

The Incredible Shrinking Neural Network: Pruning to Operate in Constrained Memory Environments

Abstract

We propose and investigate several methods for pruning artificial neural networks to operate in constrained memory environments such as mobile or embedded devices. Unlike traditional methods that compress networks by pruning their weights, we focus on pruning entire neurons, thereby eliminating all the incoming and outgoing weights from a pruned node resulting in more memory saved. All the proposed methods prune neurons based off a "ranked list" of the neurons' contribution to overall network performance. Each method proposed uses a different way of creating this ranked list. We evaluate these methods and look into their strengths and limitations. We also evaluate 2 different algorithms to perform pruning on these created ranked lists and compare their performance. In doing so, we also get to look into the correlation between neurons in a trained network and their learning representations, using which we try to answer an age old question: are all the neurons in a network the same? We argue that within this answer lies the perfect method of pruning that allows for the optimal trade-off in network size versus accuracy in order to operate within the memory constraints of a particular device or application environment.

Keywords: pruning, neural network compression, pruning neurons, learning representation

1. Introduction

Neural network pruning algorithms were first popularized by [Sietsma and Dow \(1988\)](#) as a mechanism to determine the proper size of a network required to solve a particular problem. To this day, network design and optimal pruning remain inherently difficult tasks. For problems which cannot be solved using linear threshold units alone, [Baum and Haussler \(1989\)](#) demonstrate there is no way to precisely determine the appropriate size of a neural network a priori given any random set of training instances.

Pruning algorithms, as comprehensively surveyed by [Reed \(1993\)](#), are a useful set of heuristics designed to identify and remove network parameters which do not contribute significantly to the output of the network and potentially inhibit generalization performance. At the time of Reed's writing, reducing network size was also a practical concern, as smaller networks are preferable in situations where computational resources are scarce. In this paper we are particularly concerned with application domains in which space is limited and network size constraints must be imposed with minimal impact on performance.

Neural network over-fitting is fundamentally a problem arising from the use of too many free parameters. Regardless of the number of weights used in a given network, as [Segee and Carter \(1991\)](#) assert, the representation of a learned function approximation is almost never evenly distributed over the hidden units, and the removal of any single hidden unit at random can actually result in a total network fault. [Mozer and Smolensky \(1989b\)](#) suggest that only a subset of the hidden units in a neural network actually latch on to the invariant

or generalizing properties of the training inputs, and the rest learn to either mutually cancel each other out or begin over-fitting to the noise in the data. Determining which elements are unnecessary and removing them outright is therefore a well-founded approach to improving network generalization, and simultaneously provides a way to reduce their size in memory. The generalization performance of neural networks has been well studied, and apart from pruning algorithms many heuristics have been used to avoid overfitting, such as dropout (Srivastava et al. (2014)), maxout (Goodfellow et al. (2013)), and cascade correlation (Fahlman and Lebiere (1989)), among others. However, these algorithms do not explicitly prioritize the reduction of network memory footprint as a part of their optimization criteria per se, (although in the opinion of the authors, Fahlman’s cascade correlation architecture holds great promise in this regard.) Computer memory size and processing capabilities have improved so much since the introduction of pruning algorithms in the late 1980s that space complexity has become a relatively negligible concern. The proliferation of cloud-based computing services has furthermore enabled mobile and embedded devices to leverage the power of massive data and computing centres remotely. In this domain, however, it is also reasonable to suggest that certain performance-critical applications running on low-resource devices could benefit from the ability to use neural networks locally.

At present there are few (if any) mechanisms specifically designed to shrink neural networks down in order to meet an externally imposed constraint on byte-size in memory. Without explicitly removing parameters from the network, one could use weight quantization to reduce the number of bytes used to represent each weight parameter, as investigated by Balzer et al. (1991), Dundar and Rose (1994), and Hoehfeld and Fahlman (1992). Of course, this method can only reduce the size of the network by a factor proportional to the byte-size reduction of each weight parameter.

Recently, some other network compression methods have been proposed. One such method which has recently gained popularity (Prabhavalkar et al. (2016)) uses the singular values of a trained weight matrix as basis vectors from which to derive a compressed hidden layer. Some other recent works like Oland and Raj (2015) have tried successfully to achieve compression through weight quantization followed by encoding while some others like Han et al. (2016) have tried to build on top of that by adding weight-pruning as a preceding step to quantization and encoding.

If we wanted to continually shrink a network to its absolute minimal size in an optimal manner, we might accomplish this using any number of off-the-shelf pruning algorithms, such as Skeletonization (Mozier and Smolensky (1989a)), Optimal Brain Damage (LeCun et al. (1989)), or later variants such as Optimal Brain Surgeon (Hassibi and Stork (1993)). In fact, we borrow much of our inspiration from these antecedent algorithms, with one major variation.

The aforementioned strategies all focus on the targeting and removal of *weight* parameters. Scoring and ranking individual weight parameters in a large network computationally expensive, and generally speaking the removal of a single weight from a large network is a drop in the bucket in terms of reducing a network’s core memory footprint. We therefore train our sights on the ranking and removal of entire neurons along with their associated weight parameters. We argue that this is more efficient computationally as well as practically in terms of quickly reaching a target reduction in memory size. Our approach also attacks the angle of giving downstream applications a realistic expectation of the minimal increase in

error resulting from the removal of a specified percentage of neurons. Such trade-offs are unavoidable, but performance impacts can be limited if a principled approach is used to find candidate neurons for removal.

2. Methodology

The general approach taken to prune an optimally trained neural network here is to create a ranked list of all the neurons in the network based off of one of the 3 proposed ranking criteria: Brute Force approximation (which we use as our ground truth), linear approximation and quadratic approximation of the neuron’s impact on the overall performance of the network. We then test the effects of removing neurons on the accuracy and error of the network.

2.1. Brute Force Removal Approach

This is perhaps the most naive yet the most accurate method for pruning the network. It is also the slowest and hence unusable on large-scale neural networks with thousands of neurons. The idea is to manually check the effect of every single neuron on the output. This is done by running a forward propagation on the validation set K times (where K is the total number of neurons in the network), turning off exactly one neuron each time (keeping all other neurons active) and noting down the change in error. Turning a neuron off can be achieved by simply setting its output to 0. This results in all the outgoing weights from that neuron being turned off. This change in error is then used to generate the ranked list.

2.2. Taylor Series Representation of Error

Let us denote the total error from the optimally trained neural network for any given validation dataset by E . E can be seen as a function of O , where O is the output of any general neuron in the network. This error can be approximated at a particular neuron’s output (say O_k) by using the 2nd order Taylor Series as,

$$\hat{E}(O) \approx E(O_k) + (O - O_k) \cdot \left. \frac{\partial E}{\partial O} \right|_{O_k} + 0.5 \cdot (O - O_k)^2 \cdot \left. \frac{\partial^2 E}{\partial O^2} \right|_{O_k}, \quad (1)$$

When a neuron is pruned, its output O becomes 0. From equation 1, the contribution $E(0)$ of this neuron, then becomes:

$$\hat{E}(0) \approx E(O_k) - O_k \cdot \left. \frac{\partial E}{\partial O} \right|_{O_k} + 0.5 \cdot O_k^2 \cdot \left. \frac{\partial^2 E}{\partial O^2} \right|_{O_k} \quad (2)$$

Replacing O by O_k in equation 1 shows us that the error is approximated perfectly by equation 1 at O_k . Using this and equation 2 we get:

$$\Delta E_k = \hat{E}(0) - \hat{E}(O_k) = -O_k \cdot \left. \frac{\partial E}{\partial O} \right|_{O_k} + 0.5 \cdot O_k^2 \cdot \left. \frac{\partial^2 E}{\partial O^2} \right|_{O_k}, \quad (3)$$

where ΔE_k is the change in the total error of the network when exactly one neuron (k) is turned off.

2.3. Linear Approximation Approach

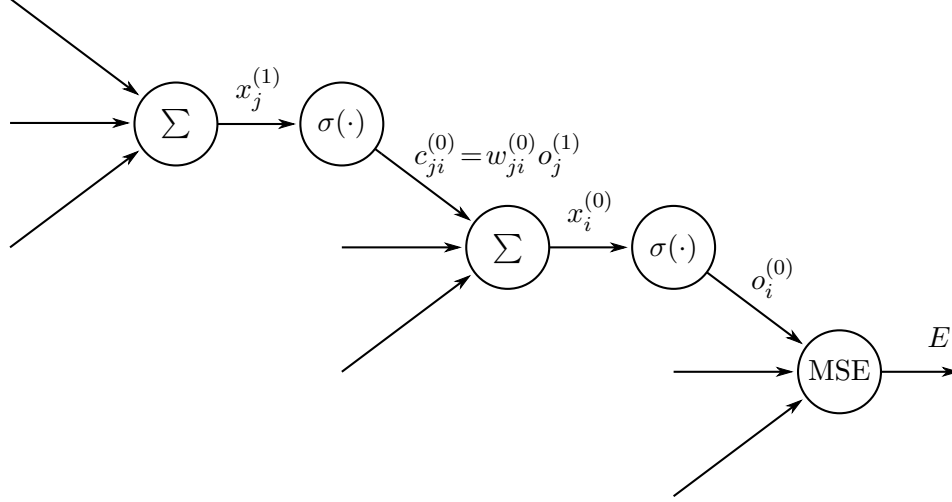


Figure 1: A computational graph of a simple feed-forward network illustrating the naming of different variables, where $\sigma(\cdot)$ is the nonlinearity, MSE is the mean-squared error cost function and E is the overall loss.

We define the following network terminology here which will be used in this section and all subsequent sections unless stated otherwise. Figure 1 can be used as a reference to the terminology defined here:

$$E = \frac{1}{2} \sum_i (o_i^{(0)} - t_i)^2 \quad o_i^{(m)} = \sigma(x_i^{(m)}) \quad x_i^{(m)} = \sum_j w_{ji}^{(m)} o_j^{(m+1)} \quad c_{ji}^{(m)} = w_{ji}^{(m)} o_j^{(m+1)} \quad (4)$$

Superscripts represent the index of the layer of the network in question, with 0 representing the output layer. E is the squared-error network cost function. $o_i^{(m)}$ is the i th output in layer m generated by the activation function σ , which in this paper is the standard logistic sigmoid. $x_i^{(m)}$ is the weighted sum of inputs to the i th neuron in the m th layer, and $c_{ji}^{(m)}$ is the contribution of the j th neuron in the $(m+1)$ th layer to the input of the i th neuron in the m th layer. $w_{ji}^{(m)}$ is the weight between the j th neuron in the $(m+1)$ th layer and the i th neuron in the m th layer.

We can use equation 3 to get the linear error approximation of the change in error due to the k th neuron being turned off and represent it as ΔE_k^1 as follows:

$$\Delta E_k^1 = -o_k \cdot \left. \frac{\partial E}{\partial o_j^{(m+1)}} \right|_{o_k} \quad (5)$$

The derivative term above is the first-order gradient which represents the change in error with respect to the output of a given neuron o_j in the $(m+1)$ th layer. This term can be collected during back-propagation. The derivative term above can be calculated as follows:

$$\frac{\partial E}{\partial o_j^{(m+1)}} = \sum_i \frac{\partial E}{\partial x_i^{(m)}} \cdot w_{ji}^{(m)} \quad (6)$$

2.4. Quadratic Approximation Approach

As seen in equation 3, ΔE_k which can now be represented as ΔE_k^2 is the quadratic approximation of the change in error due to the k th neuron being turned off. The quadratic term in equation 3 requires some discussion which we provide here.

Let us reproduce equation 3 in our new terminology here:

$$\Delta E_k^2 = -o_k \cdot \left. \frac{\partial E}{\partial o_j^{(m+1)}} \right|_{o_k} + 0.5 \cdot o_k^2 \cdot \left. \frac{\partial^2 E}{\partial o_j^{(m+1)^2}} \right|_{o_k} \quad (7)$$

This equation involves the second-order gradient which represents the second-order change in error with respect to the output of a given neuron o_j in the $(m+1)$ th layer. This term can be generated by performing back-propagation using second derivatives, a full discussion of which is out of the scope of this paper but some useful results are quoted here. The second-order derivative term can be represented as:

$$\frac{\partial^2 E}{\partial o_j^{(m+1)^2}} = \sum_i \frac{\partial^2 E}{\partial c_{ji}^{(m)^2}} \left(w_{ji}^{(m)} \right)^2 \quad (8)$$

Here, $c_{ji}^{(m)}$ is one of the component terms of $x_i^{(m)}$, as follows from the equations in 4. Hence, it can be easily proved that:

$$\frac{\partial^2 E}{\partial c_{ji}^{(m)^2}} = \frac{\partial^2 E}{\partial x_i^{(m)^2}} \quad (9)$$

Now, the value of $x_i^{(m)}$ can again be easily calculated through the steps of the second-order back-propagation using Chain Rule.

$$\frac{\partial^2 E}{\partial x_i^{(m)^2}} = \frac{\partial^2 E}{\partial o_i^{(m)^2}} \left(\sigma' \left(x_i^{(m)} \right) \right)^2 + \frac{\partial E}{\partial o_i^{(m)}} \sigma'' \left(x_i^{(m)} \right) \quad (10)$$

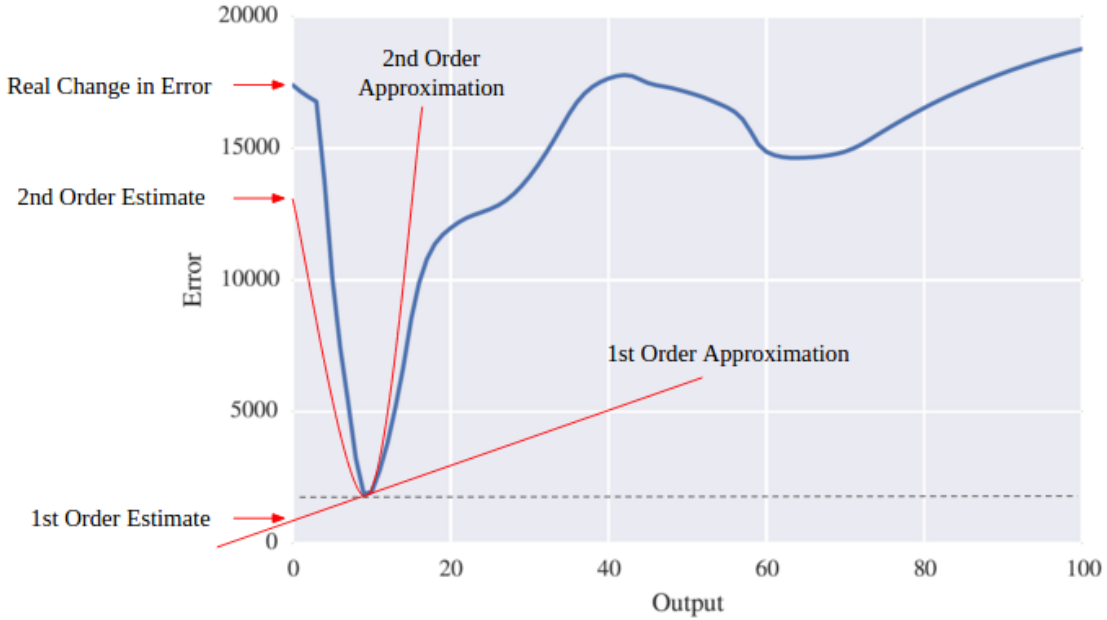


Figure 2: The intuition behind neuron pruning decision.

2.5. Proposed Pruning Algorithm

Figure 2 shows a random error function plotted against the output of any given neuron. Note that this figure is for illustration purposes only. The error function is minimized at a particular value of the neuron output as can be seen in the figure. The process of training a neural network is essentially the process of finding these minimizing output values for all the neurons in the network. Pruning this particular neuron (which translates to getting a zero output from it) will result in a change in the total overall error. This change in error is represented by distance between the original minimum error (shown by the dashed line) and the top red arrow. This neuron is clearly a bad candidate for removal since removing it will result in a huge error increase.

The straight red line in the figure represents the first-order approximation of the error using Taylor Series as described before while the parabola represents a second-order approximation. It can be clearly seen that the second-order approximation is a much better estimate of the change in error.

One thing to note here is that it is possible in some cases that there is some thresholding required when trying to approximate the error using the 2nd order Taylor Series expansion. These cases might arise when the parabolic approximation undergoes a steep slope change. To take into account such cases, mean and median thresholding were employed, where any change above a certain threshold was assigned a mean or median value respectively.

Two pruning algorithms are proposed here. They are different in the way the neurons are ranked but both of them use ΔE_k , the approximation of the change in error as the basis for the ranking. ΔE_k can be calculated using the Brute Force method, or one of the two Taylor Series approximations discussed previously.

The first step in both the algorithms is to decide a stopping criterion. This can vary depending on the application but some intuitive stopping criteria can be: maximum number of neurons to remove, percentage scaling needed, maximum allowable accuracy drop etc.

2.5.1. ALGORITHM I: SINGLE OVERALL RANKING

The complete algorithm is shown in Algorithm 1. The idea here is to generate a single ranked list based on the values of ΔE_k . This involves a single pass of second-order back-propagation (without weight updates) to collect the gradients for each neuron. The neurons from this rank-list (with the lowest values of ΔE_k) are then pruned according to the stopping criterion decided. We note here that this algorithm is intentionally naive and is used for comparison only.

Data: optimally trained network, training set

Result: A pruned network

initialize and define stopping criterion

perform forward propagation over the training set

perform second-order back-propagation without updating weights and collect linear and quadratic gradients

rank the remaining neurons based on ΔE_k

while *stopping criterion is not met* **do**

 remove the last ranked neuron

end

Algorithm 1: Single Overall Ranking

2.5.2. ALGORITHM II: ITERATIVE RE-RANKING

In this greedy variation of the algorithm (Algorithm 2), after each neuron removal, the remaining network undergoes a single forward and backward pass of second-order back-propagation (without weight updates) and the rank list is formed again. Hence, each removal involves a new pass through the network. This method is computationally more expensive but takes into account the dependencies the neurons might have on one another which would lead to a change in error contribution every time a dependent neuron is removed.

Data: optimally trained network, training set

Result: A pruned network

initialize and define stopping criterion

while *stopping criterion is not met* **do**

 perform forward propagation over the training set

 perform second-order back-propagation without updating weights and collect linear and quadratic gradients

 rank the remaining neurons based on ΔE_k

 remove the worst neuron based on the ranking

end

Algorithm 2: Iterative Re-Ranking

3. Experimental Results

For all the results presented in this section, the MNIST database of Handwritten Digits by [LeCun and Cortes \(2010\)](#) was used. It is worth noting here that since only a subset of the entire dataset was used in order to speed up experiments, the starting test accuracies reported here might differ from those reported by LeCun. This does not impact the presented results, since it’s the change in the error and accuracy of the network post pruning which is of significance here and not the starting accuracy itself.

3.1. Pruning a 1-Layer Network

The network architecture in this case consisted of 1 layer, 100 neurons, 10 outputs, logistic sigmoid activations, and a starting test accuracy of 0.998.

3.1.1. SINGLE OVERALL RANKING ALGORITHM

We first present the results for a single-layer neural network in Figure 3, using the Single Overall algorithm (Algorithm 1) as proposed in the Methodology section. (We again note that this algorithm is intentionally naive and is used for comparison only. Its performance should be expected to be poor.) After training, each neuron is assigned its permanent ranking based on the three criteria discussed previously: A brute force “ground truth” ranking, and two approximations of this ranking using first and second order Taylor estimations of the change in network output error resulting from the removal of each neuron.

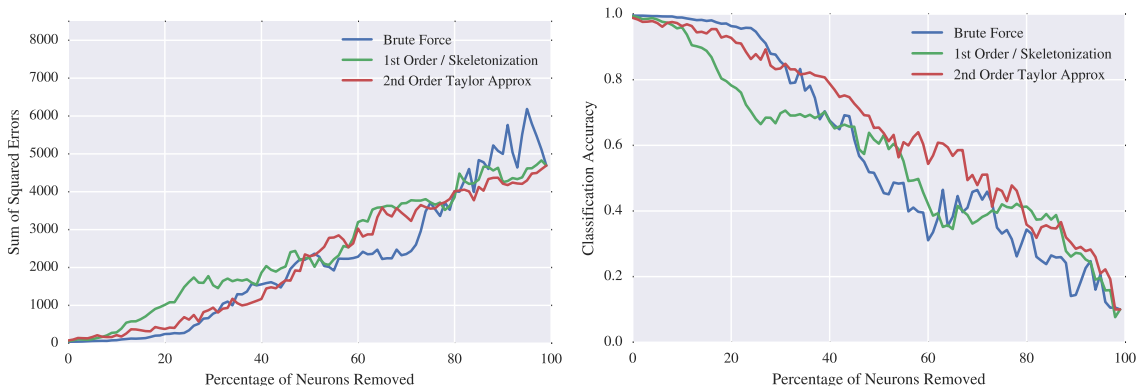


Figure 3: Degradation in squared error (left) and classification accuracy (right) after pruning a single-layer network using The Single Overall Ranking algorithm (**Network:** 1 layer, 100 neurons, 10 outputs, logistic sigmoid activation, starting test accuracy: 0.998)

An interesting observation here is that with only a single layer, no criteria for ranking the neurons in the network (brute force or the two Taylor Series variants) using Algorithm 1 emerges superior, indicating that the 1st and 2nd order Taylor Series methods are actually reasonable approximations of the brute force method under certain conditions. Of course, this method is still quite bad in terms of the rate of degradation of the classification accuracy

and in practice we would likely follow Algorithm 2 which takes into account Mozer and Smolensky (1989a)’s observations stated in the Related Work section. The purpose of the present investigation, however, is to demonstrate how much of a trained network can be theoretically removed without altering the network’s learned parameters in any way.

3.1.2. ITERATIVE RE-RANKING ALGORITHM

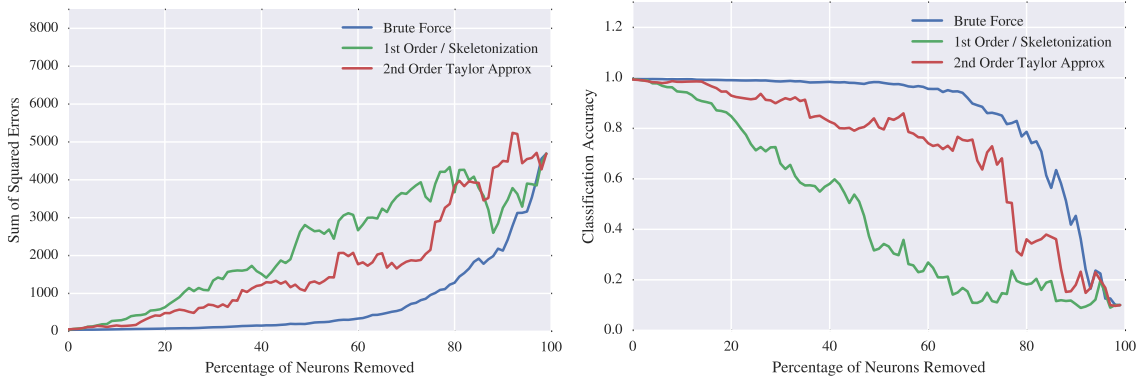


Figure 4: Degradation in squared error (left) and classification accuracy (right) after pruning a single-layer network the Iterative Re-ranking algorithm (**Network:** 1 layer, 100 neurons, 10 outputs, logistic sigmoid activation, starting test accuracy: 0.998)

In Figure 4 we present our results using Algorithm 2 (The Iterative Re-Ranking Algorithm) in which all remaining neurons are re-ranked after each successive neuron is switched off. We compute the same brute force rankings and Taylor series approximations of error deltas over the remaining active neurons in the network after each pruning decision. This is intended to account for the effects of cancelling interactions between neurons.

There are 2 key observations here. Using the Brute Force ranking criteria, almost 60% of the neurons in the network can be pruned away without any major loss in performance. The other noteworthy observation here is that the 2nd order Taylor Series approximation of the error performs consistently better than its 1st order version.

3.1.3. VISUALIZATION OF ERROR SURFACE & PRUNING DECISIONS

As explained in the Methodology section, these graphs are a visualization of the error surface of the network output with respect to the neurons chosen for removal using each of the 3 ranking criteria, represented in intervals of 10 neurons. In each graph, the error surface of the network output is displayed in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal. We create these plots during the pruning exercise by picking a neuron to switch off, and then multiplying its output by a scalar gain value α which is adjusted from 0.0 to 10.0 with a step size of 0.001. When the value of α is 1.0, this represents the unperturbed neuron output learned during training. Between 0.0 and 1.0, we are graphing the literal effect of turning the neuron off ($\alpha = 0$), and when

$\alpha > 1.0$ we are simulating a boosting of the neuron's influence in the network, i.e. inflating the value of its outgoing weight parameters.

We graph the effect of boosting the neuron's output to demonstrate that for certain neurons in the network, even doubling, tripling, or quadrupling the scalar output of the neuron has no effect on the overall error of the network, indicating the remarkable degree to which the

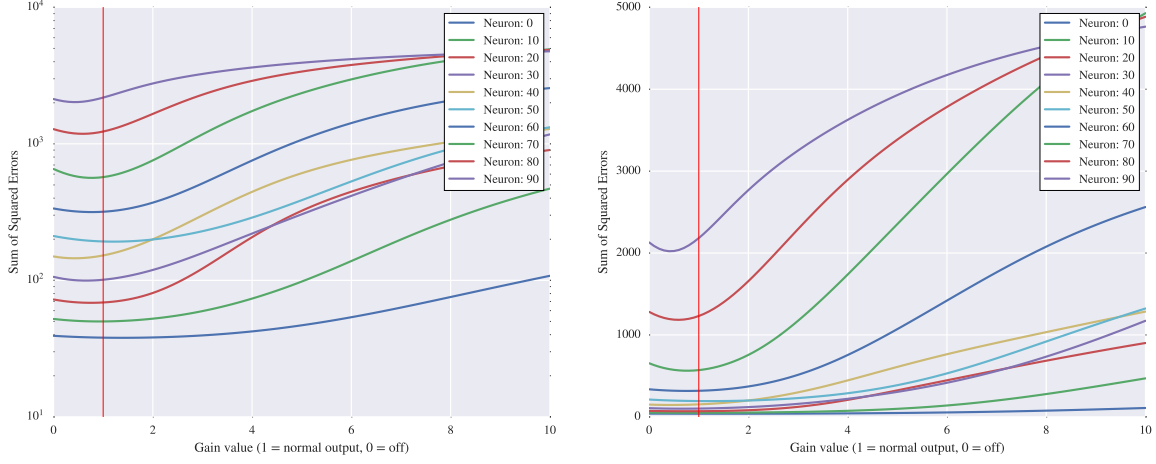


Figure 5: Error surface of the network output in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal using the brute force criterion; (**Network:** 1 layer, 100 neurons, 10 outputs, logistic sigmoid activation, starting test accuracy: 0.998)

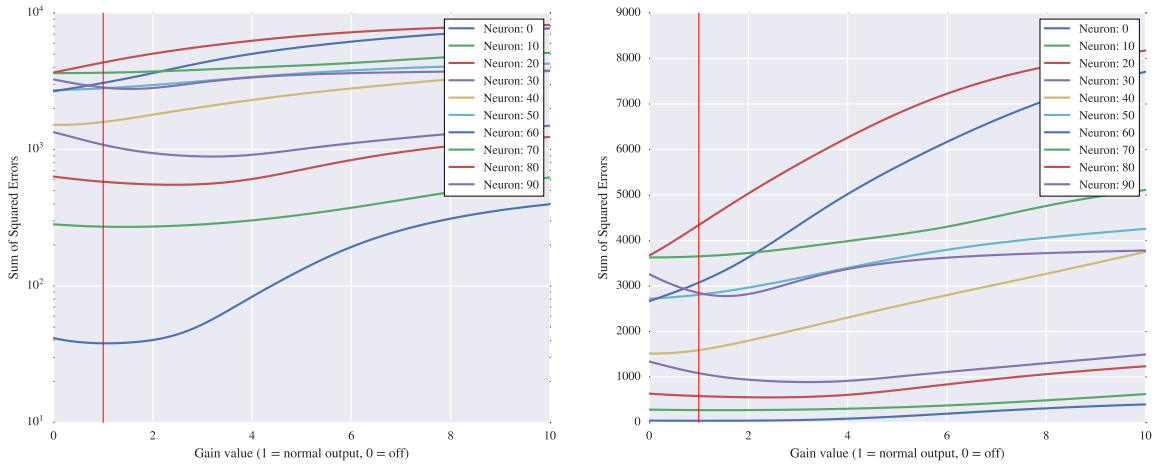


Figure 6: Error surface of the network output in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal using the 1st order Taylor Series error approximation criterion; (**Network:** 1 layer, 100 neurons, 10 outputs, logistic sigmoid activation, starting test accuracy: 0.998)

network has learned to ignore the value of certain parameters. In other cases, we can get a sense of the sensitivity of the network's output to the value of a given neuron when the curve rises steeply after the red 1.0 line. This indicates that the learned value of the parameters emanating from a given neuron are relatively important, and this is why we should ideally see sharper upticks in the curves for the later-removed neurons in the network, that is, when the neurons crucial to the learning representation start to be picked off. Some very interesting observations can be made in each of these graphs.

Brute Force Criterion

Notice how low to the floor and flat most of these curves are. It's only until the 90th removed neuron that we see a higher curve with a more convex shape (clearly a more influential piece of the network). This again validates the fact that a good majority of the neurons in a network do not contribute to the overall performance.

1st Order Approximation Criterion

It can be seen that most choices seem to have flat or negatively sloped curves, indicating that the first order approximation seems to be pretty good, but examining the brute force choices shows they could be better.

2nd Order Approximation Criterion

This method looks much more similar to the Brute Force method choices, though clearly not as good (they're more spread out). Notice the difference in convexity between the 2nd and 1st order method choices. It's clear that the first order method is fitting a line and the 2nd order method is fitting a parabola in their approximation.

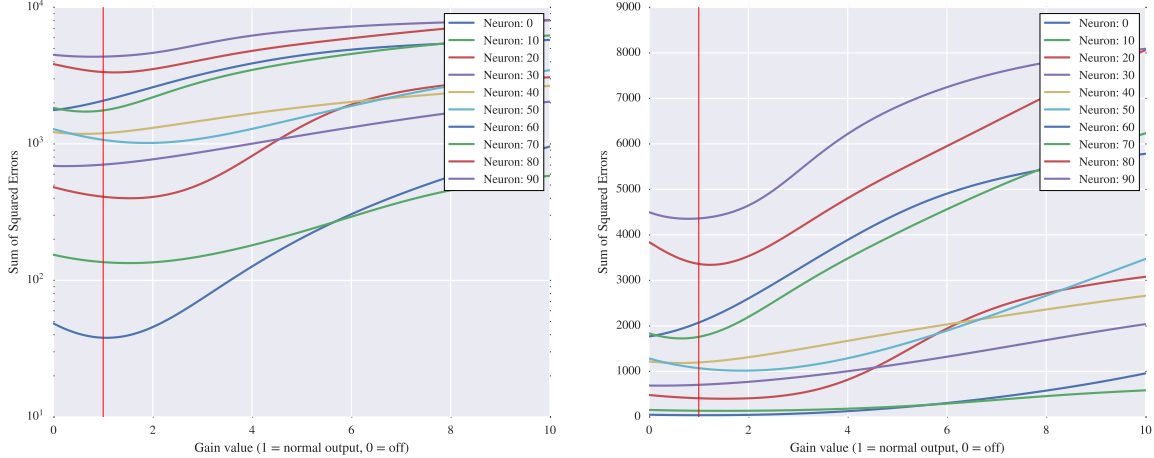


Figure 7: Error surface of the network output in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal using the 2nd order Taylor Series error approximation criterion; (**Network:** 1 layer, 100 neurons, 10 outputs, logistic sigmoid activation, starting test accuracy: 0.998)

3.2. Pruning A 2-Layer Network

The network architecture in this case consisted of 2 layers, 50 neurons per layer, 10 outputs, logistic sigmoid activations, and a starting test accuracy of 1.000.

3.2.1. SINGLE OVERALL RANKING ALGORITHM

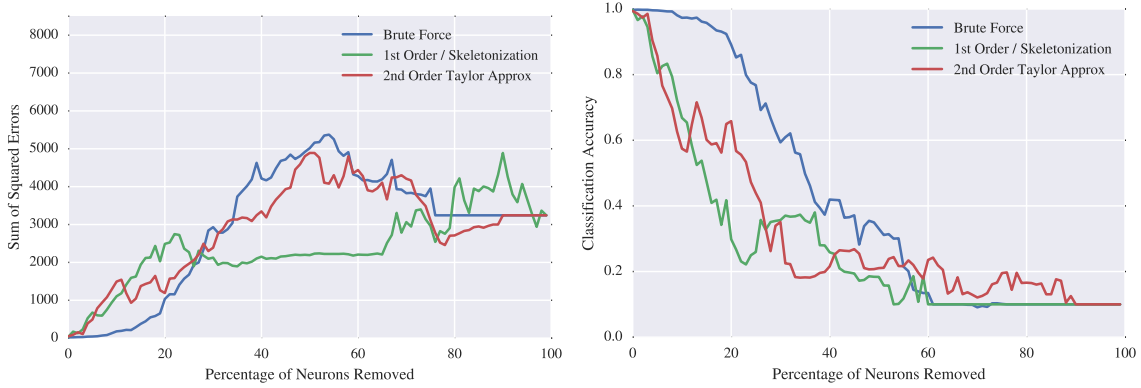


Figure 8: Degradation in squared error (left) and classification accuracy (right) after pruning a 2-layer network using the Single Overall Ranking algorithm; (**Network:** 2 layers, 50 neurons/layer, 10 outputs, logistic sigmoid activation, starting test accuracy: 1.000)

Figure 8 shows the pruning results for Algorithm 1 on a 2-layer network. The ranking procedure is identical to the one used to generate Figure 3. (We again note that this algorithm is intentionally naive and is used for comparison only. Its performance should be expected to be poor.)

Unsurprisingly, a 2-layer network is harder to prune because a single overall ranking will never capture the interdependencies between neurons in different layers. It makes sense that this is much worse than the performance on the 1-layer network, even if this method is already known to be bad, and we’d likely never use it in practice.

3.2.2. ITERATIVE RE-RANKING ALGORITHM

Figure 9 shows the results from using Algorithm 2 on a 2-layer network. We compute the same brute force rankings and Taylor series approximations of error deltas over the remaining active neurons in the network after each pruning decision used to generate Figure 4. Again, this is intended to account for the effects of cancelling interactions between neurons.

It is clear that it becomes harder to remove neurons 1-by-1 with a deeper network (which makes sense because the neurons have more interdependencies in a deeper network), but we see an overall better performance with 2nd order method vs. 1st order, except for the first 20% of the neurons (but this doesn’t seem to make much difference for classification accuracy.)

Perhaps a more important observation here is that even with a more complex network, it is possible to remove up to 40% of the neurons with no major loss in performance which is clearly illustrated by the brute force curve. This shows the clear potential of an ideal pruning technique and also shows how inconsistent 1st and 2nd order Taylor Series approximations of the error can be as ranking criteria.

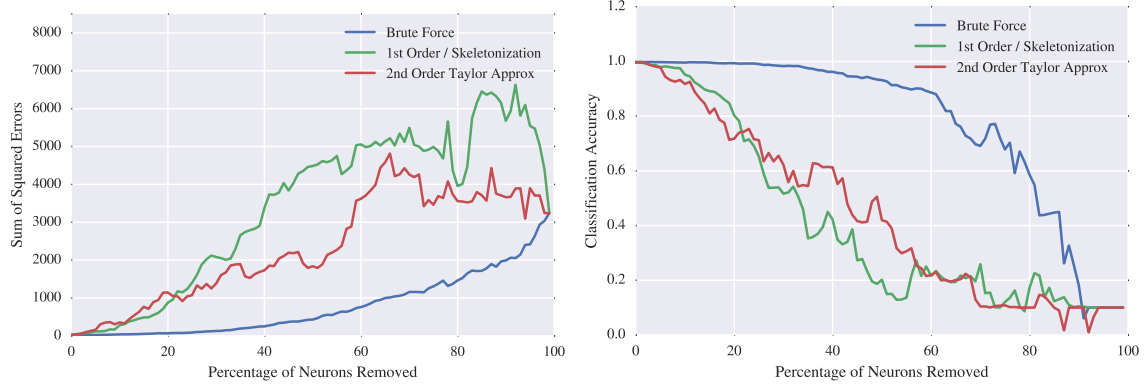


Figure 9: Degradation in squared error (left) and classification accuracy (right) after pruning a 2-layer network using the Iterative Re-ranking algorithm; (**Network:** 2 layers, 50 neurons/layer, 10 outputs, logistic sigmoid activation, starting test accuracy: 1.000)

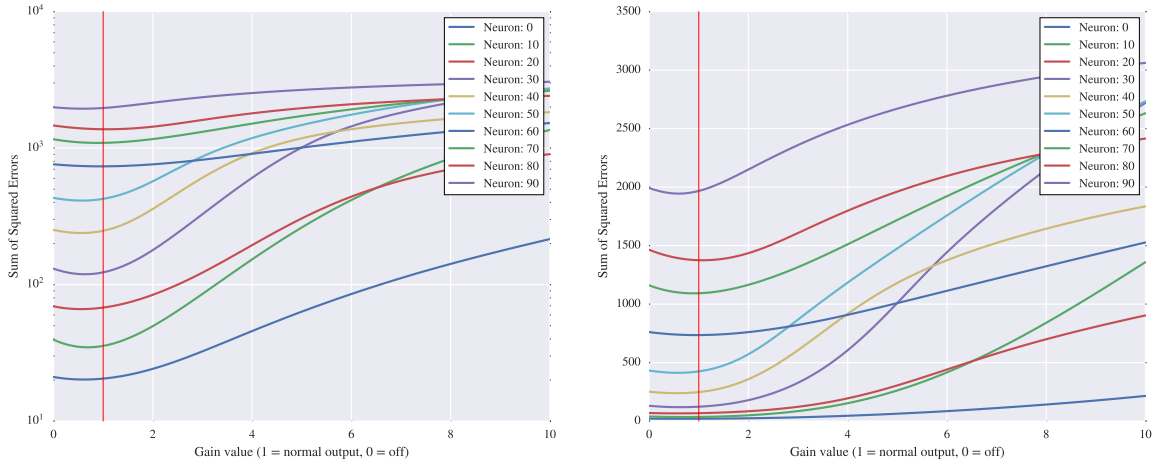


Figure 10: Error surface of the network output in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal using the brute force criterion; (**Network:** 2 layers, 50 neurons/layer, 10 outputs, logistic sigmoid activation, starting test accuracy: 1.000)

3.2.3. VISUALIZATION OF ERROR SURFACE & PRUNING DECISIONS

As seen in the case of a single layered network, these graphs are a visualization the error surface of the network output with respect to the neurons chosen for removal using each algorithm, represented in intervals of 10 neurons.

Brute Force Criterion

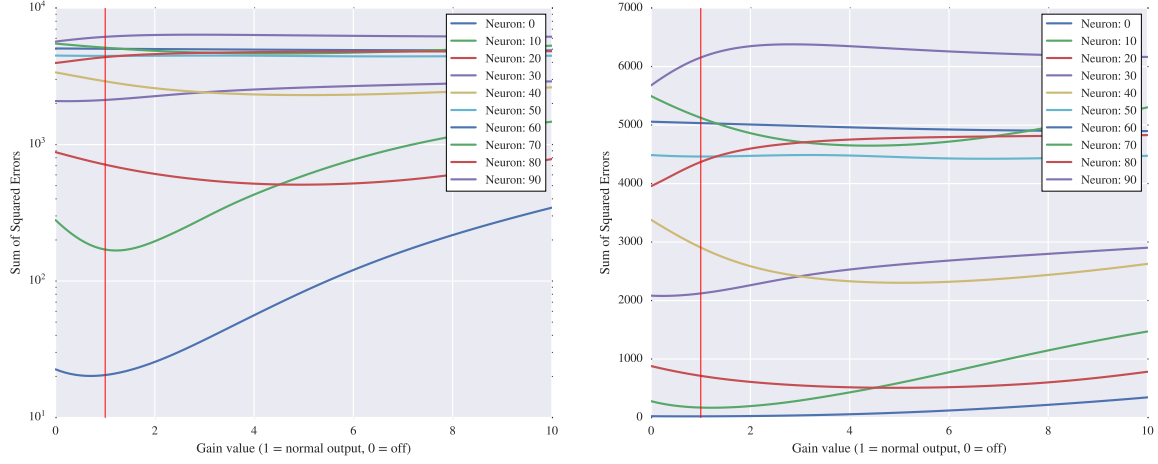


Figure 11: Error surface of the network output in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal using the 1st order Taylor Series error approximation criterion; (**Network:** 2 layers, 50 neurons/layer, 10 outputs, logistic sigmoid activation, starting test accuracy: 1.000)

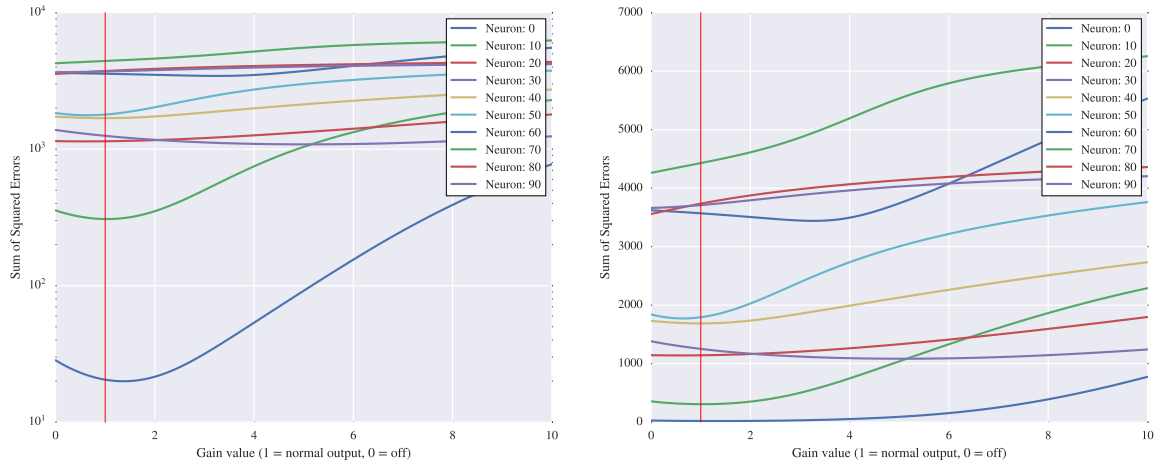


Figure 12: Error surface of the network output in log space (left) and in real space (right) with respect to each candidate neuron chosen for removal using the 2nd order Taylor Series error approximation criterion; (**Network:** 2 layers, 50 neurons/layer, 10 outputs, logistic sigmoid activation, starting test accuracy: 1.000)

It is clear why these neurons got chosen, their graphs clearly show little change when neuron is removed, are mostly near the floor, and show convex behaviour of error surface, which argues for the rationalization of using 2nd order methods to estimate difference in error when they are turned off.

1st Order Approximation Criterion

Drawing a flat line at the point of each neurons intersection with the red vertical line (no change in gain) shows that the 1st derivative method is actually accurate for estimation of change in error in these cases, but still ultimately leads to poor decisions.

2nd Order Approximation Criterion

Clearly these neurons are not overtly poor candidates for removal (error doesn't change much between 1.0 & zero-crossing left-hand-side), but could be better (as described above in the Brute Force Criterion discussion).

4. Conclusions & Future Work

Pruning neurons (instead of pruning individual weights) in a pre-trained neural network without seeing a major loss in performance is not only possible but also enables compressing networks to 40-80% of their original size, which is of great importance in constrained memory environments like embedded devices. This fact is established through the experiments using the brute force criterion, which if made computationally viable (through parallelization) or approximated more efficiently than the Taylor Series based methods discussed in this paper, can prove to be a useful compression tool. It would also be interesting to see how these methods perform on deeper networks and on some other popular and real world datasets. The experiments on the visualization of error surfaces and pruning decisions concretely establish the fact that not all neurons in a network contribute to its performance in the same way. This confirms the idea put forth by [Mozer and Smolensky \(1989b\)](#) that learning representation is not distributed uniformly across neurons. Neural networks use a few neurons to learn the function approximation, and the remaining neurons cooperate to cancel out each other's effects. This is also a strong indication of the fact that once training is done, bigger networks do not hold an advantage over smaller ones, which is similar to the idea put forth by [Hinton et al. \(2015\)](#) in their work on ensemble learning techniques.

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