Applied Deep Learning - Reviewed Notes

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1 Introduction

Definition 1.1 - Deep Representation Learning

Representation Learning is a set of techniques in machine learning where a system can automatically learn representations needed for feature detection from the raw data without the need for hand-designed feature descriptions. Deep Representation Learning is then learning to classify using this feature detection.

Remark 1.1 - Biological Inspiration

In the natural world *Neurons* are the basic working units of the brain. *Neurons* can be split into three main areas

- i). Dendrites Receives inputs from other neurons.
- ii). Axon Carries information.
- iii). Axon Terminals & Synapses = Send information to other neurons.

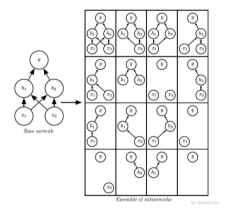
Artificial Neural Networks seek to mimic this structure.

Definition 1.2 - Neuro-Plasticity

Neuro-Plasticity is the ability of a neural system to adapt its structure to accommodate new information (i.e. Learn). This can take several forms including growth & function changes.

Proposition 1.1 - Neural Networks as an Ensemble of Sub-Networks

A Neural Network can be considered to represent many sub-networks. These sub-networks are switched between depending on which components are picked and how they are defined.

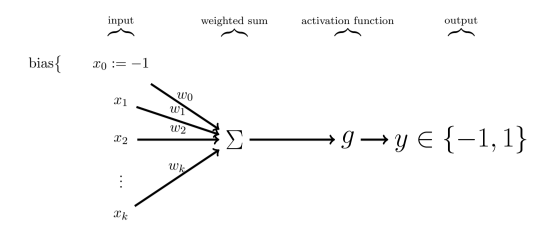


2 Structures

2.1 Perceptron

Definition 2.1 - Perceptron

A *Perceptron* is an algorithm for supervised learning of a binary classifier. A *Perceptron* defines a hyperplane which acts as a decision boundary which linearly separates the input-state space. These two regions correspond to the two-classes. A perceptron has the following structure.



 x_0 is the bias element. It is always set to -1 in the input and the actual value is defined by its weight w_0 .

 $\mathbf{x} = (x_0, \dots, x_k)$ is the input. Note that x_0 is the bias and (x_1, \dots, x_k) are the observed inputs.

 $\mathbf{w} = (w_0, \dots, w_k)$ is the weights assigned to each input (including the bias).

$$\Sigma$$
 is the weighted sum of the bias & inputs. $\Sigma := \sum_{i=0}^k w_i x_i = \boldsymbol{w}^T \boldsymbol{x}$

- g is the Activation Function which maps the result from the weighted sum Σ to $\{-1,1\}$, performing a binary classification.
- y is the output of the *Activation function*. (i.e. the classification). Typically denoted as f(x; w)

$$y = g\left(\sum_{i=0}^{k} x_i w_i\right) = g(\boldsymbol{w}^T \boldsymbol{x})$$

Remark 2.1 - Limitations of Perceptron

A *Perceptron* can only perform linear binary classification so is not useful when two classes are not linearly separable.

Proposition 2.1 - Perceptron Supervised Learning Rule

To make a perceptron learn from misclassifications, we adjust the weight vector \mathbf{w} . One learning rule for this is to update the current weights by a certain proportion of the error made $\Delta \mathbf{w}$.

$$\mathbf{w}_{t+1} = \mathbf{w}_t + \Delta \mathbf{w}$$
 where $\Delta \mathbf{w} = \begin{cases} \eta f^*(\mathbf{x}) \mathbf{x} & \text{if } \overbrace{f^*(\mathbf{x})}^{\text{ground truth prediction}} \neq \overbrace{f(\mathbf{x})}^{\text{prediction}} \\ 0 & \text{otherwise} \end{cases}$

where $\eta \in \mathbb{R}^+$ is the *Learning Rate*. Remember that $f^*(\cdot) \in \{1, -1\}$.

Remark 2.2 - Learning Rate η

The Learning Rate η defines how big the steps learning rule makes towards a minimum. The Learning Rate needs to be tuned as too small a value will mean it takes a long time (and a lot of data) to reach a minimum, whereas too great a value means the minimum can be overstepped and convergence is unlikely.

Proposition 2.2 - Training Process for a Single-Layer Perceptron Let $\{(\mathbf{x}_1, f^*(\mathbf{x}_1)), \dots, (\mathbf{x}_N, f^*(\mathbf{x}_N))\}$ be a set of training data.

The following is an algorithm for learning a good set of weights \boldsymbol{w} for a Perceptron

- i). Initialise the weight vector $\boldsymbol{w} = \boldsymbol{0}$
- ii). Consider next training datum $(\boldsymbol{x}_i, f^*(\boldsymbol{x}_i))$.
- iii). Calculate prediction $f(\mathbf{x}) := g(\mathbf{w}^T \mathbf{x})$.
- iv). Compare prediction $f(\mathbf{x})$ and ground truth $f^*(\mathbf{x})$.
- v). Update the weight vector $\mathbf{w} = \mathbf{w} + \Delta \mathbf{w}$ where $\Delta \mathbf{w} = \begin{cases} \eta f^*(\mathbf{x}) \mathbf{x} & \text{if } f^*(\mathbf{x}) \neq f(\mathbf{x}) \\ 0 & \text{otherwise} \end{cases}$
- vi). Repeat ii)-v) until the training set is exhausted.

Remark 2.3 - Arbitrary Decision Boundaries

Multiple preceptrons can be connected to form networks which are able to define arbitrary decision boundaries, rather than just a linear boundary. See Section 4 for details on these network architectures.

2.1.1 Activation Functions

Proposition 2.3 - Activation Function

There are several choice for the Activation Function including:

• The Sign activation function binarily assigns values depend on whether they are positive or negative. Sign is not differentiable.

$$g_{sign}(x) := \begin{cases} 1 & x \ge 0 \\ -1 & x < 0 \end{cases} = \frac{x}{|x|}$$

• tanh function is a differentiable activation-function^[1]

$$\tanh(x) := \frac{e^{2x} - 1}{e^{2x} + 1}$$
$$\tanh'(x) := 1 - \tanh(x)^2$$

• The *Rectifying Linear Unit* (ReLU) function is a differentiable non-linear activation function.

$$g_{ReLU}(x) := \max\{0, x\}$$
 $g'_{ReLU}(x) := \begin{cases} 1 & \text{if } s \ge 0\\ 0 & \text{otherwise} \end{cases}$

Remark 2.4 - Activation Functions should be Non-Linear

If an *Activation Function* is <u>not</u> non-linear then the deep network can be represented by a shallow network, with a single hidden layer, <u>and</u> the network would not be able to model non-linear data, which is not desirable.

 $^{^{[1]}}$ tanh is not as popular as ReLU as it is saturating (output is bounded above and below) meaning early layers learn slowly.

Remark 2.5 - Limitation of ReLU

Using ReLU introduces a problem of Dying Neurons where a large gradient flowing through ReLU may force the neuron never to activate again (as it pushes the incoming signal to 0). This is bad, as these neurons will no longer contribute to learning anymore.

2.2 Artificial Neural Network



Figure 1: Abstract Diagram of the Structure of an Artifical Neural Network with K hidden layers.

Definition 2.2 - Artificial Neural Network

An Artificial Neural Network is a graph which loosely models the neural network of a brain and is used for classification/regression.

An Artificial Neural Network is structured as a set of layers with each layer only takes input from the layer immediately before, and passes its signal to the layer immediately after. There are three groups of layers:

- i). Input Layer. Observed values.
- ii). Hidden Layers. A collection of perceptrons which are grouped into layers. How the layers pass their output/take an input is defined by the architecture of the neural network
- iii). Output Layer. The classification of the network. This may be a single node which represents a numerical value being predicted, or a collection of N nodes used to classify to one of N classes.

See Figure 1 for an diagram of the structure of an ANN.

Definition 2.3 - Forward Propagation

Forward Propagation is the process a neural network calculating its output, once it is given a set of inputs.

This is done by inserting the observed values into the correct nodes of the input layer. This values are then passed forwards, throught the network, one hidden layer at a time. The flow of data is unidirectional.

Several values are calculated during Forward Propagation

- Signal $s_j^l := (w^l)^T f^{l-1}$. The weighted sum calculated by the perceptron from its inputs, before it is passed to the activation function.
- Output $f_j^l := g_j^l(s_j^l)$. The result from applying the activation function g_l^j to the signal s_j^l .

Here l denotes a layer and j the specific node in that layer.

3 Deep Neural Networks

Definition 3.1 - Deep Neural Network

A Deep Neural Network is an Artifical Neural Network with multiple hidden layers. These multiple hidden layers allow Deep Neural Networks to learn complex non-linear relationships as later layers can compose features identified by earlier layers.

Remark 3.1 - Which DNN Architecture to use?

Input Data	Architecture
Grid-Ordered Data	Convolution DNN
Sequential Data	Recurrent DNN

3.1 General

3.1.1 Utility

Remark 3.2 - Advantages of Deep Neural Networks

Here are some advantages *DNNs* offer over neural networks with a single hidden layer.

- Hierarchical Automatic Modularisation. A deep neural network has many layers, and the information of each layer is available to the succeeding layer. This means each layer can be considered to extract slightly more precise features (e.g. pixel colours → edges → corners → object parts → object class). These modular layers are generated automatically during training.
- Practical Performance. Greater depth gives greater performance than greater width. Note that large networks (either by width or depth) require more training time and larger data sets.
- The Oscillation Argument. There are functions f that can be represented by a <u>deep</u> ReLU network with a polynomial number^[2] of neurons, whereas a <u>shallow</u> network would require exponentially many units.

3.1.2 Overfitting

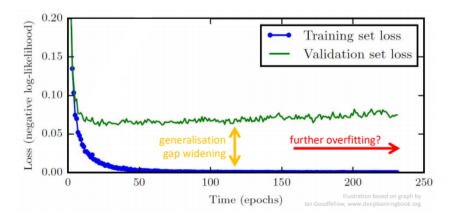


Figure 2: An example of a bad relationship between training and testing loss

^[2] Polynomial wrt input width.

Proposition 3.1 - Overfitting

Deep Neural Networks have lots-and-lots of parameters, meaning they have high degrees of freedom and thus prone to Overfitting (ie the model is fitted too closely to the training data and does not fit unseen data well).

Figure 2 provides an example of how *Overfitting* can be identified graphically as the distance between training and testing loss increases with each time step, and the training loss converges to zero. Ideally the lines for training and testing loss would be fairly similar.

Overfitting can be combatted by: Using more data; Using data which represents the full sample space better; And, strategic sampling of data.

Remark 3.3 - Characterisation of Overfitting in a DNN

Overfitting in a DNN is characterised by weight values being very large. This is due to large weights making the network unstable, as small changes in the input (inc. noise) will have a large affect on the output.

Definition 3.2 - Data Augmentation

Collecting data can be expensive and hard. *Data Augmentation* is a set of techniques to increase the available set of data, without having to collect more data, by slightly modifying already observed data. For images this may involve: cropping; translating; adding noise; shifting the hue...

Definition 3.3 - Regularisation

A Regularisation is any modification made to a learning algorithm which is intended to reduced its generalisation error, <u>but</u> not its training error. This is typically done by introducing more information and making changes to the Loss Function.

L-Regularisation places constraints on the weight space, thus reducing the searchable area for the model.

 L_1 -Regularisation extends the cost function $J(\cdot, \cdot)$ to include a sum of the absolute values of all the weights, with a dampening factor λ .

This gives the following cost function J_{L1} and weight update rule

$$\begin{array}{lcl} J_{L1}(X;W) & := & J(X,W) + \lambda \sum_{w \in W} |w| \\ \\ W^l & \leftarrow & W^l - \eta \left(\nabla J + \lambda \cdot \text{sign}(W^l) \right) \end{array}$$

 L_2 -Regularisation extends the cost function $J(\cdot, \cdot)$ to include a sum of the square value of each weights, with a dampening factor λ .

This gives the following cost function J_{L2} and weight update rule

$$J_{L2}(X; W) := J(X, W) + \frac{\lambda}{2} \sum_{w \in W} w^2$$
$$W^l \leftarrow W^l - \eta \left(\nabla J + \lambda W^l \right)$$

Remark 3.4 - L_1 Regularisation vs L_2 Regularisation

Both L_1 Regularisation and L_2 Regularisation encourage the model to learning the smallest-valued set of weights which minimise the normal cost function. Moreover, they discourage extreme weight values.

The key difference is that, in L_1 Regularisation the less important features are shrunk to zero, removing that feature altogether.

Definition 3.4 - *Dropout*

Dropout is a training procedure which builds on the idea that Neural Networks are an ensemble of sub-networks (See Proposition 1.1) and seeks to allow each sub-network to train somewhat independently.

Dropout does this by, each training loop, choosing a random set of nodes and setting the inbound weight to each of these nodes to 0. This effectively immobilises this set of nodes during this training loop

During validation all weights are turned on, so the output of the network is significantly greater. To combat this weights are set to pW in order to reduce their magnitude (where p is the probability of a particular node being immobilised in a particular training loop).

Dropout decreases training accuracy but should help the model generalise. Without any Dropout it is possible for a model to perfectly classify the training data. Generally p = .5 is a good balance.

Definition 3.5 - *DropConnect*

DropConnect is a variation on *Dropout* where <u>connections</u> are dropped, rather than nodes. This a more fine-grained approach to ensemble learning than *Dropout*.

3.1.3 Extensions

Proposition 3.2 - Extensions of Neural Networks

Here I suggest some areas which could be explored to extend implementation of Neural Networks

- Learn an ensemble of deep networks and then compress them into a single shallow-network. (This is sometimes possible).
- Learn a set of networks which each deal with a specific subtask of the problem and then learn another network which decides which of these specialised networks (or which combination) to use.

3.2 Fully Connected DNN

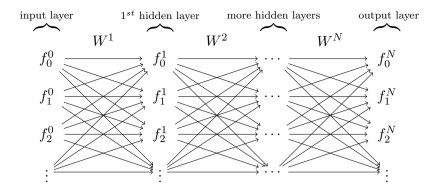
Definition 3.6 - Fully Connected DNN

A Fully Connected DNN is an Artificial Neural Network where each node in a layer takes an input from $\underline{\text{every}}$ node in the preceding layer and passes its output to $\underline{\text{every}}$ node in the next layer.

The output of each perceptron is the weighted sum of its inputs, passed through an activation function.

Below is a diagram of a $MLP^{[3]}$ of depth N (i.e. there are N layers of computation)

^[3] Fully Connected DNNs are also known as Multi-Layer Perceptrons



Note that each layer can have a different number of nodes (AKA width).

For each consecutive pair of layers $\mathbf{f}^i, \mathbf{f}^j$ (of widths n_i, n_j respectively) there is an associated weight matrix $W \in \mathbb{R}^{n_i \times n_j}$ st $\mathbf{f}^j = W^T \mathbf{f}^i$.

Remark 3.5 - Discriminating Power of Different Depth Fully Connected DNNs

A Fully Connected DNN with a <u>single</u> hidden layer is sufficient to represent any boolean or continuous function, althought the <u>layer</u> may be exponentially wider than the input.

A $Fully\ Connected\ DNN$ with two hidden layers is sufficient to represent any mathematical function.

Proposition 3.3 - MLPs as Computation Graphs

$$\begin{array}{lll} \mathbf{s}^{j} & := & (W^{j})^{T}\mathbf{f}^{j-1} & \text{weighted sum of the i^{th} node of the j^{th} hidden layer} \\ \Longrightarrow & \frac{\partial s_{i}^{j}}{\partial w_{ii}^{j}} & = & f_{i}^{j-1} \\ & & f_{i}^{j} & := & g_{i}^{j}(s_{i}^{j}) & & & & & & & & & & \\ \Longrightarrow & \frac{\partial f_{i}^{j}}{\partial s_{i}^{j}} & = & \text{depends on def of g_{i}^{j}} \end{array}$$

Proposition 3.4 - Output Layer for Classification Problem

When using a Fully Connected DNN for classification the output layer represents a probability distribution for each possible class. This means, in the output layer, the node values are in [0,1] and they sum to 1.

This distribution is achieved by using a *Softmax Neuron Group* in the last layer with activation function. Defined as

$$g_j^N(s_j^N) := \frac{e^{s_j^N}}{\sum_{i \in \text{Group}} e^{s_i^N}}$$

This has gradients

$$\begin{array}{rcl} g_{j}^{N'}(s_{j}^{N}) & = & f_{j}^{N}(1-f_{j}^{n}) \\ g_{j}^{N'}(s_{i}^{N}) & = & -f_{j}^{N}f_{i}^{N} & \text{for } i \neq j \end{array}$$

$$g_j^{N'}(s_i^N) = \begin{cases} f_j^N(1 - f_j^n) & \text{if } i = j \\ -f_j^N f_i^N & \text{if } i \neq j \end{cases}$$

3.3 Convolution DNN

Remark 3.6 - This is actually the Cross-Correlation operation, but is what is used in practice. Convolution flips the kernel.

Definition 3.7 - Channels

When we have multiple data readings per instance (e.g. for each pixel of an RGB image) we consider each of these data fields to be a *channel*. When we apply a convolution they must have the same dimension as the number of channels, and they can applied to both space & channels.

Definition 3.8 - Convolution DNN (CNNs)

A Convolution DNN is an Artificial Neural Network which contains a Convolutional Layer (it may also contain fully connected layers). Pooling Layers are common in CNNs in order to reduce dimensionality.

• A Convolutional Layer takes an n-dimensional grid-like topology (e.g. an image or video) as an input and applies several convolutions to this input (rather than matrix multiplication). The output from the convolution in each position of the input is passed through an Activation Funtion and then to the node in the layer at the equivalent position. The convolutional layer is three dimensional with X, Y representing spatial data and Z the convolution applied.

A Convolutional Layer has two possible additions Zero Padding and Stride Length. See Definition 4.4 and Definition 4.5.

• A *Pooling Layer* takes an *n*-dimensional grid-like topology as an input and for each position it outputs a summary of the values in the neighbourhood of that position. This summary is typically the: mean, min or max value; and the size of the neighbourhood is user defined. *Pooling* is applied after the convolution and the activation function, and reduces the output size.

Rather than fitting/learning a weights matrix, here we are learning the *Kernel* (AKA *Tensor*) of the convolution.

Definition 3.9 - Zero Padding

The output from *Convolution* is smaller than the original input. This is bad as eventually the dataset would disappear. To avoid this *Zero Padding* is used. *Zero Padding* adds rings zeros to the outside of the input st the input and output are the same size after *Convolution* is applied.

The number of rings added depends on the size of the Kernel. For a Kernel of size $N \times M$: M-1 rows are added to the top & bottom of the input; and, N-1 columns are added to the left and right of the input.

Definition 3.10 - Stride Length

Stride Length defines how far the convolution operation steps each time. This applies to both horizontal and vertical steps. The greater the Stride Length, the smaller the output will be.

In practice *Stride Length* is rarely set to one as this requires a lot more weights to be fitted and often little is gained by looking at values which are adjacent.

Example 3.1 - CNN Architecture

Figure 3 gives an example of an architecture for a convolution DNN. In this example

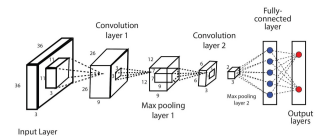


Figure 3: Example of an Architecture for a Convolution DNN

- Input Layer has dimension $36 \times 36 \times 3$. This could be an RGB image with 36×36 pixels.
- Convolution Layer 1 applies 9 different convolutions, each with a kernel of dimension $11 \times 11 \times 3$. In this example a stride of 1 is used and no zero-filtering. This means the output has dimension 26×26 and a depth of 9 (due to 9 convolutions being used).
- Max Pooling Layer 1 applies the max pooling operation to each $3 \times 3 \times 1$ set of values in Convolution Layer 1. In this example a stride of 2 is used, meaning the output has dimension $12 \times 12 \times 9$.
- Convolution Layer 2 applies 3 different convolutions, each with a kernel of dimension $7 \times 7 \times 9$.
- Max Pooling Layer 2 applies the max pooling operation to each $3 \times 3 \times 1$ set of values in Convolution Layer 2.
- A Fully-Connected Layer is the used before the final classification.

Remark 3.7 - Attraction of CNNs

Here are some features which distinguish Convolution Layers away from Fully Connect Layers

- i). Sparse Interactions. In CNNs a node in one layer does not necessarily connected to every node in the next layer. This means that changing this node will not affect every node in the next layer. This is ideal due to the grid structure of the input, where there is implicit relationships between adjacent cells/cell groups (e.g. adjacent pixels in an image).
- ii). Parameter Sharing. The same parameter/weight is used for more than one function in the network. This can be considered as tying two parameters together st that have the same value. We use prior knowledge to decide which nodes to tie together (rather than doing it randomly). This reduces the number of parameters which need to be optimised (which means less data is required for training). The number of reduced parameters increases for later layers. This does not affect the runtime of the forward pass, but significantly reduces the memory requirements of the model.
 - The sharing occurs as the same filter is applied to multiple parts of the image, producing several outputs for its layer. This can be seen as equivalent functions being applied to each location.
- iii). Equi-variant Representations. If the input changes/shifts in a certain way (e.g. translation), then the output changes in exactly the same way. This is as a result of the previous two properties. However, CNNs are <u>not</u> equivariant to rotation or scaling.

Remark 3.8 - Implementing CNNs

In practice there are several difficulties in implementing CNNs, including

• Care needs to be taken when backpropagating CNNs with zero padding or stride greater than 1.

Remark 3.9 - Training CNNs

- The most expensive part of training CNNs is training the convolutional layers. The fully-connected layers at the end are relatively inexpensive as they have a small number of features.
- When performing gradient descent, every gradient step requires a complete run of feedforward propagation and backward propagation through the entire network.

3.3.1 Residual Networks (ResNets)

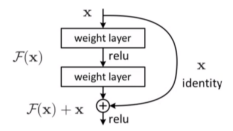


Figure 4: Example of a Residual Network Layer

Proposition 3.5 - Residual Networks (ResNet)

ResNets are a version in CNNs where filters are applied to the input and then merged back (using addition) with the input before being passed to the activation function. This leads to faster convergence by searching for weights which deviate only slightly from the identity. This allows for deeper networks.

See Figure 4 for an example of the architecture of a ResNet layer.

3.4 Recurrent DNN

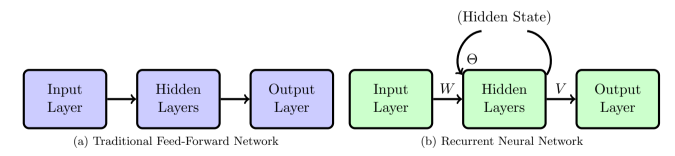


Figure 5: Comparison of the generalised architecture of a traditional *Feed-Forward Network* and a *Recurrent Neural Network*.

Definition 3.11 - Recurrent DNN (RNN)

A Recurrent DNN is an Artificial Neural Network architecture which is designed for sequential data.^[4] This is done carrying part of previously processed data into future processing, The

^[4] When there is likely to be a relationship between a data-point and those immediately before/after it. (e.g. audio, text, time-series data.)

data carried forward is known as the *Hidden State* (See Figure 5).^[5]

Proposition 3.6 - Weights in RNNs

An RNN can be considered to have three sets of weights:

- Input-to-Hidden, W Controls data from the Input Layer to the Hidden Layer.
- Hidden-to-Hidden, Θ Controls data from the Hidden State to the Hidden Layer.
- Hidden-to-Output, V Controls data from the Hidden Layer to the Output Layer.

The same weights are applied to each element in the input sequence, once the weights have be learnt.

Proposition 3.7 - Equations of an RNN

Consider an RNN with a Model $f(\cdot;\cdot)$ and Activation Function $g(\cdot;\cdot)$. Let x_t be the t^{th} datapoint in the input sequence, then

$$\begin{array}{ll} \text{Hidden State} & h_t := f(x_t, h_{t-1}; W, \theta) \\ \text{Outputted Prediction} & \hat{y}_t := g(h_t; V) \end{array}$$

Note that h_0 needs to be initialised in some way, typically to a series of 0s.

Remark 3.10 - An RNN vs A 1D-CNN

A 1D-CNN accepts the same sequential data as an RNN. Here are some differences between the two

- A 1D-CNN only allows for shallow parameter sharing across time, whereas an RNN applies the same parameters (weights) to each element of the input $^{[6]}$.
- The output of a 1D-CNN is a function of the neighbouring elements of the input (before and after), where an RNN only considers the preceding elements of the input.

$$h_0 \xrightarrow{\hat{y}_1} \begin{array}{cccc} \hat{y}_1 & \hat{y}_{t-1} & \hat{y}_t & \hat{y}_{t+1} \\ \uparrow V & \uparrow V & \uparrow V & \uparrow V & \uparrow V \\ h_0 \xrightarrow{\longrightarrow} h_1 \xrightarrow{\longrightarrow} h_1 \xrightarrow{\longrightarrow} h_{t-1} \xrightarrow{\longrightarrow} h_t \xrightarrow{\longrightarrow} h_{t+1} \xrightarrow{\longrightarrow} \\ \uparrow W & \uparrow W & \uparrow W & \uparrow W \\ x_1 & x_{t-1} & x_t & x_{t+1} \end{array}$$

Figure 6: A visualisation of unwrapping an RNN. This shows the dependencies between input data $(x_1, x_2, ...)$ and each hidden state $(h_1, h_2, ...)$.

Proposition 3.8 - Unrolling an RNN

Unrolling an RNN is the process of expanding the recurrence in the Model $f(\cdot)$ over a finite number of steps. This is possible as the Model f and Weight Parameters W, Θ, V all maintain the same size through all steps.

^[5]In this module we assume the *Hidden State* is the output of the *Model f* before the *Activation Function g* has been applied.

^[6] This is deep computational sharing

Consider unwrapping the first three steps (See Figure 6)

$$h_1 = f(x_1, h_0; W, \Theta)$$

$$h_2 = f(x_2, h_1; W, \Theta)$$

$$= f(x_2, f(x_1, h_0); W, \Theta)$$

$$h_3 = f(x_3, h_2; W, \Theta)$$

$$= f(x_3, f(x_2, f(x_1, h_0)); W, \Theta)$$

Remark 3.11 - Purpose of Unrolling

Unrolling creates an explicit expression for each Hidden State in terms of the: Model, f; Weight Parameters, $W\theta$; and the initial Hidden State, x_0 . This expression is then used to learn all these features.

Remark 3.12 - Properties of RNN

Here are some notable properties of an RNN:

- RNNs scale better to much longer sequences of data than other ANN architectures.
- RNNs are Lossy The Model f is a lossy summary [7] of the input x, this means that some information is lost each time the model is applied. Moreover, the influence x_{t-n} has on h_t decreases as n increases, which may not be desirable. [8] [9]

 $Gated\ RNNs$ attempt to mitigate this issue by controlling what information is passed onto the next layer.

Variations of RNNs

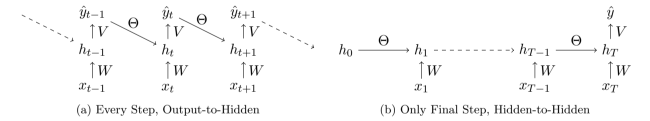


Figure 7: A visualisation of the unwrapping of different RNN architectures.

Proposition 3.9 - Simple Variations of RNNs

Some simple variations on the architecture of an RNN are to vary the recursive relationship between layers^[10] and how often predictions are made:

	Freq. Predictions	Recursive Relationship	Figure	Comment
I	Every Step	Hidden-to-Hidden	Figure 6	Computationally Complete
II	Every Step	Output-to-Hidden	Figure 7 (a)	Less powerful
III	Only Final Step	Hidden-to-Hidden	Figure 7 (b)	Only used to produce summaries.

^[7] It is a lower dimensional projection of the input, meaning not all information from the input is carried forward.

^[8] This is a realisation of the *Vanashing gradient problem* as gradients propagated over long periods of time tend to either vanish or explode. Neither of which is useful.

^[9] Begin lossy results in a Short-Term Memory where more recently seen data has a greater affect on the output.

^[10] Change what is used as the *Hidden State*.

Proposition 3.10 - Bi-Directional RNN

When processing input element x_t a Bi-Directional RNN utilises information from before and after x_t . This is implemented as two RNNs each with their own set of weight parameters: One moving forward; and, the other moving backwards. The hidden values determined for x_t by each of these RNNs is used to make the final prediction \hat{y}_t .

Suppose the forwards RNN has weight sets V_h and calculates hidden values $\{h_1, \ldots, h_T\}$, and, similarly, the backwards RNN has V_k and $\{k_1, \ldots, k_T\}$. Then

$$\hat{y}_t = g(V_h h_t, V_k k_t + b)$$

Bi-Directional RNNs are popular for text problems such as translation and speech recognition, as the context to a word can come before and after it.

Proposition 3.11 - Encode-Decoder RNN

An Encoder-Decoder RNNs map from one variable-length sequence to another variable-length sequence^[11]. Encoder-Decoder RNNs "encodes" the whole input, in one go, to a fixed sized vector c (The Context). c is then "decoded" to produce the output, each time an output element is produced c is updated and this continues until a stopping criteria wrt c is reached. [12]

Encode-Decoder RNNs are used for translation as a sentences are of variable length and it is likely equivalent sentences in different languages will contain a different number of words.

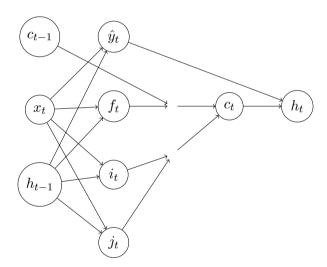


Figure 8: Example of the architecture of an LSTM

Definition 3.12 - Gated RNN

Gated RNNs attempt to mitigate the short-term memory problem mentioned in Remark 3.11. This is achieved by having "gates" which create several paths for the input sequence, allowing the network to accumulate some specified information and forget others, over a long period of time. Connection weights generally vary wrt time.

Long Short-Term Memory RNNs (LSTMs) are the most popular form of Gated RNN, while Gated Recurrent Unit (GRU) are a simpler version of an LSTM which is similarly powerful.

Proposition 3.12 - LSTM RNN

 $^{^{[11]}}$ The input and output can be sequences of any length, and not necessarily the same length.

^[12]When predicting output \hat{y}_t the context c and all previous outputs $\hat{y}_1, \dots, \hat{y}_{t-1}$ as used.

A Long Short-Term Memory RNN (LSTM) is a popular example of a Gated RNN (See Figure 8). LSTMs has the normal input x_t , hidden states h_{t-1} and output \hat{y}_t nodes, but some new "gates" as well:

- Forget Gate, f Determines what old information from h_{t-1} should be forgotten (ie not included in h_t).^[13]
- External Input Gate, i Determines what new information from x_t should be carried forward h_t .^[13]
- State Gate, c Updates what is currently stored.

Proposition 3.13 - GRU

A Gated Recurrent Unit (GRU) is a simplified LSTM, although likely to be as powerful. GRUs just have an Update Gate (What to forget) and a Reset Gate (What to learn), meaning only a single equation is required to calculate the hidden value h_t in a GRU

Training an RNN

Proposition 3.14 - Training an RNN

A variation of Backpropagation, called Backpropagation Through Time, is used to train an RNN. Computing the gradient of the Loss Function L is expensive in an RNN as it requires a forward pass and an Unrolling of the network (despite the same parameter values W, Θ, V be used at each time step) and then a backward pass. This can not be reduced by parallelisation as the forward pass is sequential and all values of the forward pass are required for the backward pass. The run-time and memory costs are both O(T).

4 Algorithms

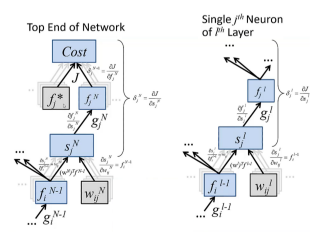


Figure 9: Computational Graph of an ANN

Remark 4.1 - Figure 9

• $J(\cdot, \cdot)$ is the cost function.

^[13]Uses a the input x_t and hidden state h_{t-1} to calculate a value in [0,1] to determine how much information to remember/forget.

- $s_j^l := (w^l)^T f^{l-1}$ is the signal of the j^{th} node in the l^{th} layer.
- $g_i^l(\cdot)$ is the activation function of the j^{th} node in the l^{th} layer.
- $f_j^l := g_j^l(s_j^l)$ is the output of the j^{th} node in the l^{th} layer.

4.1 Backpropagation

Definition 4.1 - Backpropagation

Backpropagation is an algorithm for training feedforward neural-networks. Backpropagation uses Reverse Auto-Differentiation (See Section 0.2) o calculate the derivative of the loss function wrt each weight. The value of the derivative then defines how much to update each weight by.

Proposition 4.1 - Backpropagation Algorithm

For each training sample $(\mathbf{f}^0, \mathbf{f}^*)$ where \mathbf{f}^0 is the input and \mathbf{f}^* is the true output.

i). Read the input & perform a forward pass through the network to calculate signals s_j^l and outputs f_i^l for all nodes in all layers.

$$s_j^l := (\mathbf{w}_j^l)^T \mathbf{f}^{l-1} \quad f_j^l := g_j^l(s_j^l)$$

- ii). Calculate the cost function value $J(f_j^*, f_j^N)$ between each output layer neuron f_j^N and its target value f_j^* .
- iii). Calculate the error derivatives δ_j^{N+1} of the cost function J wrt each output of the last hidden layer f_j^N (ie the values of the output layer).

$$\delta_j^{N+1} := \frac{\partial J}{\partial f_j^N}$$

iv). Compute the error derivative δ^N_j of the cost function wrt the signals of the last hidden layer s^N_j .

$$\delta_j^N := \frac{\partial J}{\partial s_i^N} = g_j^{N'}(s_j^N) \cdot \delta_j^{N+1}$$

NB - This is preparing for Auto-Differentiation.

v). Layer-by-layer: Calculate the error derivatives δ_i^{l-1} of the cost function wrt the signal s_j^{l-1} of each neuron in the next layer, using the error derivatives δ_i^l of the layer above.

$$\delta_i^{l-1} := \frac{\partial J}{\partial s_i^{l-1}} = g_i^{l-1'}(s_i^{l-1}) \sum_{j=1}^{d(l)} w_{ij}^l \delta_j^l$$

where d(l) is the width of the l^{th} layer.

NB - This is applying Auto-Differentiation.

vi). Calculate the error derivates $\frac{\partial J}{\partial w_{ik}^l}$ wrt to the weights of each neuron w_{ik}^l using the error derivatives of the neuron activities δ_i^l .

$$\frac{\partial J}{\partial w_{ij}^l} := \frac{\partial J}{\partial s_j^l} \frac{\partial s_j^l}{\partial w_{ij}^l} = \delta_j^l f_i^{l-1}$$

NB - This is applying the Chain Rule.

vii). Update all weights w_{ij}^l based on the deltas and neuron activities using the Gradient Descent learning rule

$$\begin{aligned} w_{ij}^l &\leftarrow & w_{ij}^l - \eta f_i^{l-1} \delta_j^l \\ \mathbf{w}^l &\leftarrow & \mathbf{w}^l - \eta (\mathbf{f}^{l-1})^T \boldsymbol{\delta}^l \end{aligned}$$

where η is a pre-defined learning rate.

NB - This is the *Gradient-Descent* update rule.

Proposition 4.2 -
$$Step \ iii)$$
 - $\frac{\partial J}{\partial f_i^N}$

The derivative of the Cost Function J wrt the outputs \mathbf{f}^N depends on the definition of the Cost Function and thus on the Distance Measure $L(\cdot, \cdot)$ it uses.

Here are some examples

• Empirical Risk and Quadratic Loss Function

$$\begin{split} L(x,y) &:= (x-y)^2 \\ \text{and } J(\mathbf{f}^*,\mathbf{f}^N) &:= \frac{1}{m} \sum_{i=1}^m (f_i^* - f_i^N)^2 \text{ where } |\mathbf{f}^*| = |\mathbf{f}^N| = m \\ & \Longrightarrow \frac{\partial L}{\partial y} = -2(x-y) \\ & \Longrightarrow \frac{\partial J}{\partial \mathbf{f}^N} = \left(-\frac{2}{m} (f_1^* - f_1^N) & \dots & -\frac{2}{m} (f_m^* - f_m^N) \right) \\ & \Longrightarrow \delta_j^{N+1} &:= \frac{\partial J}{\partial f_j^N} = \frac{-2}{m} (f_j^* - f_j^N) \end{split}$$

• Empirical Risk and Absolute Loss Function

$$L(x,y) := |x-y|$$
and $J(\mathbf{f}^*, \mathbf{f}^N) := \frac{1}{m} \sum_{i=1}^m |f_i^* - f_i^N| \text{ where } |\mathbf{f}^*| = |\mathbf{f}^N| = m$

$$\Rightarrow \frac{\partial L}{\partial y} = -\frac{(x-y)}{|x-y|}$$

$$\Rightarrow \frac{\partial J}{\partial \mathbf{f}^N} = \left(-\frac{(f_1^* - f_1^N)}{|f_1^* - f_1^N|} \dots - \frac{(f_m^* - f_m^N)}{|f_m^* - f_m^N|}\right)$$

$$\Rightarrow \delta_j^{N+1} := \frac{\partial J}{\partial f_j^N} = -\frac{(f_j^* - f_j^N)}{|f_j^* - f_j^N|}$$

Proof 4.1 - Derivation of δ_i^{l-1}

$$\begin{split} \delta_i^{l-1} &:= & \frac{\partial J}{\partial s_i^{l-1}} \\ &= & \sum_{j=1}^{d(l)} \underbrace{\frac{\partial J}{\partial s_j^l}}_{\delta_j^l} \underbrace{\frac{\partial s_j^l}{\partial f_j^{l-1}}}_{w_{ij}^l} \underbrace{\frac{\partial f_i^{l-1}}{\partial s_i^{l-1}}}_{g_i^{l-1'}(s_i^{l-1})} \\ &= & \sum_{j=1}^{d(l)} \delta_j^l w_{ij}^l g_i^{l-1'}(s_i^{l-1}) \\ &= & \underbrace{g_i^{l-1'}(s_i^{l-1})}_{\text{independent}} \sum_{j=1}^{d(l)} w_{ij}^l \delta_j^l \end{split}$$

Definition 4.2 - ∇J

The derivative for layer l defined below

$$\nabla J^{l} := \begin{pmatrix} f_{1}^{l-1} \delta_{1}^{l} & f_{1}^{l-1} \delta_{2}^{l} & \dots \\ f_{2}^{l-1} \delta_{1}^{l} & f_{2}^{l-1} \delta_{2}^{l} & \dots \\ \vdots & \vdots & \ddots \end{pmatrix}$$

Remark 4.2 - Limitations with Backpropagation Algorithm

Here are some limitations to the *Backpropagation Algorithm*. These are part of the reason why the practice of deep learning did not start earlier.

- The Vanishing Gradient Problem. Gradients are unstable/noisy when you backpropagate gradients in a very deep network, meaning that when the true value of the gradient gets very small it is lost in noise.
- Descent-based optimisation techniques need to work <u>accurately and fast</u> in practice, despite large training data sets. This was not possible <u>before GPU</u> parallelisation and improved optimisers.
- Regularisation techniques are critical to achieve good generalisation beyond the training data available (avoid overfitting).

Remark 4.3 - Activation Functions need to be Differentiable & Non-Linear

In the *Backpropagation Algorithm* the derivative of each activation function is used, so each activation function must be differentiable.

The step-function does not fulfil this. This is addressed (usually) by using the ReLU activation function.

4.2 Gradient Descent

Definition 4.3 - Gradient Descent

Gradient Descent is an iterative algorithm for finding a local minimum of a differentiable function. In an ANN this is done by learning a set of weight values \boldsymbol{w} which produce a local minimum for a given cost function J. The update rule for gradient descent is

$$\boldsymbol{w}_{t+1} = \boldsymbol{w}_t - \underbrace{\eta \cdot \nabla J(X; \boldsymbol{w}_t)}_{\Delta \boldsymbol{w}}$$

where η is a specified learning rate and $\nabla J(X; \boldsymbol{w}_t)$ is the partial derivative of the cost function wrt to the weights (ie the direction of fastest descent). We calculate the i^{th} component of $\Delta \boldsymbol{w}$ after observing as $(\boldsymbol{x}, f^*(\boldsymbol{x}))$

$$[\Delta \boldsymbol{w}]_i = \eta x_i (\underbrace{\boldsymbol{w}_t^T \boldsymbol{x}}_{f(\boldsymbol{x}; \boldsymbol{w}_t)} - f^*(\boldsymbol{x}))$$

Definition 4.4 - Online Gradient Descent

Below is an online algorithm for *Gradient Descent* (ie you can pass it new data without having to restart the process).

- i). Initialise all weights W randomly.
- ii). for each training sample (x, f^*) do:
 - (a) Forward propagate to calculate network output values.
 - (b) Back propagate to calculate ∇J^l for each layer l
 - (c) Update weights for each layer $l W^l \leftarrow W^l \eta \nabla J^l$.
 - (d) if (stopping criteria met) break loop.
- iii). return final weights W^l for all layers.

Definition 4.5 - Simulated Annealing

Simulated Annealing is a process for setting the learning rate η by testing τ learning rates in the interval $[\eta_{\tau}, \eta_0]$. Annealing transitions from η_0 to η_{τ} .

- i). Initialise all weights W randomly.
- ii). for $k = 0, \ldots, \tau$ do:
 - (a) $\eta_k := \left(1 \frac{k}{\tau}\right) \eta_0 + \frac{k}{\tau} \eta_\tau$
 - (b) for each training sample (x, f^*) do:
 - i. Forward propagate to calculate network output values.
 - ii. Back propagate to calculate ∇J^l for each layer l
 - iii. Update weights $W^l \leftarrow W^l \eta_k \nabla J^l$.
 - iv. if (stopping criteria met) break loop.
 - (c) return final weights W^l .

Remark 4.4 - Limitations of Online Gradient Descent

• Sample Size. Using single samples at a time to find the minimum point of the cost function will only roughly approximate aspects of the cost function gradient. This leads to a very noisy gradient descent which may not find the global minimum at all. This is exasperated if the learning rate η is set too high as the minimum may be overshot.

This addressed by Deterministic Gradient Descent and Stochastic Gradient Descent.

• Constant Learning Rate. Having the same learning rate η for the whole learning process is bad as you cannot skip over shallow gradient, nor focus more iterations in areas of steep gradient. We would rather have weights with a shallow gradient have a greater learning rate, and weights with a steeper gradient have a lesser learning rate.

This is addressed by Learning via Momentum.

• Monotonic Learning Rate. Having the same learning rate η for all weights is problematic as weights have different gradients and thus should be learning at different rates as they will hit shallow/steep areas at different times.

This is addressed by Adaptive Gradient Algorithm and Root-Mean-Square Propagation.

Remark 4.5 - Ideal Step Length

When looking for minima we want to make <u>longer</u> steps in areas of shallow gradient, and <u>shorter</u> steps in area of steep gradient.

This means less time is spent in regions which do not help us towards our goal, and more time in those that might.

4.2.1 Stochastic & Deterministic Gradient Descent

Definition 4.6 - Alternatives to Online Gradient Descent

As Online Gradient Descent's use of a single sample at a time is bad, multiple samples can be used by using mean ∇J . Here are two approaches

- Deterministic Gradient Descent (DGD) where <u>all</u> training samples (X, F^*) are used at once. Given a small enough learning rate η this will process to the true local minimum, but at high computational cost.
- Stochastic Gradient Descent (MiniBatch) where a <u>small subset</u> of training samples (X, F^*) are used each iteration. This is still good at finding a minimum, and much less computationally costly.

For the average of ∇J we use

$$\nabla J = \frac{1}{|X|} \nabla_W \sum_{j} L(\underbrace{f(\mathbf{x}_j, W)}_{\text{prediction}}, f^*)$$

4.2.2 Momentum

Definition 4.7 - Learning via Momentum

Momentum is a extension to the Gradient Descent weight update rule. Rather than a fixed learning rate η , a velocity term v_t is introduced which defines the step length and direction. The closer the direction of the previous step is aligned to the direction of the current step, the longer the step length is.

$$W^l \leftarrow W^l + \underbrace{v_{t+1}^l}_{\text{upper transform}} \quad \text{where} \quad v_{t+1}^l := \alpha v_t^l - \eta \nabla J(X; W_t^l)$$

where α, η are hyperparameters for momentum and learning rate, respectively.

Moreover, $\alpha \in [0,1]$ can be considered a *Frictional Force* as it prevents the steps increasing without limit.

Proposition 4.3 - Nesterov Accelerated Gradient (NAG)

Nesterov Accelerated Gradient is an extension of Learning via Momentum which, instead of calculating the gradient at the current position, looks-ahead at the gradient of the target. This is since Momentum will carry us towards the next location anyway.

Formally we now define weight updates as

define weight updates as
$$W^l \leftarrow W^l + v^l_{t+1} \quad \text{where} \quad v^l_{t+1} = \alpha v^l_t - \eta \nabla J(X; \underbrace{W^l + \alpha v^l_t}_{\text{perview}})$$

NAG is consistently better that Learning via Momentum in practice.

Remark 4.6 - Building Momentum

In $Learning\ via\ Momentum\ and\ NAG,\ Momentum\ builds$ up as we take steps in the same direction and dissipates as we deviate from this direction.

Remark 4.7 - Interpretting v_{t+1}^l

The two terms in the definitions of v_{t+1}^l for both Learning via Momentum and NAG can be interpreted as

 αv_t^l Velocity already in the system.

 $\eta \nabla J$ Acceleration being added to the system.

Remark 4.8 - Limitations of Momentum Methods

Methods which use momentum progress very slowly in shallow plateau regions of the cost function state space as momentum is not able to build up. This can be rectified by tuning the learning rate.

Remark 4.9 - Learning via Momentum vs. NAG

NAG reduces momentum earlier, decreasing the chance of overshooting.

4.2.3 Newton's Method

Proposition 4.4 - Newton's Method

Newton's Method removes all hyperparameters (η and α) and instead uses curvature to rescale the gradient by multiplying the gradient by the inverse Hessian of the current cost function $H(J(X; W_t))$.

Newton's Method takes aggressive steps in directions of shallow curvature, and shorter steps in directions of steep curvature, however it is attracted to saddle points (bad!).

$$W^l \leftarrow W^l - H(J(X; W^l))^{-1} \nabla J(X; W^l)$$

Computing and inverting the Hessian is computationally and space expensive.

Remark 4.10 - The more parameters there are the more likely saddle points are

Saddle points occur when the hessian has both positive & negative eigenvalues. This is more likely when we have more parameters (as the probability of all eigenvalues being positive is low).

Random Matrix Theory states that the lower the cost function J is (ie the closer it is to the global minimum), the more likely it is to find positive eigenvalues. This means that if we find a minimum it is likely to be a good one (i.e. low cost).

Thus, most critical points with higher cost function values are likely to be saddle points, which we can escape using symmetry-breaking descent methods.

4.2.4 Adaptive Gradient Algorithm (AdaGrad)

Definition 4.8 - Adaptive Gradient Algorithm (AdaGrad)

Adaptive Gradient Algorithm (AdaGrad) is an extension to the Gradient Descent weight update rule st each weight can have a different learning rate. Scaling the learning rate η for each weight wrt the past gradients for that weight.

For each weight w_i the following update rule is used

$$w_i \leftarrow w_i - \frac{\eta g_t}{\sqrt{M_t} + \varepsilon}$$
 where $M_t := \sum_{i=1}^t g_i^2$

where η is a hyper-parameter for the learning rate; g_t is the gradient for this weight in this iteration; M_t is the square-sum of all observed gradients for this weight; and, $\varepsilon > 0$ is a small constant to prevent dividing by zero.

Remark 4.11 - Result

The AdaGrad algorithm decreases the learning rate for parameters whose weights have been high in the past, and increases the learning rate for weights which have had small gradients in the past. This is desirable.

A limitation is that all previous observations is given the same influence on the learning rate, we probably want more recent observations to have greater influence.

Further, if the learning rate is dramatically increased/decreased in the early learning iterations then it is hard for it to recover.

4.2.5 Root-Mean-Square Propagation (RMSProp)

Definition 4.9 - Root-Mean-Square Propagation (RMSProp)

Root-Mean-Square Propagation (RMSProp) is an extension to the AdaGrad weight update rule which introduces a smoothing parameter β to combat the aggressive reduction of learning speed.

For each weight w_i the following update rule is used

$$w_i \leftarrow w_i - \frac{\eta g_t}{\sqrt{M_t} + \varepsilon}$$
 where $M_t := (1 - \beta)g_t^2 + \beta M_{t-1}$

where η is a hyper-parameter for the learning rate; g_t is the gradient for this weight in this iteration; M_t is an exponentially decaying average^[14] of the square of all previously observed gradients of this weight; and, $\varepsilon > 0$ is a small constant to prevent dividing by zero.

Remark 4.12 - Result

The *RMSProp* algorithm decreases the learning rate for parameters whose weights have been high in the past, and increases the learning rate for weights which have had small gradients in the past. This is desirable.

The advantage of RMSProp over AdaGrad is that the influence of early observations decreases-exponentially over time.

4.2.6 Adaptive Moment Estimation (AdaM)

Definition 4.10 - Adaptive Moment Estimation (AdaM)

Adaptive Moment Estimation (AdaM) which aims to smooth the incoming gradient g_t in RMSProp. This is done by introducing an exponentially decaying average of all previously observed gradients for this weight G_t (Not just their squares M_t)^[15].

^[14]The influence of each gradient in M_t decays over time. The closer β is to $\frac{1}{2}$ the quicker this occurs.

 $^{^{[15]}}G_t$ is an estimate of the first moment (ie mean) of the gradients. M_t is an estimate of the second moments (ie un-centred variance) of the gradients.

For each weight w_i the following update rule is used

$$G_t := (1 - \alpha)g_t + \alpha G_t$$

$$\bar{G} := \frac{G_t}{1 - \alpha^{t-1}}$$

$$M_t := (1 - \beta)g_t^2 + \beta M_{t-1}$$

$$\bar{M} := \frac{M_t}{1 - \beta^{t-1}}$$

$$w_i \leftarrow w_i - \frac{\eta \bar{G}}{\sqrt{\bar{M}} + \varepsilon}$$

where α, β, η are hyper-parameters for the learning and decay rates; g_t is the gradient for this weight in this iteration; G_t, M_t are an exponentially decaying averages for previously observed gradients of w_i , and their squared value; and, $\varepsilon > 0$ is a small constant to prevent dividing by zero.

 \bar{G}, \bar{M} are bias-corrected versions of G_t, M_t .

Remark 4.13 - \bar{G} and \bar{M}

 \bar{G} forces fast startups.

Using the square-root of \bar{M} in the update to w_i amplifies shallow gradients (since \bar{M} is small) and dampens steep gradients (since \bar{M} is large).

Remark 4.14 - Using AdaM

Applying AdaM to a ReLU-based network is sufficient to perform deep learning but there are other features which need to be decided.

- i). What to set hyperparameters $\alpha, \beta, \varepsilon$ and the size of each training batch.
- ii). How to initialise the network.
- iii). How to avoid overfitting.
- iv). Which loss function to use.

Tuning and trial-and-error of these features is often required to achieve good results from deep learning.

4.3 Cost Functions

Definition 4.11 - Cost Function, $J(\cdot, \cdot)$

A Cost Function is a real-valued measure of how inaccurate a classifier is for a given input configure (ie test data $[X, f^*(X)]$ and weights W). Greater values imply the classifier is less accurate.

Neural networks us training to learn a set of weights which minimise the *Cost Function*, given the training data.

4.3.1 Numerical Cost Functions

Proposition 4.5 - Distance Measures for Numerical Quantities $L(\cdot,\cdot)$

Let $x, x^* \in \mathbb{R}$ with x representing an estimate and x^* the true value.

Here are some popular measures of distance between real-valued quantities.

Identity Distance
$$L(x, x^*) := \mathbb{1}\{x \equiv x^*\} = \begin{cases} 0 & \text{if } x \equiv x^* \\ 1 & \text{otherwise} \end{cases}$$

Absolute Distance $L(x, x^*) := |x - x^*|$

Quadratic Distance $L(x, x^*) := (x - x^*)^2$

Proposition 4.6 - Cost Functions for Numerical Quantities J

Let $[X, f^*(X)]$ be a set of training data, $f_W(\cdot)$ be the values our model predicts when using weight set W and $L(\cdot, \cdot)$ be some Loss Function.

Here are some Cost Functions for classifiers which classify real-valued quantities.

Expected Loss $J(X; W) = \mathbb{E}[L(f_W(X), f^*(X))]$

Empirical Risk
$$J(X; W) = \frac{1}{|X|} \sum_{\mathbf{x} \in X} L(f_W(\mathbf{x}), f^*(\mathbf{x}))$$

4.3.2 **Classification Cost Functions**

Remark 4.15 - Non-Numerical Cost Measures

In non-numerical classification there is no obvious way to define the distance between classification. Thus we cannot define a cost function.

How we combat this depends on how the output layer is defined. When using the setup of the output layer defined in Proposition 4.2, which uses Softmax, the Cross-Entropy Cost Function Function

Definition 4.12 - Cross-Entropy Cost Function J

Let f_j^* be the ground truth for output node j and f_j^N be our predicted value for the ground truth for output node j.

The Cross-Entropy Cost Function J and its derivative δ_i^N wrt signal i is defined as

$$J := -\sum_{j \in \text{Group}} f_j^* \ln(f_j^N)$$

$$\delta_i^N = f_i^N - f_i^*$$

The cost function derivative δ_i^N is just the difference between the predicted value and the true value.

Proof 4.2 - Derivation of Derivative δ_i^N of Cross-Entropy Cost Function

The steepness of the cost function derivative $\frac{\partial J}{\partial f_i^N}$ exactly cancels the shallowness of the softmax

derivative $\frac{\partial f_j^N}{\partial s_i^N}$, leading to an MSE-Style Delta δ_i^N which is propagated backwards from layer

N.

$$\begin{split} \delta_i^N &:= \frac{\partial J}{\partial s_i^N} \\ &= -\sum_{j \in \text{Group}} f_j^* \frac{\partial \ln(f_j^N)}{\partial s_i^N} \\ &= -\sum_{j \in \text{Group}} f_j^* \frac{1}{f_j^N} \frac{\partial f_j^N}{\partial s_i^N} \quad \text{where } f_j^N := \frac{e^{s_j^N}}{\sum_{i \in \text{Group}} e^{s_i^N}} \\ &= -\sum_{j = i} f_j^* \frac{1}{f_j^N} \underbrace{f_j^N (1 - f_j^N)}_{\frac{\partial f_j^N}{\partial s_i^N} \text{ for } i = j} -\sum_{j \neq i} f_j^* \underbrace{f_j^N}_{\frac{\partial f_j^N}{\partial s_i^N} \text{ for } i \neq j} \\ &= -f_i^* (1 - f_i^N) + \sum_{j \neq i} f_j^* \underbrace{f_j^N f_i^N}_{f_j^N} \\ &= -f_i^* + f_i^* f_i^N + \sum_{j \neq i} f_j^* f_i^N \\ &= -f_i^N \sum_{j \in \text{Group}} f_j^* \\ &= f_i^N \underbrace{\sum_{j \in \text{Group}} f_j^*}_{=1} \\ &= f_i^N - f_i^* \end{split}$$

5 Training

Remark 5.1 - Training a Model

Once an architecture for an ANN is decided, there are still decisions to be made about how to training the model. This decisions generally trade-off between: Training Accuracy, Testing Accuracy, and Computational Resources.

Remark 5.2 - Which layers learn most?

The later layers of a network learn "the most", as in their weights will change most from the values they were initialised with.

This is due to gradients of each layer depending upon the gradients of the layer after them (due to direction of backpropagation), meaning gradients shrink over time and thus the earlier layers are not updated as much.

Proposition 5.1 - Testing & Training Data Sets

When training a model on a dataset D we randomly partition the D in two, creating a: training dataset ($\sim 75\%$ of D); and, a testing dataset ($\sim 25\%$ of D).

The training dataset is used to fit the model (ie learn model parameters) and the testing dataset is used to test the generalised predictive power of the fitted model. It is important that the testing dataset remains unseen during model training.

Definition 5.1 - Model Checkpointing

Some models take a long time (weeks) to train. *Model Checkpointing* is the practice of periodically saving model parameters (ie weights) so that if the script crashes then training can be resumed. Further, the model can be backtracked to a version which had better generalised performance (avoiding overfitting).

A *Model Checkpoint* typically only contains parameter values and nothing to describe the computation/architecture of the model, so is only useful with the source code for this model.

5.1 Metrics

Definition 5.2 - Accuracy

Accuracy is a metric of how good a model is at classification. Accuracy is the proportion of a data set which are correct classified by the model (generally stated as a percentage).

For a dataset $\{(x_1, y_1), \dots, (x_N, y_N)\}$ and model $f(\cdot)$, Accuracy is defined as

Accuracy :=
$$\frac{1}{N} \sum_{i=1}^{N} \mathbb{1}\{y_i \equiv f(x_i)\}$$

Definition 5.3 - Per Class Accuracy

Per Class Accuracy is the same as the Accuracy metric, except it considers each class individually. This allows for more nuanced understanding of our models performance

For a dataset $\{(x_1, y_1), \dots, (x_N, y_N)\}$ and model $f(\cdot)$, Per Class Accuracy for class c is defined as

Accuracy_c :=
$$\frac{\sum_{i=1}^{N} \mathbb{1}\{y_i \equiv c \& f(x_i) \equiv c\}}{\sum_{i=1}^{N} \mathbb{1}\{y_i \equiv c\}}$$

5.2 Data Augmentation

Definition 5.4 - Common Transformation

Here is a description of several transformation which can be applied to data

- Brightness Making an image uniformly brighter or darker. There can be an issue if values are clipped at 0 or 255.
- Colour Variation Change the colour profile of an image. Easily done by swapping channel values or using grayscale.
- Cropping/Scaling Removing parts of the image. Be careful to not cut the object out of the image.
- Flipping/Rotating An image can be flipped or rotated on either of its axes and keep its training label.
- Occlusion Obstructing part of an image.
- Padding Pad the image with random noise. Learns to classify regardless of position.
- Random Noise Randomly vary the value of each pixel in an image. Generally only small variations otherwise the underlying image is lost.

- Translation Shifting an image and adding random noise to new empty space.
- Viewpoint Minor geometric transformation to an object

See torchvision.transforms

Definition 5.5 - *Invariant Transformations*

An *Invariant Transformation* of a piece of data, is a transformation which does not invalidate the label applied to that data. Which transformations are invariant depends on the type of data being transformed and the problem which is being tackled.

Here is a table of common problems and some *Invariant Transformations* for them.

Problem	Invariant To
Object Recognition	Translation, Rotation, Scaling/Cropping, Viewpoint
Number Plate Recognition	Translation, Rotation
Action Recognition	Translation, Rotation, Scaling/Cropping, Viewpoint, Speed

Definition 5.6 - Data Augmentation

Data Augmentation is the process of applying Invariant Transformations to training data (It is common to chain transformations). This increases the size of the training set without having to actually collect more data (which is often expensive). Data Augmentation can be done either Online or Offline

Proposition 5.2 - Online v Offline Data Augmentation

In Offline Augmentation data is transform and saved before then training on the larger data set. Offline Augmentation is typically used when transformation operations are expensive.

In Online Augmentation whenever a new training sample is read into the training process, a transformation is applied with some given probability. Online Augmentation leads to greater diversity as a sample is likely to be different each time it is used.

5.3 Tuning

Remark 5.3 - Parameter Initialisation

The Cost Function of a DNN is non-convex, meaning the outcome of optimisation algorithms for the cost function depend on the initial parameter values.

Random initialisations are easy, but in practice it is better to start with a pre-trained model from a relevant problem which has larger data sets. (For images *ImageNet* is a common pre-trained model.)

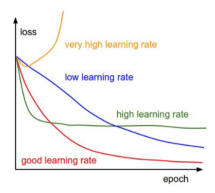


Figure 10: Relationship between learning rate and training accuracy after each training epoch.

Proposition 5.3 - Learning Rate

Learning Rate is the most important hyper-parameter to tune as it affects the rate of convergence. If the Learning Rate is too <u>small</u> then convergence will take a long time to occur, but if the Learning Rate is too <u>high</u> then convergence may never occur as the steps taken are too large. Figure 10 shows the relationship between different learning rates and training accuracy after each epoch.

Batch Normalisation allows for greater Learning Rates to be used.

Proposition 5.4 - Batch Size

Batch Size is the number of training datapoints propagated through the network at any one time, with members of each batch chosen randomly.

Smaller batches are better able to utilise a GPU's parallel processing capability. Although, they likely do not sufficiently represent the gradient of the entire dataset, so the parameter updates overfit to that batch.

Larger batches generally make execution quicker (due to fewer passes) and training accuracy greater (as they better represent the gradient of the whole training set). However, larger batches result in fewer weight updates due to the fewer passes and the increases in training accuracy are not linear with batch size.

Batch size is often limited by the GPU's memory capacity and typically are powers of 2. For small-sized data batches of 32-256 are common, for large-sized data batches of 8-16 are common.

Definition 5.7 - Batch Normalisation

Batch Normalisation normalises layer inputs so the distribution of these inputs does not vary between features while training and adjusting parameters. This speeds up convergence by changing each input distribution to have zero mean and unit variance. The equation used is

$$\hat{x}_{i,c,x,y} = \gamma_c \cdot \frac{x_{i,c,x,y} - \mu_c}{\sqrt{\sigma_c^2 + \epsilon}} + \beta_c$$

where $x_{i,c,x,y}$ is the input to the layer, $\hat{x}_{i,c,x,y}$ is the transformed value which will be used as an input, i,c,x,y define the batch,channel,width & height position of the input, μ_c and σ_c^2 are the mean and variance for each training batch; ϵ is a small constant for numerical stability (ie when σ_c^2 is 0).

 γ_c and β_c are introduced to ensure the ANN can still represent all mathematical functions. Batch Normalisation is typically added after each convolution or fully connected layer, and before the activation function. Batch Normalisation should not be applied to the output layer for classification.

Remark 5.4 - Advantages of Batch Normalisation

Batch Normalisation means all features have the same distribution. In turn this makes the Cost Function more symmetric allowing for a greater learning rate and means the initial parameter values matter less.

0 Reference

Definition 0.1 - Convolution *

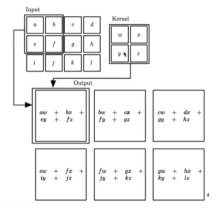
Convolution is an operation which takes two functions f & g and produces a third function (f * g) which shows how one function affects the other.

Consider the 2D discrete space where f, g are real valued 2D matrices (often representing images). Convolution in this case is defined as

$$(f * g)(x,y) = \sum_{i \in I} \sum_{j \in J} f(x-i,y-j) \cdot g(i,j)$$

where I is the horizontal index of g and J is the vertical index of g.

Typically a *Convolution* is then shift right until the RHS of the kernel surpasses the RHS of the input, it will then shift down a row. This will continue until the bottom row has been completed



Note that *Convolution* reduces the size of the input. To avoid this *Zero-padding* is used, where a ring on zeros is placed around the input.

Definition 0.2 - Hessian Matrix $H(\cdot)$

$$H(J) = \begin{pmatrix} \frac{\partial J^2}{\partial w_1^2} & \frac{\partial J^2}{\partial w_1 \partial w_2} & \cdots & \frac{\partial J^2}{\partial w_1 \partial w_n} \\ \frac{\partial J^2}{\partial w_2 \partial w_1} & \frac{\partial J^2}{\partial w_2^2} & \cdots & \frac{\partial J^2}{\partial w_2 \partial w_n} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial J^2}{\partial w_n \partial w_1} & \frac{\partial J^2}{\partial w_n \partial w_2} & \cdots & \frac{\partial J^2}{\partial w_n^2} \end{pmatrix}$$

0.1 Feed-Forward Network

Definition 0.3 - Feed-Forward Network

A Feed-Forward Network is an artificial neural network where the connections between nodes are uni-directional. Data is provided to the input layer and then an output is returned from the output layer, no layers are visited twice.

0.2 Auto-Differentiation

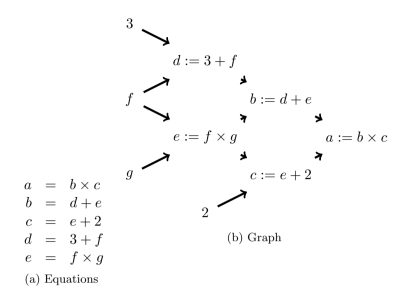


Figure 11: Example of a Feedforward Computational Graph

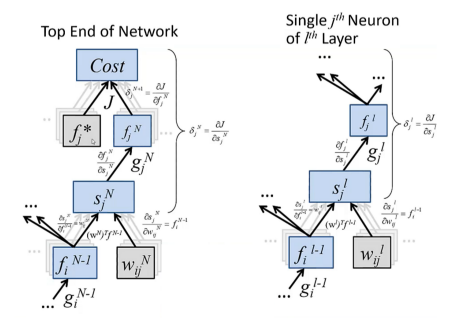


Figure 12: A Neural Network as a Feedforward Computational Graph. J is the cost function, s_j^l , $g_j^l(\cdot)$, f_j^l are the signal, activation function and output of the j^{th} node in the l^{th} layer respectively.

Definition 0.4 - Feedforward Computational Graph

A Feedforward Computational Graph is a uni-directional graph which represents a series of equations. In a Feedfoward Computational Graph nodes represent variables or constants and edges represent mathematical operations (and thus dependence between nodes).

If values are set for all leaves of the graph, then values can be calculated for all nodes in the graph. Moreover, if values are defined for all nodes at a given depth then values can be calculate for all variables higher up the tree.

Figure 11 provides an example of a Feedforward Computational Graph.

Definition 0.5 - Auto-Differentiation

Auto-Differentiation is a technique for calculating partial derivatives of a Feedforward Computational Graph and is used by Gradient Descent.

Consider two nodes in a computational graph x, y. Use the following process to calculate the partial derivative $\frac{\partial x}{\partial y}$.

- i). Establish all the paths from y to x in the graph.
- ii). Calculate the partial derivatives of each step of these graphs. (i.e. if there is a path $y \to a \to x$ calculate $\frac{\partial a}{\partial y}, \frac{\partial x}{\partial a}$).
- iii). Apply the chain rule along each path (i.e. For $y \to a \to x$ calculate $\frac{\partial a}{\partial y} \cdot \frac{\partial x}{\partial a}$).
- iv). Sum these calculations together to get the final result $\frac{\partial x}{\partial u}$.
- v). Substitute variables to make computation easier.

Example 0.1 - Auto-Differentiation using a Feedforward Computational Graph Consider calcualte $\frac{\partial a}{\partial f}$ for the graph in Figure 11.

- i). There are three paths from f to a in the graph: (1) $f \to d \to b \to a$; (2) $f \to e \to b \to a$; and, (3) $f \to e \to c \to a$.
- ii). We need to calculate the following sets of partial derivatives: $\frac{\partial d}{\partial f}, \frac{\partial b}{\partial d}, \frac{\partial a}{\partial b}$ for (1); $\frac{\partial e}{\partial f}, \frac{\partial b}{\partial e}, \frac{\partial a}{\partial b}$ for (2); and, $\frac{\partial e}{\partial f}$, $\frac{\partial c}{\partial e}$, $\frac{\partial a}{\partial c}$ for (3).

iii). Applying the chain rule to each path gives

$$(1) \quad \frac{\partial a}{\partial f} \frac{\partial b}{\partial d} \frac{\partial a}{\partial b} \quad = \quad 1 \cdot 1 \cdot c = c$$

$$(2)$$
 $\frac{\partial e}{\partial f} \frac{\partial e}{\partial f} \frac{\partial b}{\partial c} = g \cdot 1 \cdot 1 \cdot c = gc$

$$\begin{array}{lll} (1) & \frac{\partial d}{\partial f} \frac{\partial b}{\partial d} \frac{\partial a}{\partial b} & = & 1 \cdot 1 \cdot c = c \\ (2) & \frac{\partial e}{\partial f} \frac{\partial e}{\partial f} \frac{\partial b}{\partial e} & = & g \cdot 1 \cdot 1 \cdot c = gc \\ (3) & \frac{\partial e}{\partial f} \frac{\partial c}{\partial e} \frac{\partial a}{\partial c} & = & g \cdot 1 \cdot b = gb \end{array}$$

iv). Summing the terms together we get

$$\frac{\partial a}{\partial f} = c + gc + gb$$

v). By substitution we get a final expression

$$\frac{\partial a}{\partial f} = 2 + 5g + 2fg + 2fg^2$$

So when f = 4, g = 2 we have that a = 150 and $\frac{\partial a}{\partial f} = 60$.

Proposition 0.1 - Using Hierarchical Dependency

By the chain rule we have that $\frac{\partial x}{\partial z} = \frac{\partial x}{\partial y} \frac{\partial y}{\partial z}$. So, if $\frac{\partial x}{\partial y}$ is already known then we just need to multiply that value by $\frac{\partial y}{\partial z}$ to get $\frac{\partial x}{\partial z}$.

This can be utilised to ease the computational load of a calculation. In particular, calculating the derivatives one layer at a time is a good strategy.

Remark 0.1 - Usefulness of Auto-Differentiation

Auto-Differentiation allows us to mathematical quantify the affect one variable has on another, which is good. However, the number of paths in a network grows exponentially with the number of nodes, thus this can be computational hard. (*Hierarchical Dependence* can be used to mitigate this)