COMS30035, Machine learning: Introduction to Neural Networks

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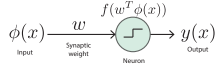
Acknowledgement

► These slides are adapted from ones originally created by Rui Ponte Costa and later edited by Edwin Simpson.

Agenda

- Perceptron
- ► Neural networks (multi-layer perceptron)
 - Architecture
 - ► The backpropagation algorithm
 - ► Gradient descent

- ▶ It is the very beginning of neural network models in ML!
- It is directly inspired by how neurons process information:



It is an example of a linear discriminant model given by $v(x) = f(w^T \phi(x))$

with a nonlinear activation function
$$f(a) = \begin{cases} +1, & a \geq 0 \\ -1, & a < 0 \end{cases}$$

- ▶ Here the target $t = \{+1, -1\}$.
- And we aim to minimise the following error $-\sum_{n=1}^{\infty} \mathbf{w}^T \phi_n t_n^{-1}$

¹Intuitively we want to improve our chances of having $t_n = y_n = -1$ or $t_n = y_n = 1$, which will both decrease our error function.

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$$\phi(x) \xrightarrow[\text{Input}]{\text{Synaptic}} \psi(x) \xrightarrow[\text{Neuron}]{\text{Seuron}} y(x)$$

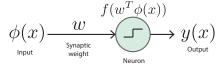
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Example



The Perceptron of Rosenblatt (1962)

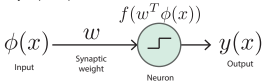
Perceptrons started the journey to the current *deep learning* revolution! Frank Rosenblatt used IBM and special-purpose hardware for a parallel implementation of perceptron learning.

Marvin Minksy, showed that such models could only learn *linearly* separable problems.

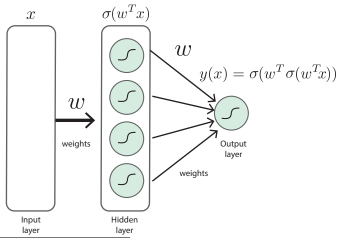
However, this limitation is only true in the case of single layers!

source: Bishop p193.

From a single layer perceptron:



To a Multiple Layer Perceptron (MLP) 2:



 $^{^2 \}mbox{Although, we call it perceptron, it typically uses logistic sigmoid activation functions (continuous nonlinearities), instead of step-wise discontinuous nonlinearities. NB the first and second weight vectors will be different, despite both being denoted by where$

- Neural networks are at heart composite functions of linear-nonlinear functions.
- ▶ Deep learning³ refers to neural networks (or MLPs) with more than 1 hidden layer
- ► They can be applied in any form of learning, but we will focus on supervised learning and classification in particular
- ► MLP recipe ⁴:
 - ▶ Define architecture (i.e. how many hidden layers and neurons) ⁵
 - Define cost function (e.g. mean squared error)
 - Optimise network using backprop:
 - 1. Forward pass calculate activations; generate y_k
 - 2. Calculate error/cost function
 - Backward pass use backprop to compute error gradient
 - 4. Use gradient to improve parameters

⁵Note that this makes them parametric models.



³If you would like to learn more take our Applied Deep Learning unit in your 4th year.

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Neural networks – forward pass step-by-step

- 1. Calculate activations of the hidden layer h: $a_j = \sum_{i=1}^{D} w_{ji}^{(h)} x_i + w_{j0}^{(h)}$ [linear]
- 2. Pass it through a nonlinear function: $z_j = \sigma(a_j)$ [nonlinear⁶]
- 3. Calculate activations of the output layer o: $a_k = \sum_{j=1}^{moderate} w_{kj}^{(o)} z_j + w_{k0}^{(o)}$ [linear]
- 4. Compute predictions using a sigmoid: $y_k = \sigma(a_k)$ [nonlinear⁷]
- 5. All together: $y_k = \sigma \left(\sum_{i=1}^{D} w_{kj} \, \sigma \left(\sum_{i=1}^{D} w_{ji} x_i^{(h)} + w_{j0}^{(h)} \right) + w_{k0}^{(o)} \right)$

⁶In MLP we typically use sigmoid functions.

⁷For classification problems we use a sigmoid at the output, where each output neuron codes for one class.

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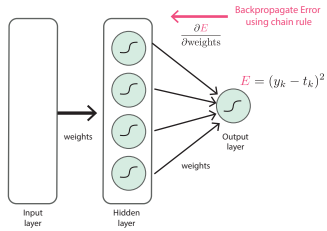


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Neural networks – backward pass

We now need to optimise our weights, and as before we use derivatives to find a solution. Effectively backpropagating the output error signal across the network – backpropagation algorithm.



This is a special case where the output layer has a single node. In the general case we could have e.g. $E = \sum_k (y_k - t_k)^2$.

Total error as a sum of datapoint errors

Typically the error for an entire dataset breaks down into a sum of errors for each individual datapoint:

$$E(\mathbf{w}) = \sum_{n=1}^{N} E_n(\mathbf{w})$$

▶ This means that the error gradient for the entire dataset with respect to some weight w_{ji} will just be the sum of error gradients for each datapoint:

$$\frac{\partial E}{\partial w_{ji}} = \sum_{n=1}^{N} \frac{\partial E_n}{\partial w_{ji}}$$

▶ So we focus attention on computing the $\frac{\partial E_n}{\partial w_{ii}}$ values.

- Although all values in the network depend on the datapoint n we're focusing on, we will (like Bishop) omit the subscript n from network values to reduce clutter.
- Consider an arbitrary unit j (aka 'node') in an neural network.
- ▶ We will associate the *bias* for a unit with a dummy input fixed to 1.
- \triangleright The unit computes a value z_i by first computing a_i , a weighted sum of its inputs (from the previous layer), and then sending a_i to some nonlinear activation function h:

$$a_j = \sum_i w_{ji} z_i \tag{1}$$

$$z_j = h(a_j) \tag{2}$$

(3)

 \triangleright Since E_n (the error for the *n*th datapoint) only depends on weight w_{ii} via a_i we can use the chain rule to write:

$$\frac{\partial E_n}{\partial w_{ii}} = \frac{\partial E_n}{\partial a_i} \frac{\partial a_j}{\partial w_{ii}}$$

$$\frac{\partial E_n}{\partial w_{ji}} = \frac{\partial E_n}{\partial a_j} \frac{\partial a_j}{\partial w_{ji}}$$

► Since $a_j = \sum_i w_{ji} z_i$, $\frac{\partial a_j}{\partial w_{ji}}$ is easy to compute:

$$\frac{\partial a_j}{\partial w_{ii}} = z_i$$

As for $\frac{\partial E_n}{\partial a_j}$ let's just introduce some notation for now: $\delta_j \equiv \frac{\partial E_n}{\partial a_j}$

$$\frac{\partial E_n}{\partial w_{ii}} = \delta_j z_i \tag{4}$$

- ▶ We compute the $\delta_j \equiv \frac{\partial E_n}{\partial a_j}$ backwards, starting with the output units.
- For example, if the error (for a single datapoint) is $E_n = \frac{1}{2} \sum_k (y_k t_k)^2$ then:

$$\delta_k = y_k - t_k \tag{5}$$

► For the hidden units we use (a multivariable version of) the chain rule:

$$\delta_j \equiv \frac{\partial E_n}{\partial a_j} = \sum_k \frac{\partial E_n}{\partial a_k} \frac{\partial a_k}{\partial a_j}$$

"where the sum runs over all units k to which unit j sends connections" (Bishop). "we are making use of the fact that variations in a_j give rise to variations in the error function only through variations in the variables a_k ." (Bishop).

$$\delta_j \equiv \frac{\partial E_n}{\partial a_j} = \sum_k \frac{\partial E_n}{\partial a_k} \frac{\partial a_k}{\partial a_j}$$

can be reformulated to give the backpropagation formula:

$$\delta_j = h'(a_j) \sum_k w_{kj} \delta_k \tag{6}$$

Error Backpropagation

As given in Bishop:

- 1. Apply an input vector \mathbf{x}_n to the network and forward propagate through the network using (1) and (2) to find the activations of all the hidden and output units.
- 2. Evaluate the δ_k for all the outputs units using e.g. (5).
- 3. Backpropagate the δ 's using (6) to obtain δ_i for each hidden unit in the network.
- 4. Use (4) to evaluate the required derivatives.

 \triangleright A key point is that computing a δ_i value helps us compute **several** other $\delta_{i'}$ values.

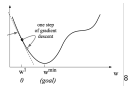
Backpropagation in practice

- It is crucial to organise the computation of all these partial derivatives (i.e. the gradient $\partial E/\partial \mathbf{w}$) efficiently.
- ▶ The quantities required will be organised in tensors (generalisation of matrices beyond 2 dimensions).
- ► The PyTorch approach is called *Autograd*. There's lots of useful info/explanation of how it works on the PyTorch site.

PyTorch's Autograd feature is part of what make PyTorch flexible and fast for building machine learning projects. It allows for the rapid and easy computation of multiple partial derivatives (also referred to as gradients) over a complex computation. This operation is central to backpropagation-based neural network learning. The Fundamentals of Autograd

Neural networks – gradient descent ⁹

In many ML methods it is common to iteratively update the parameters by descending the gradient.



In our neural network this means to update the weights using

- $w_{ji} = w_{ji} \Delta w_{ji}$, where $\Delta w_{ji} = \sigma'(a_k)(y_n t_n)w_{kj}\sigma'(a_j)x_i$
- $w_{kj} = w_{kj} \Delta w_{kj}$, where $\Delta w_{kj} = \sigma'(a_k)(y_n t_n)z_j$
- ▶ This is often done in mini-batches using a small number of samples to compute Δw .

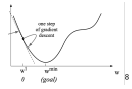
⁹Its called descent because we are minimising the cost function, so descending on the function landscape, which can be quite hilly!



⁸ Figure from https://mc.ai/an-introduction-to-gradient-descent-2/ (link no longer working)

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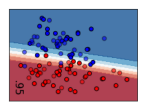
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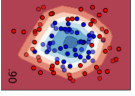


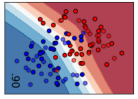
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Example

Using sklearn we fitted a MLP classifier to the data:



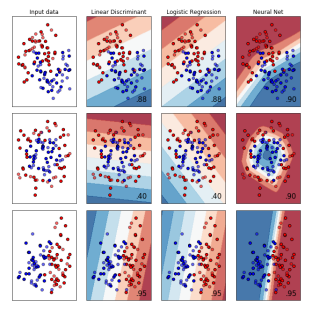




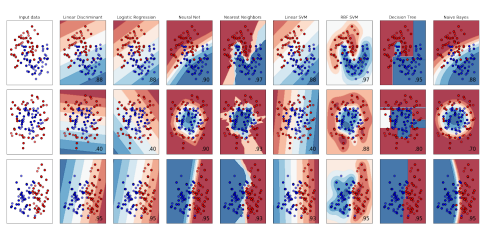
An MLP with one hidden layer can perform well in nonlinear classification problems.

However, because MLPs are highly flexible they can easily *overfit*. Solutions: *early stopping* (stop when test performance starts decreasing) and *regularisation* methods such as *dropout* (randomly turn off units during training).

Classification methods – overall comparison



Classification methods — overall comparison [we don't cover all these methods]



Reading

- ▶ Bishop §5.1 except §5.1.1
- ► Bishop §5.2 except §5.2.2
- ▶ Bishop §5.3 up to §5.3.2
- ► Murphy §13.1-§13.2
- ► Murphy's description of backpropagation in §13.3 is interesting since it is closer to how it is actually implemented, but more complex than Bishop's presentation.

Problems and quizzes

- ▶ Bishop Exercise 4.12
- ▶ Bishop Exercise 5.6 (Tip: Use the result you derived in Exercise 4.12 to help you.)
- Quizzes:
 - Week 2: Neural Networks