Neural Networks and Non-convex Optimisation

Machine Learning Theory (MLT) Edinburgh Rik Sarkar

Course matters

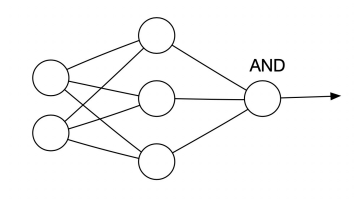
- Please attend tutorials!
- Solutions to tutorial 1 are up
- Coursework will be released by this weekend

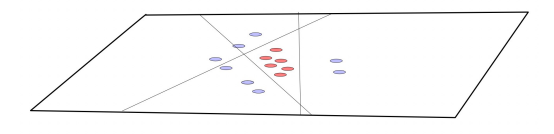
Today's topics

- What neural networks can do: splitting the plane into cells by step activation
- Expressive power of deep neural networks with ReLU
- Softmax function and cross entropy loss
- Generalization and overfitting in neural networks
- Pruning
- Shape of minima flat and sharp

Creating shapes

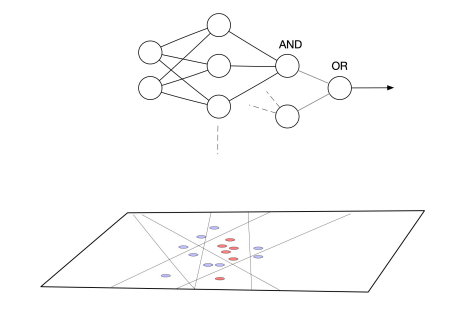
- Neurons with step activation
- Each hidden layer neuron identifies a half-plane
- The output neuron performs AND
- We can create arbitrary polygons this way

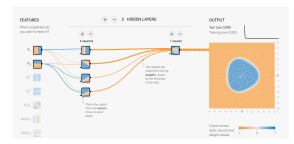




More complex shapes

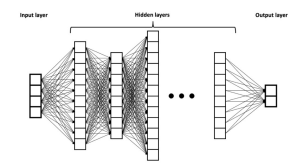
- Compute binary relations with respect to suitable half planes
- Use a Boolean formula to create the right area
- An optimization algorithm finds the right weights
 - to make the half planes
 - And make the Boolean expression





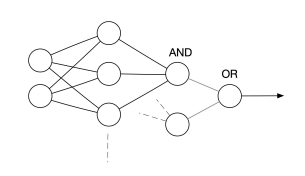
Optimisation in NNs

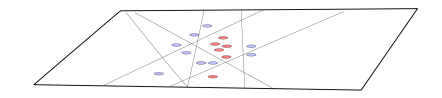
- Usually via SGD and its variants
- Computing gradient:
 - Via chain rule of differentiation
 - Can be done for any sequence of operations
 - In NNs, each layer computes a function of output of the previous layer
 - Chain rule is applied via *backpropagation algorithm*
- Note: ReLU is not differentiable
 - Usually gradient computed via some local analysis
 - There exisit approximate variants that are differentiable

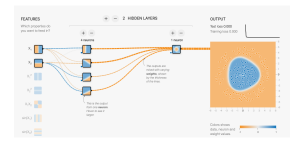


Expressive power of neural networks

- Complex classification requires splitting the space into small cells (polygons)
 - And assigning a class to each
- Larger networks can create more smaller cells and make fine distinctions
 - Great ability or "expressive power"
 - Also greater chances of overfitting

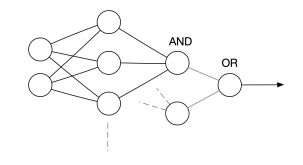


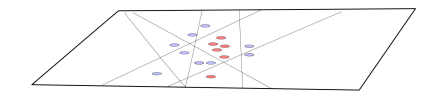


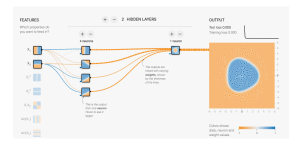


Observation: Non-convexity

- There are many possible assignments of weights
 - To separate the red points
- E.g.
 - Use a different set of lines
 - Same set of lines realised by different neurons
- There are many minima in the space of models (edge weights)
- The optimisation problem is non-convex

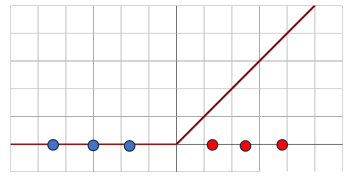


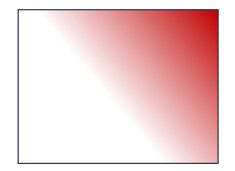




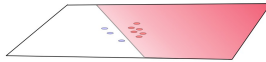
Common activation: ReLU

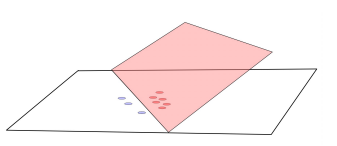
- ReLU
 - $f(x) = \max(0, x)$





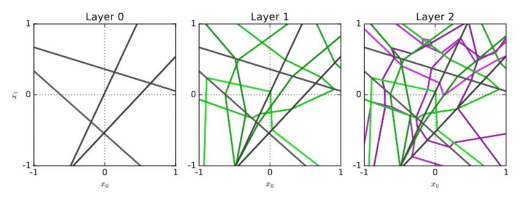
- ReLU does 2 things
 - Creates a half plane separation
 - Gives a score of how deep (far from boundary)
- Effectively, a "score" function on the plane





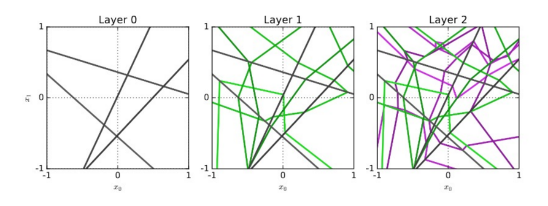
What happens when we train a multi-layer ReLU?

- A ReLU layer
 - Splits domain into cells
 - Also outputs distance from the boundary
 - Acts as coordinate within the cell
- Next ReLU layer further splits each cell!
 - Creating even smaller cells
- Further layers split them into even smaller cells etc...



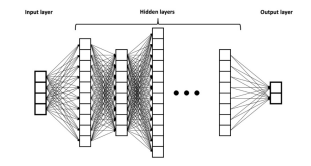
Expressive power of deep ReLU networks

- The complexity of the function that a ReLU neural network can compute grows exponentially with depth
 - It can create more cells, create fine divisions among points
 - And assign them suitable scores
- Raghu et al. On expressive power of deep neural networks.
 - ICML 2017.



Popular Neural network structure

- Use ReLU or similar activation functions
 - More compatible with gradients
 - Easy to compute



- The middle layers produce a vector y of "scores" for each class, called logit values
- Final layer: apply "softmax" to logits:
 - $softmax(y_i) = \frac{e^{y_i}}{\sum e^{y_j}}$ (improved the notation from the lecture)
 - And output the softmax score for each class

Question: Why softmax?

Hard max or exact max

- Take a vector of values eg. [2,3,5,2,6,4,9,2,2,4]
- Make one indicating the position of the max eg. [0,0,0,0,0,0,0,1,0,0,0]

Softmax

- Substitute for hard-max, but differentiable
- Normalized, can be treated as probability p_i for each class

Cross entropy loss

• Neural networks are usually trained on the cross entropy loss of their output p

• Given:

- Data point *x*
- Probability estimate vector p
- Truth label vector *t*: indicator vector or onehot encoding where only the true class has value 1.
- Cross entropy loss: $\ell_{CE} = -\sum t_i \ln p_i$
 - Measures difference between the two probability distributions

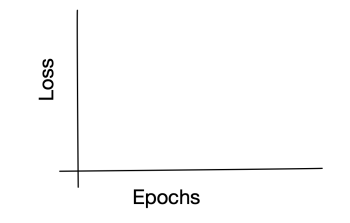
p=[0.1, 0.5, 0.2, 0.2] t= [0.0, 1.0, 0.0, 0.0]

Generalisaiton gap

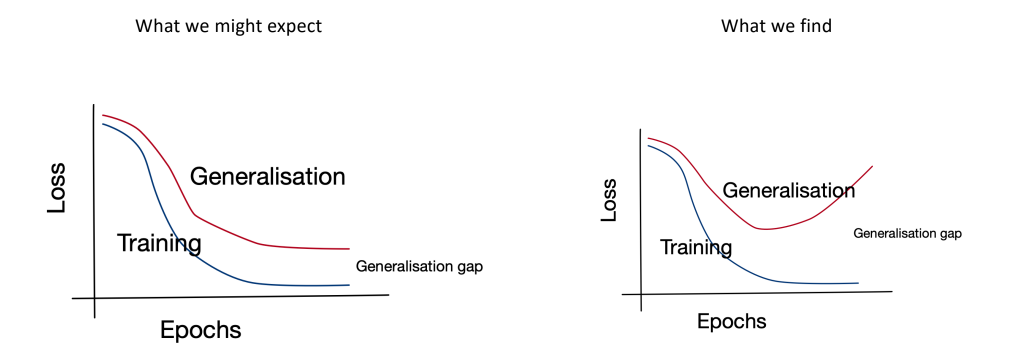
- We can compute the training loss with cross entropy
- And the test loss with cross entropy
- Generalisation gap: difference between the two

Generalisation gap for neural networks

- More training should create better models
- Suppose we plot training loss and test loss
- What do curves look like?



Generalisation gap for neural networks



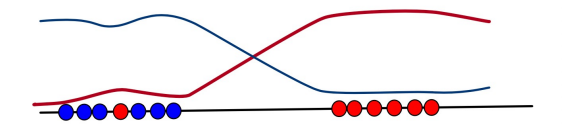
Overfitting in neural networks

- The role of cross entropy loss
- Consider probability outputs for this data and this data space
 - One curve for each class
- What should the curves look like?



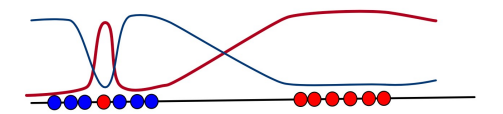
Probability curves for classification

- A reasonable model sacrifices the outlier for better generalization
- But what is the cross entropy loss at the outlier?



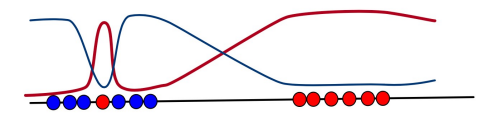
Overfitting

- Optimiser tries to modify probability curves
- Such that large CE losses become smaller



Can this happen with a single neuron?

• Or two hidden ReLU neurons: Red and blue?

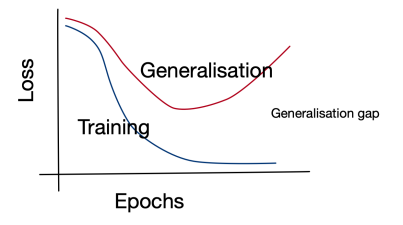


NN and overfitting

- The NN architecture restricts the possible arrangements of hyperplanes
- The architecture and activation functions restrict the scores possible in any one cell
- Smaller networks have implicit regularisation
- In large networks, overfitting does occur, with a small peak at every point
 - Cross entropy loss is not zero until the vector is similar to [0, 1, 0, 0,...]
 - Optimiser continues improving loss even at correctly classified points!

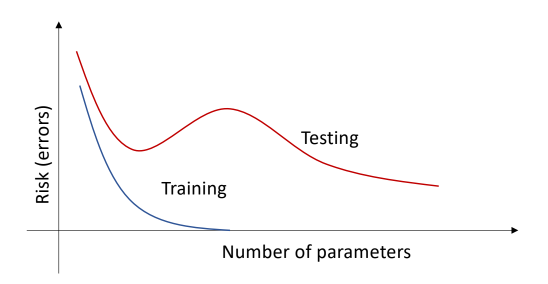
Overparameterised neural networks

- Recap of statistical ML: data requirements grow with parameters/complexity
- Modern neural networks:
 - Many more parameters than data points
 - High complexity and therefore high estimation error
 - We expect heavy overfitting and high test/generalization loss/error



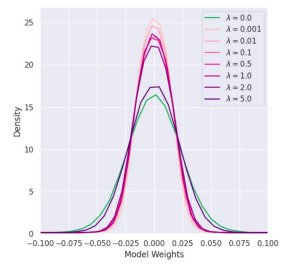
Paradox of Double descent

- With very large number of parameters (more than number of data points) testing performs well again!
- Out of many possible models with low training loss, SGD is finding ones that have low test loss!



Distribution of weights on trained NNs

- A large fraction of weights are close to zero
- Small fraction is far from zero
- Observation:
 - Zero weight edges have no effect do not conduct information
 - Almost zero weights: Little effect
- Conclusion: While NNs have large number of parameters, after training, many of them have little to no effect!



Histogram of weights

Pruning

- Idea: take all the edges that are tiny weights, and remove them!
- Observations
 - Can sometimes remove 80% 90% of edges
 - Retains comparable performance and sometimes better generalization

Lottery ticket hypothesis

• Hypothesis: A randomly initialised dense NN already contains a subnetwork (a winning ticket) that can give good performance.

• Algorithm to find the winning ticket

- Initialise a network to random weights
- Train for some iterations
- Prune p% of edges with small weights
- Reset the remaining edge to their original random weights
- Works surprisingly well on MNIST, CIFAR with test performance comparable to a well trained network [Frankle and Carbin, 2019]

Standard pruning methods

- One shot:
 - Train
 - Remove small weights
 - Return to initialization weights and retrain
 - Stop
- Iterative
 - Set random weights
 - Train
 - Remove edges with small weights
 - Start over

Other results

- Theoretical proofs (special cases, few layers etc)
 - [Malach et al. 2020, Bartoldson et al. 2020]
- Pruning and finding winning tickets without data
 - [Wang et al. 2020, Tanaka et al. 2020]

Pruning and dimension

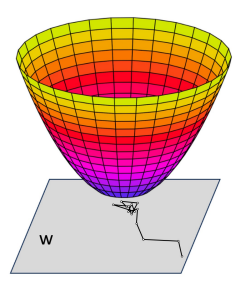
- The dimension of ${\mathcal H}$ is determined by the number of parameters
- The pruning and lottery tickets papers suggest that there are lower dimensional subspaces of ${\mathcal H}$ that contain good solutions

Question

• If a small network is good enough, why are we using a large one?

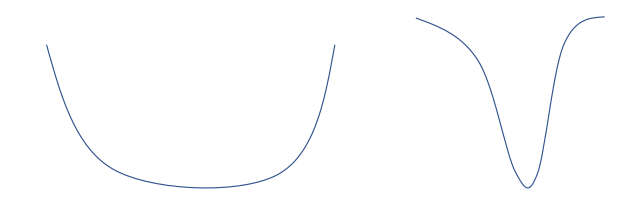
Shape of minima

- Recap:
 - In the space of models \boldsymbol{W}
 - SGD or similar methods try to find a "good" model
 - A minimum of the loss function
- Why it is important
- Hessians and epsilon curvature



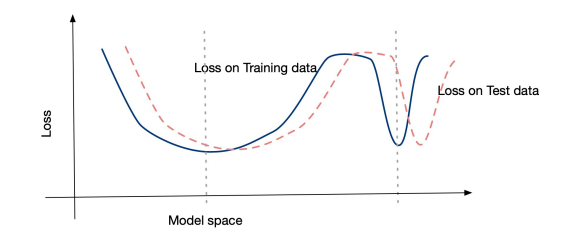
Flat and sharp minima

- A minimum of the loss function can be flat or sharp
- Which is better?

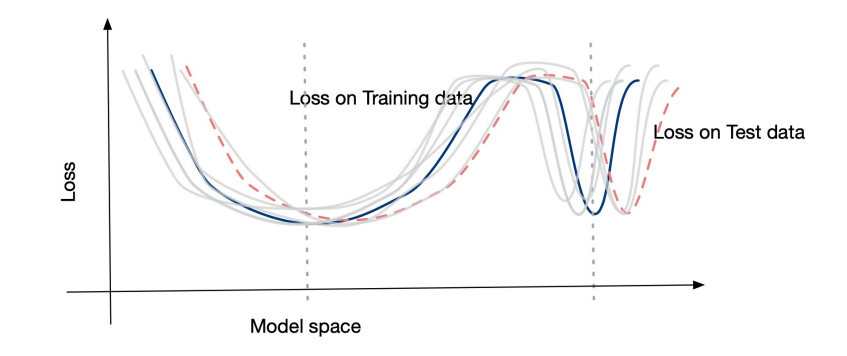


Flat and sharp minima

- Flat minima generalize better
- Sharper minima likely to represent overfitting
 - If we take a slightly different model or slightly different data
 - The loss will jump



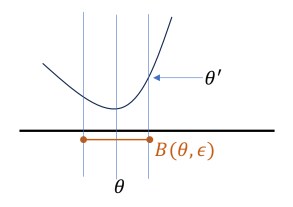
Flat minima are also more likely to be stable



ϵ -Sharpness

- At min θ take ball $B(\theta, \epsilon)$ of radius ϵ
 - Set of all points within a distance ϵ of θ
- Sharpness is:

$$\frac{\max_{\theta' \in B_2(\epsilon,\theta)} \left(L(\theta') - L(\theta) \right)}{1 + L(\theta)}$$



Model spaces are high dimensional

- ϵ Sharpness definition applies directly
- Alternative definition: Curvature
 - Requires considering the Hessian high dim representation of 2nd derivative

Curvature as a sharpness measure

- Mathematical notion of curvature
- For the min of a real valued function in 1-D we can measure curvature as the second derivative

•
$$\frac{d^2y}{dx^2}$$

• For loss over models

•
$$\frac{d^2L}{dw^2}$$

• Larger second derivative => sharper min

Partial derivatives

- Suppose *f* is a function of many variables *x*, *y*, *z*, ...
- We can ask how f changes with x. This is written as $\frac{\partial f}{\partial x}$
 - Same as $\frac{df}{dx}$, but implying that there are other variables to potentially consider
 - And we can write the curvature along x as $\frac{\partial^2 f}{\partial x^2}$: how $\frac{\partial f}{\partial x}$ changes with x

Partial derivatives

- Now we can also ask how $\frac{\partial f}{\partial x}$ changes with y
- This is written as $\frac{\partial^2 f}{\partial y \partial x} = \frac{\partial}{\partial y} \left(\frac{\partial f}{\partial x} \right)$
- Hessian is just a collection of all these written as a matrix
- With two variable models (with *f* as loss):

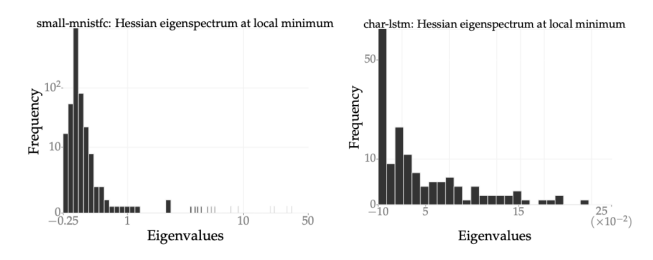
$$\left(\begin{array}{ccc} \frac{\partial^2 f}{\partial w_1^2} & \frac{\partial^2 f}{\partial w_1 \partial w_2} \\ \frac{\partial^2 f}{\partial w_2 \partial w_1} & \frac{\partial^2 f}{\partial w_2^2} \end{array}\right)$$

Curvature directions

- The problem is that strongest directions of curvature may not align exactly with w_1, w_2 etc
- So, we need to take eigen values and eigen vectors of the hessian
- The eigen values represent the principal curvatures
 - Corresponding eigen vectors represent the directions of these curvatures
- Larger eigen values of hessian imply sharper minima
- (Think Principal components of curvature matrix)

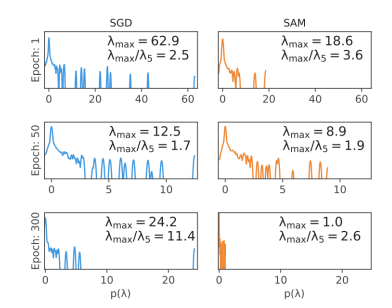
So, the method is

- Take the hessian
- Compute its eigen values
- Look at their distributions
- If there are more of large values, that implies a sharper min



Algorithms

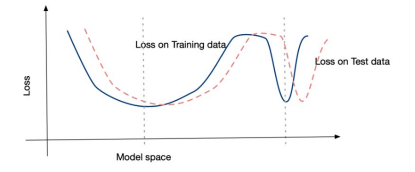
- Shapness aware minimization
 - Use ϵ sharpness
 - Minimize $L(\theta) + [L(\theta + \epsilon') L(\theta)]$
- Entropy SGD
 - Optimise a different function
 - Computationally very expensive
- Stochastic weight averaging
 - Average the weights of the last *c* models
 - Shown to produce flat minima



Flat minima

- Current topic of research
- While flat minima are generally agreed to be good, the full picture is not clear
- There are some works showing that sometimes sharp minima can work well
- Neural nets are highly redundant (e.g. symmetric) and many possible weight assignments achieve the same effective function
 - It is possible to reconfigure weights such that the effective prediction function is same, therefore loss is same, but the curvature is different

SGD and Flat minima



- SGD is known to have a bias toward flat and well generalizable min
- Large batch sizes and small learning rate approximates a smooth gradient
 - And more likely to find a sharp min
- Small batch sizes and larger learning rate makes a more random, jumpy trajectory that can skip over sharp min.
 - Also easy to jump away from sharp min neighborhood since that is likely a small region
- However, a flat min means that even after step away from it, SGD is likely in the same basin and will eventually converge to it

- We have done
 - Statistical ML
 - Neural networks
 - Optimization
- Next
 - Privacy
 - Fairness
 - Explainability