

Understanding Neural Networks

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LIP Summer-Student Tutorials

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Seminar Questions

- What are artificial neural networks?
- How do they work?
- How can we improve them?
- Why use them in the first place?

Introduction and history



mita	Antainer chin		
mite	container snip	motor scooler	leopard
mite	container ship	motor scooter	leopard
black widow	amphibian	go-kart	Jaguar
cockroach	amphibian	moped	cheetah
tick	fireboat	bumper car	show leopard
starfish	drilling platform	goircart	Egyptian cat
grille	mushroom	cherry	Madagascar cat
convertible	agaric	dalmatian	squirrel monkey
grille	mushroom	grape	spider monkey
pickup	jelly fungus	elderberry	titi
beach wagon	gill fungus	ffordshire bullterrier	indri
fire engine	dead-man's-fingers	currant	howler monkey





Mark I Perceptron – Rosenblatt, 1957

 First machine to run the single-layer perceptron algorithm

$$f(x) = egin{cases} 1 & ext{if } w \cdot x + b > 0 \ 0 & ext{otherwise} \end{cases}$$

- Weights (w) set using potentiometers
- Used for image recognition, but didn't live up to expectations; couldn't learn properly



- Widrow and Hoff 1960

- (Multi-layer) perceptron machine
- Still hardware-based
- Used a slightly more advanced algorithm to learn the correct weights

Still failed to perform as well as expected



Back propagation – 1960-1986

- Weight-learning based on chain-rule differentiation
- Basics, <u>Kelley 1960</u> and Bryson 1962
- First applied to ANNs in <u>1982 by Werbos</u>
- Shown to be useful in multi-layer ANNs by <u>Rumelhart, Hinton, and Williams in</u> <u>1986</u>
- However, ANNs still underperformed, and were limited in size; training would get stuck
 - Interest in ANNs diminishes

Neural Network Renaissance - 2006

- <u>Hinton and Salakhutdinov</u> develop a layer-by-layer pre-training method
- Allowed backpropagation to work for deep neural-networks
- In 2010 deep neural-networks begin outperforming other methods in speech recognition [<u>Acero, Dahl, Deng, and Yu, 2010</u>]
- Reinvigorated research in NNs

Overview

Example

Example

- Say we want to predict the class (orange or blue) of points according to their position
- We want to draw decision boundaries in our feature space



FEATURES

Which properties do you want to feed in?

X₂²

X₁X₂

sin(X1)

sin(X₂)















Main components of a neural network

1. Neurons

2. The network

3. Training

Overview

Neurons

What is a neuron?

- Quite simply, it is a mathematical transformation:
- It takes vector of inputs <u>x</u>
- Weights each input element
- Applies an *activation function*, e.g sigmoid: f
- And passes its output forwards in the network



What is a neuron?

- The function applied by the neuron can be any continuous mathematical function of the inputs
- However there are several 'standard' ones which are used
- Sigmoid was a common choice



Overview

Networks

 As seen earlier, a network is simply many layers of neurons



- A single neuron applies a basic function to the inputs
- By connecting layers of neurons together, more complex functions can be constructed



 The aim is to learn a function which maps the inputs to the desired outputs



- Each neuron applies the same basic function
- But the weights each neuron applies can be different
- create the map by altering the weights



Overview

Optimisation

Towards training

- How do we alter the weights?
- Could test random settings, but unlikely to arrive at good settings for anything but tiny networks
- Need to alter the weights intelligently, i.e. train the network
- To do this, we need to quantify the performance of the network

Quantifying performance - Loss

- This measure of performance is called a loss function
- It quantises the difference between the network's prediction for a data point, and the actual value of the data point
- Since the inputs are can be thought of has being drawn from a probability density function, rather than an analytic function, the performance of a NN is stochastic
- By evaluating the loss using several sets of inputs (a *mini-bαtch*) a more general value may be computed

Quantifying performance - Loss

• One example is the mean squared-error (minibatch size of *N*):



33

Quantifying performance - Loss

• For classification, the cross-entropy is better:



Network optimisation

- Armed with a quantified measure of performance
- Our aim now is to minimise the loss function ⇒ an optimisation problem
- Lots of advanced algorithms exist: Genetic, Metropolis-Hastings, et cetera
- But the parameter space is huge! \Rightarrow long convergence time

Network optimisation

• Turns out, the gradient descent algorithm works just fine


Network optimisation

- The loss function contains many local minima
- But each is about as optimal as the others
- We simply need to reach to bottom of a high-dimensional bowl
- We do this by moving down the gradient



Gradient evaluation - numerical method

- In order to move down the slope, we first need to know the gradient of the loss function at a given point: $abla \mathcal{L}$
- This can be estimated numerically by varying each weight in the network by a small amount, h, and seeing how the output changes :

$$\frac{\partial f\left(x,y\right)}{\partial x}\approx\frac{f\left(x+h,y\right)-f\left(x,y\right)}{h}$$

This works, but is time-consuming to compute: we can hundreds of thousands of weights to evaluate!

What is a neuron?

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Gradient evaluation - Analytical method

- Because each neuron applies a continuous function, the entire network is differentiable
- We can compute the gradient analytically !

Enter back-propagation

- Essentially, this method of analytical evaluation is a two-step process
- First we do a *forward pass* of a data point, to evaluate the loss
- Next we do a *backwards pass* through the network of the gradient of loss at that parameter point
- We then know exactly how the each weight affect the loss function and can adjust them accordingly
- This is called *back-propagation*

Back-propagation

Example







- Aim is to make decrease the value of *g*(*x*,*y*,*x*)
- Say we have an example data point: *x*=3, *y*=-4, *z*=2
- Let's do a forward pass through our network









- So for our test point, the output is -10
- Now let's back-propagate the gradient
- This will tell us how we should alter the inputs in order to decrease the output



• The output's effect on itself, just one





• Input *z* exerts a force of 1 on the output





• As does the value of *f*(*x*,*y*)





• Let's use the chain-rule:
$$\frac{\partial g}{\partial x} = \frac{\partial g}{\partial f} \times \frac{\partial f}{\partial x}$$













- So, we now know each variable's effect on the output
- Now let's take one step down the gradient
- We'll use a step size (µ) of 0.1









• Having updated our inputs, we find that the output has decreased by 2.72



- In actual implementation we can't change our input data
- Instead we weight the incoming signals
- This is just another 'sub-neuron'
- Meaning we can back-propagate the gradient into it

• Let's generalise and recap



- Let's generalise and recap
- We have a neuron in a network



- Let's generalise and recap
- We have a neuron in a network
- It receives inputs, applies a function, and produces an output



- We have a neuron in a network
- It receives inputs, applies a function, and produces an output
- These inputs come from neurons in the previous layer


- It receives inputs, applies a function, and produces an output
- These inputs come from neurons in the previous layer
- And the outputs are passed to the next layer



- These inputs come from neurons in the previous layer
- And the outputs are passed to the next layer
- At the same time as calculating its output, the neuron can also compute its local gradients



- And the outputs are passed to the next layer
- At the same time as calculating its output, the neuron can also compute its *local gradients*
- Eventually the loss function gets computed



- Eventually the loss function gets computed
- The gradient of the loss eventually gets back-propagated to our neuron
- The neuron sees the effect of its output on the loss



- The neuron sees the effect of its output on the loss
- Having already calculated its local gradients, the neuron simply times this by the incoming gradient (chain-rule)
- The new gradient propagates on to the next layer



- Having already calculated its local gradients, the neuron simply times this by the incoming gradient (chain-rule)
- The new gradient propagates on to the next layer
- Having calculated all the analytic gradients we can update the weights by stepping down the gradient

Problems with neural networks

Back propagation – 1960-1986

- Weight-learning based on chain-rule differentiation
- Basics, Keely 1960 and Bryson 1962
- First applied to ANNs in 1982 by Werbos
- Shown to be useful in multi-layer ANNs by Rumelhart, Hinton, and Williams in 1986
- However, ANNs still underperformed, and were limited in size; training would get stuck
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Problems

- Even with back-propagation, NNs would get stuck during training
- Why did it take another 28 years for them to become useful?

Problems with neural networks

Activation function

Problem 1: Activation function

- The sigmoid function was used because it was smooth between the bounds of zero and one
- Early 'connectionist' interpretations of NNs likened it to the firing rate of a biological neuron
- But it has several problems...



1: It can kill gradients during back-prop

- When |x| is large, the local gradient drops close to zero
- The saturated neuron effectively passes zero loss-gradient back to previous layers
- This stops them from updating their weights



2: The outputs are not zero-centred

- Outputs are always positive
- Gradients propagated to the weights are therefore either always positive or always negative
- If the optimum set of weights is a mixture of positive and negative weights, then this can only be reached by zigzagging towards the optimum position



3: Expensive to compute

- The sigmoid function contains the exponential function
- This requires a lot of CPU time to compute, compared to other functions
- Only a slight slowdown, but a slowdown nonetheless
- Especially once networks start to get large

$$\sigma\left(x\right) = \frac{1}{1 + e^{-x}}$$

An improvement: tanh



The solution



The solution

- Still non-zero centred
- Still kills gradients in negative region
- Depending on initialisation of weights, can sometime never activate (dead ReLu)



Problems with neural networks

Initialisation

Problem 2: Initialisation

- How exactly do we initialise the weights in a network?
- Could set them all to the same value; they'd all respond the same way
- We need something 'symmetry breaking'

Problem 2: Initialisation

- Default was to sample a Gaussian distribution and times by some factor
- If the factor were too large then the neurons would saturate (for sigmoid and tanh); gradients go to zero, nothing trains
- If the factor is too small, the output of the network becomes zero
- Factor must be set carefully by hand

The solution

- Mathematically sensible solution proposed by <u>Bengio and Glorot in 2010</u>: Glorot initialisation
- Scales the Gaussian distribution by $\sqrt{\frac{2}{N_{\text{in}} + N_{\text{out}}}}$
- For neurons with fewer connections, the weights are higher
- For neurons with many connections, the weights are lower
- Similar levels of outputs throughout the network

The solution

- This was derived assuming a linear activation function
- Works well for sigmoid and tanh
- Doesn't work for ReLu; results in lots of dead neurons
- Instead, only the number of inputs should be considered : $\sqrt{}$
- <u>He et al, 2015</u>



Problems with neural networks

Convergence

Problem 3: Convergence time

- Gradient descent is able to optimise the weights
- However, it can easily slow down in narrowly sloping 'valleys'



How GD moves

Ideal moves



Solution 1: Add momentum

- Instead, allow velocity to accumulate:
- Should help move quickly down shallow slopes

$$\label{eq:vt-point} \begin{array}{c} \text{Momentum} \\ \text{coefficient} \\ v_{t+1} = \alpha v_t - \mu \, \nabla \! \mathcal{L}(\theta_t) \\ \theta_{t+1} = \theta_t + v_{t+1} \end{array}$$

Solution 1: Add momentum



- We saw a large speed up in convergence with momentum
- But the method also overshot the target
- The momentum update consists of a momentum step, and a gradient step



Momentum step

- Since we know we'll make the momentum step
- Let's make it first before evaluating the gradient
- Then we'll be evaluating the gradient at the position after the momentum step

Momentum step



- This one-step-lookahead allows for • reduced overshooting
- Allows for quicker convergence •
- $v_{t+1} = \alpha v_t \mu \nabla \mathcal{L}(\theta_t + \alpha v_t)$ $\theta_{t+1} = \theta_t + v_{t+1}$ Referred to as Nesterov momentum

Evaluate gradient after momentum step



Solution 3: Adapt the learning rate

- For steep gradients we want a small learning rate
- For shallow ones, a high learning rate
- Let's give each parameter its own learning rate
- And scale them according to past gradients
- ADAGRAD; Duchi, Hazan, and Singha 2011



Square sum of past gradients

Solution 3: Adapt the learning rate

Over time, the learning rate will drop to zero

- Not so good for deep networks
- Let's allow the store of past gradients to decay
- Effectively keeping a moving average of past gradients
- RMSProp; <u>Hinton & Tieleman, 2012</u>

$$a_{i,t+1} = \lambda a_{i,t} + (1 - \lambda) \nabla \mathcal{L}(\theta_{i,t})^2$$

$$\mu_{i,t+1} = \frac{\mu_0}{\sqrt{a_{i,t+1}}}$$
Leaking store of past gradients

Decay rate

$$\theta_{i,t+1} = \theta_{i,t} - \mu_{i,t+1} \nabla \mathcal{L}_{\theta_{i,t}} D_{\theta_{i,t}}$$

Solution 3: Adapt the learning rate



Final step: Combine them

- Both methods of adding momentum and adapting the learning rate are seen to offer improvements
- No reason why they can't be combined
- This is called ADAM; <u>Ba & Kingma 2014</u>
- And with Nesterov momentum NADAM; <u>Dozat 2015</u>

Improvements
Improvements - Ensembling

- A single model is unlikely to be optimal for all possible inputs
- By training multiple copies of the same model
- Then combining their predictions
- The ensembled model is likely to be more performant in a wider range of input regions
- Effectively a guaranteed improvement!
- Can experiment with different weighting schemes, combinations of architectures, ML algorithms, *et cetera*

Improvements - Dropout

- Slightly counter-intuitive
- Involves randomly dropping (masking) neurons per training iteration
- Means that during that iteration, the dropped neurons are never used
- Hinton et al, 2014



Improvements - Dropout

- Prevents the network from becoming over reliant on certain inputs
- Forces it to generalise to the data
- Effectively trains many sub-networks, i.e. internal ensembling
- Speeds up training (fewer things to evaluate)



Improvements - Dropout

- One subtlety:
- During training perhaps only half the network is used
- During application, all the network is used
- Need to scale outputs during training to maintain similar levels of activation in each regime



Advantages of neural networks

- Direct access to nonlinear responses
- Many previous ML methods have a linear response
- Ensembling them (e.g. random forest; an ensemble of decision trees) could allow for non-linear fitting
- By using a nonlinear activation function, NNs can directly apply nonlinear fitting





NN

Test loss 0.094 Training loss 0.056



NN



predictions of GB (all 200 trees)

train loss: 0.105 test loss: 0.207

BDT

Test loss 0.094 Training loss 0.056





BDT

test loss: 0.207

train loss: 0.105



predictions of GB (all 200 trees)

train loss: 0.096 test loss: 0.162







predictions of GB (all 200 trees)

train loss: 0.129 test loss: 0.275

BDT+rotation



train loss: 0.058 test loss: 0.088

BDT

119

- Power of linear classifiers relies on computing appropriate features beforehand
- These effectively warp the feature-space to make data classes be linearly separable



- HEP example might be the invariant mass of a resonance
- *High-level features* which are nonlinear combinations of other features
- For linear models, these must be calculated by hand and fed in; feature engineering
- High reliance on *domain knowledge*



Summary

- Neural networks are powerful implementations of Machine Learning
- Are able to make use of high-dimensional patterns in data
- Reduced feature engineering
- Must be built with care

Further resources - Lecture & Courses

- Fast.Al Introduction to Machine Learning for Coders
- Fast.Al Practical Deep Learning for Coders
- Fast.Al Practical Deep Learning from the Foundations
- <u>Stanford CS231n</u> lecture series

Further resources - Lecture & Courses

- A disciplined approach to neural network hyper-parameters: Part 1 -learning rate, batch size, momentum, and weight decay - <u>Smith 2018</u>
- SGDR: Stochastic Gradient Descent with Warm Restarts <u>Loshchilov &</u> <u>Hutter, 2016</u>
- Entity Embeddings of Categorical Variables <u>Guo & Berkhahn, 2016</u>
- Regularization for Deep Learning: A Taxonomy <u>Kukačka, Golkov, &</u> <u>Cremers, 2017</u>