>
<td>8</td>
<td>0.5( n_2 ) + 0</td>
</tr>
<tr>
<td>Czarnigowski et al. 2007 [17]</td>
<td>5</td>
<td>4</td>
<td>0.5( n_2 ) + 1</td>
</tr>
<tr>
<td>Deh Kiani et al. 2010 [18]</td>
<td>25</td>
<td>25</td>
<td>0.5( n_2 ) + 0</td>
</tr>
<tr>
<td>Janikiranam et al. 2006 [23]</td>
<td>10</td>
<td>20</td>
<td>0.5( n_2 ) – 5</td>
</tr>
<tr>
<td>Kurniajan et al. 2007 [19]</td>
<td>15</td>
<td>12</td>
<td>0.5( n_2 ) – 1</td>
</tr>
<tr>
<td>Nikzadfar and</td>
<td>10</td>
<td>10</td>
<td>0.5( n_2 ) + 0</td>
</tr>
<tr>
<td>Shamekhi 2014 [24]</td>
<td>10</td>
<td>8</td>
<td>0.5( n_2 ) + 1</td>
</tr>
<tr>
<td>Ozgoren et al. 2013 [25]</td>
<td>11</td>
<td>7</td>
<td>0.5( n_2 ) + 2</td>
</tr>
<tr>
<td>Roy et al. 2014 [10]</td>
<td>8</td>
<td>8</td>
<td>0.5( n_2 ) + 0</td>
</tr>
<tr>
<td>Taghavifar et al. 2014 [9]</td>
<td>19</td>
<td>17</td>
<td>0.5( n_2 ) + 1</td>
</tr>
<tr>
<td>Wu et al. 2004 [20]</td>
<td>8</td>
<td>8</td>
<td>0.5( n_2 ) + 0</td>
</tr>
<tr>
<td>Wu et al. 2005 [21]</td>
<td>10</td>
<td>10</td>
<td>0.5( n_2 ) + 0</td>
</tr>
</tbody>
</table>

**III. EXPERIMENTAL METHOD**

**A. Experimental Environment**

All experiments were carried out using Matlab R2014b with the ‘fitnet’ function available in the neural network toolbox.

**B. Datasets**

The datasets used in these experiments were selected because of their availability in the public domain, and their suitability for a function approximation tasks. They were obtained from the following sources:

- **UCI Machine learning Repository** [26]: Abalone, Airfoil Self-Noise, Concrete Compressive Strength, White Wine.
- **BU Function Approximation Repository** [27]: Kinematics, Mortgage.
- **UP Regression Datasets** [28]: Delta elevators.
- **Matlab**: chemical_dataset, engine_dataset, simplefit_dataset. Further information about these can be found by typing ‘help’ followed by the dataset name at the Matlab command line.

**C. Data Preparation**

In all cases, the data was split into three subsets: Training, Validation, and Test as summarised in Table II. The Validation set was used to stop the training process when the validation error starts to rise, and the Test set was used exclusively as an estimate of the generalisation error. Prior to this, the Engine dataset, which has two inputs and two outputs was reorganised into three inputs and a single output as in [14].

**TABLE II. DATASET SUMMARY**

<table>
<thead>
<tr>
<th>Dataset Name</th>
<th>Inputs</th>
<th>Samples</th>
<th>Training</th>
<th>Val &amp; Test (each)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abalone</td>
<td>8</td>
<td>4177</td>
<td>3341</td>
<td>418</td>
</tr>
<tr>
<td>Airfoil Self-Noise</td>
<td>5</td>
<td>1503</td>
<td>1201</td>
<td>151</td>
</tr>
<tr>
<td>Chemical</td>
<td>8</td>
<td>498</td>
<td>398</td>
<td>50</td>
</tr>
<tr>
<td>Concrete</td>
<td>8</td>
<td>1030</td>
<td>824</td>
<td>103</td>
</tr>
<tr>
<td>Delta Elevators</td>
<td>6</td>
<td>9517</td>
<td>7613</td>
<td>952</td>
</tr>
<tr>
<td>Engine</td>
<td>3</td>
<td>1199</td>
<td>959</td>
<td>120</td>
</tr>
<tr>
<td>Kinematics</td>
<td>8</td>
<td>8292</td>
<td>6652</td>
<td>820</td>
</tr>
<tr>
<td>Mortgage</td>
<td>15</td>
<td>1049</td>
<td>839</td>
<td>105</td>
</tr>
<tr>
<td>Simplefit</td>
<td>1</td>
<td>94</td>
<td>64</td>
<td>15</td>
</tr>
<tr>
<td>White Wine</td>
<td>11</td>
<td>4898</td>
<td>3918</td>
<td>490</td>
</tr>
</tbody>
</table>

For any given dataset, exactly the same subsets were used for every single network created in the experiments (220,000 per dataset and thus 2.2 million in total). By eliminating any bias in the error surface that may have resulted from a different random split for each network, it was ensured that they were all competing on the same playing field. The only random element at play was thus the initial randomisation of the weights. This initial starting point determines which local minimum in the error surface the training might get stuck in and thus has a direct impact on the generalisation error. For complex error surfaces, it is extremely unlikely that the global minimum will be found.

**D. Training Algorithm**

In all cases, data preprocessing was ‘mapminmax’ for both inputs and outputs, the transfer function was ‘tansig’ and the global minimum will be found.

\[
NRMSE_i = \frac{1}{\hat{y}_{\text{max}} - \hat{y}_{\text{min}}} \sqrt{\frac{\sum_{i=1}^{N} (y_i - \hat{y}_i)^2}{N}},
\]

where \( N \) represents the number of samples, \( \hat{y}_i \) is the target value, and \( y_i \) is the actual value.
The Levenberg-Marquardt training algorithm ‘trainlm’ was deployed with early stopping for all experiments, using its default training parameters. Notably: 1000 epochs, training goal of 0, min. gradient of $10^{-3}$, and 6 validation failures.

E. Method

It was found in [14] that the difference in generalisation error between adjacent nodes about the axis of symmetry becomes less significant as $n_h$ is increased. This can be seen quite clearly in Fig. 2, where the 34th isonode contour is almost flat. For this reason it was decided to experiment using only two values of $n_h$, small ($n_h = 8$) and medium ($n_h = 16$) for this study. Referring to Fig. 4, for the 8th isonode ($n_h = 8$), the 7 possible topologies are represented by squares, and the process is as follows:

1. For each topology, find the medium generalisation error from 100 networks resulting in the set, $\varepsilon$
2. The best generalisation error is $\varepsilon_{\text{min}} = \min(\varepsilon)$
3. Evaluate the set of relative errors $\varepsilon_{\text{rel}} = \varepsilon - \varepsilon_{\text{min}}$. This is the raw data for a single run shown in Fig. 5
4. For all topologies such that $\varepsilon_{\text{rel}} < \varepsilon_{\text{th}}$, register a hit in its column. In Fig. 5 this is columns 4, 5 and 6
5. Repeat for 100 runs for each of the 10 datasets
6. The probability that a particular topology will yield a generalisation error less than the threshold is given by $p(\varepsilon_{\text{rel}} < \varepsilon_{\text{th}}) = k / r$, where $k$ is the number of hits for that topology, and $r$ is the total number of runs
7. The results are plotted on a ‘probability map’, where each column has a probability between 0 and 1.

The process is identical for the 16th isonode ($n_h = 16$), but in this case the 15 topologies denoted by circles are considered.

IV. RESULTS

The results are presented as the average and worst case probability across the ten datasets for the 8th and 16th isonodes. The error threshold values ($\varepsilon_{\text{th}}$) were adjusted, in each case, until the peak worst case probability reached 0.5. At these values, even the worst performing dataset has a 50% chance of being less than the error threshold $\varepsilon_{\text{th}}$. This occurred when the value of $\varepsilon_{\text{th}}$ was 0.18% and 0.13% for the 8th and 16th isonodes respectively. In the probability maps shown in Figs. 6 and 7, the x-axis represents the number of nodes in the first hidden layer $n_1$ (the number of nodes in the second hidden layer $n_2 = n_h - n_1$), and the y-axis shows the average and worst case probability that the relative error is less than $\varepsilon_{\text{th}}$, or in other words $p(\varepsilon_{\text{rel}} < \varepsilon_{\text{th}})$. 

Fig. 4. Experimental topologies for 8th and 16th isonodes.

Fig. 5. Raw data for a single run with error threshold $\varepsilon_{\text{th}}$.

For each topology, find the median generalisation error from 100 networks resulting in the set, $\varepsilon$.

The best generalisation error is $\varepsilon_{\text{min}} = \min(\varepsilon)$.

Evaluate the set of relative errors $\varepsilon_{\text{rel}} = \varepsilon - \varepsilon_{\text{min}}$. This is the raw data for a single run shown in Fig. 5.

For all topologies such that $\varepsilon_{\text{rel}} < \varepsilon_{\text{th}}$, register a hit in its column. In Fig. 5 this is columns 4, 5 and 6.

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The probability that a particular topology will yield a generalisation error less than the threshold is given by $p(\varepsilon_{\text{rel}} < \varepsilon_{\text{th}}) = k / r$, where $k$ is the number of hits for that topology, and $r$ is the total number of runs.

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Fig. 5. Raw data for a single run with error threshold $\varepsilon_{\text{th}}$.

Fig. 6. Probability map for the 8th isonode, with $\varepsilon_{\text{th}} = 0.18\%$.

Fig. 7. Probability map for the 16th isonode, with $\varepsilon_{\text{th}} = 0.13\%$.

Fig. 8. Average relative error map.
Both the average and worst case probabilities peak at \( n_1 = 5 \) and \( n_1 = 9 \) for the 8th and 16th isonodes respectively. This is consistent with the previous findings of the Authors [14], which concluded that the best ratio of nodes was given by the heuristic \( n_1 = int(0.5n_h + 1) \). The probability maps of the individual datasets (not shown here), revealed that some were less susceptible to variations in the node ratio than others, about 50% of them having a fairly flat probability map.

Examining the actual generalisation errors relative to the best generaliser (\( \epsilon_{rel} \)), a similar picture emerged. Figs. 8 and 9 show maps of the average and worst case \( \epsilon_{rel} \), respectively, against the number of nodes in the first hidden layer. In these maps, the x-axis represents an offset from 0.5\( n_h \) for ease of comparison – in other words \( n_1 = 0.5n_h + x \). From these maps it can be seen that on average, at \( n_1 = 0.5n_h + 1 \), the relative generalisation error \( \epsilon_{rel} \) is less than 0.1% for both the 8th and 16th isonodes. Furthermore \( \epsilon_{rel} \) is at most 0.15% and 0.23% for the 16th and 8th isonodes respectively.

V. CONCLUSIONS

Training feedforward neural networks using backpropagation algorithms is a very probabilistic affair. The initial random allocation of weights dictate the starting position on the error surface, and this in turn dictates the end point. With a perfect training algorithm, the endpoint would always be the global minimum, in which case it wouldn’t matter what the initial weights were. Unfortunately, backpropagation algorithms are far from perfect and the endpoint will almost certainly be a local minimum. The consequence of this is that the final generalisation error will vary from training session to training session, and that any search for the ‘optimal’ topology is more than likely to yield a different outcome on each occasion. In reality, therefore, there is no such thing as an optimal topology, only a probability that a particular topology will yield the optimal generalisation errors.

The same constraints apply to this study, where it is concluded that although the heuristic \( n_1 = int(0.5n_h + 1) \) does not guarantee that the best generaliser will be found, there is a high probability that it will – at the very least within the context of the datasets tested here. The findings are that using this heuristic, even the worst performing dataset has a probability of 0.5 of finding a network within 0.18% (\( n_h = 8 \)) and 0.13% (\( n_h = 16 \)) of the best generaliser, but that on average these probabilities are much higher at 0.85 and 0.89 respectively. Furthermore, in the worst case, the relative errors are within just 0.23% and 0.15% respectively, but typically less than 0.1% in both cases.

These findings are very significant, as they can be applied to reduce the complexity of a quadratic (raster) search through \( n_1, n_2 \) to a linear search through \( n_h \) with very little penalty in the generalisation error. Such a search would follow the staircase trajectory shown in Fig. 10. Alternatively, only even values of \( n_h \) could be considered, in which case the trajectory would be \( n_2 = n_1 - 2 \), shown as the shaded node allocations in Fig. 10. This latter trajectory would reduce a search through \( n^2 \) nodes to one through \( n - 2 \) nodes. This is equivalent to a reduction in complexity from \( O(n^2) \) to \( O(n) \).

A further application of these findings is in providing an isonode topology ‘transform’ or ‘mapping’ between single and double hidden layer networks. In this context, given a particular number of hidden nodes, \( n_h \), equation (10) is used to map a single hidden layer topology on to an ‘equivalent’ two hidden layer topology. Fig. 11 shows this topology mapping applied to the Airfoil Self-Noise dataset. It compares the average generalisation error of SLFN and TLFN networks with an equal number of hidden nodes. In this case, node for node, two hidden layers clearly perform better than a single hidden layer beyond \( n_h = 5 \).

VI. FURTHER WORK

It is possible that the complexity could be further reduced to \( O(\log_2(n)) \) using the predictive method developed by the Authors in [29]. There are two possible approaches:

1. Use the predictive method ‘as is’ to find the optimal SLFN in \( O(\log_2(n)) \). Subsequently, apply an isonode topology mapping to generate an equivalent TLFN.

2. Alternatively, the predictive method could be adapted to use the relationship in equation (10) directly. It was tentatively shown in [14] that the isonode topology mapping could well be suitable for use in such a predictive approach.
It is believed at this stage that the former approach would probably be the faster of the two. However, it is suspected that the latter would yield better results. Pending the results of further experimentation, either approach would reduce the complexity of finding TLFNs from $O(n^2)$ to $O(\log(n))$. This significant reduction in complexity could potentially find ‘optimal’ TLFNs in a matter of minutes rather than days.

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REFERENCES


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