#### Introduction

#### Seminar Series Neural Networks for Finance

#### Lecture 2

Aim

Neural networks in depth: capacity and optimization

24-02-2022

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Deep versus Shallow slides written by Nikolaj Mücke

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# Agenda

- Neural networks: ANNs, DNNs, activations
- Universal Approximation Theorem
- 3 Stochastic Gradient Descent
- 4 Automatic differentiation: descending into a computer
- 5 Gradient based optimization problems
- 6 Deep versus Shallow
  - Shallow Networks

## (Fully-Connected) Feedforward Artificial Neural Networks

• A fully-connected, feedforward artificial neural network is simply a hierarchical composition of the transformations above, mapping an input  $x \in \mathbb{R}^d$  to the output  $\Phi \in \mathbb{R}^q$  whose i'th coordinate reads as follows

$$\Phi_{i}(x|\theta) := \varphi_{i}^{(2)} \left( \sum_{j=1}^{p_{1}} W_{ij}^{(2)} \varphi_{j}^{(1)} \left( \sum_{k=1}^{d} W_{jk}^{(1)} x_{k} + b_{j}^{(1)} \right) + b_{i}^{(2)} \right)$$
where  $\theta := (\theta^{(1)}, \theta^{(2)}) := (W^{(1)}, b^{(1)}, W^{(2)}, b^{(2)}) \in$ 

$$\mathbb{R}^{p_{1} \times (d+1) + q \times (p_{1}+1)}$$
(1)

• The total number of parameters in the statistical model is

$$p = p_1 \times (d+1) + q \times (p_1 + 1) \tag{2}$$

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#### Disclaimer

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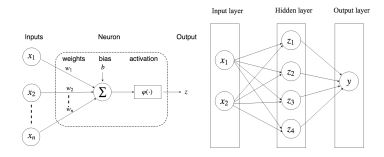
From this lecture on – as is usual in the field of statistics (and also machine learning) – we drop the bold face notation for vectors and matrices (and tensors). Tensor valued entries should be interpreted by context. For example:  $x \leftarrow \mathbf{x} \in \mathbb{R}^d$ ,  $\Theta \leftarrow \mathbf{\Theta} \in \mathbb{R}^p$ ,  $W^{(I)} \leftarrow \mathbf{W}^{(I)} \in \mathbb{R}^{p_I \times p_{I-1}}$ , . . .

## (Fully-Connected) Feedforward Artificial Neural Networks

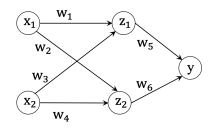
• Denoting the *element-wise* non-linearities by  $\varphi^{(n)}(z^{(n)}) := (\varphi_1^{(n)}(z_1^{(n)}), \dots, \varphi_{p_n}^{(n)}(z_{p_n}^{(n)})) \in \mathbb{R}^{p_n}, \text{ in vector notation this can be written as follows}$ 

$$\Phi(x|\Theta) := \varphi^{(2)} \circ z^{(2)}(\cdot|\theta^{(2)}) \circ \varphi^{(1)} \circ z^{(1)}(x|\theta^{(1)})$$
 (3)

• FCFF ANNs can be thought of as directed, acyclic graphs

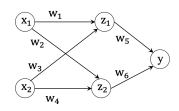


## Example: XOR revisited



- Unlike with SLP, adding a *hidden layer*, and parametrizing with ANNs solves the problem
- Let  $h_1(z_1)$  and  $h_2(z_2)$  be perceptrons with  $b_1 = b_2 = 0$
- Fix  $w_1 = -1$ ,  $w_2 = 1$ ,  $w_3 = 1$ ,  $w_4 = -1$
- The desired output is obtained by setting  $w_5 = w_6 = 1$  and  $b_3 = -1$

# Example: XOR revisited



•  $\widehat{\Theta} = (w_1 = -1, w_2 = 1, w_3 = 1, w_4 = -1, b_1 = 0, b_2 = 0, w_5 = 1, w_6 = 1, b_3 = -1)$  gives and the XOR problem is solved

$x_1$	$x_2$	$z_1$	$z_2$	$a_1$	$a_2$	У
-1	-1	2	-2	1	-1	1
-1	1	0	0	-1	-1	-1
1	-1	0	0	-1	-1	-1
1	1	-2	2	-1	-1 -1 -1 1	1

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Activations: the key feature of non-linearity

#### **Activation Functions**

- The key component of *non-linearity* is given by the **activation** functions  $\varphi_j^{(n)}: \mathbb{R} \to \mathbb{R}, j=1,\ldots,p_n$
- ullet For SLP we had  $arphi_1^{(1)}(x)=2\mathbb{1}_{x-b>0}(x)$ -1
- Most often, application functions within the same layer are chosen the same  $\varphi_1^{(n)}(x) = \cdots = \varphi_{p_n}^{(n)}(x)$
- In regression applications the output activations are usually chosen to be the identity function  $\varphi^{(2)}(z^{(2)}) = z^{(2)}$
- The parametrization inherits continuity and differentiability properties from the chosen activations
- The non-linear activation functions are hyperparameters of the statistical model, appropriate choices vary depending on the problem

### **Activation Functions**

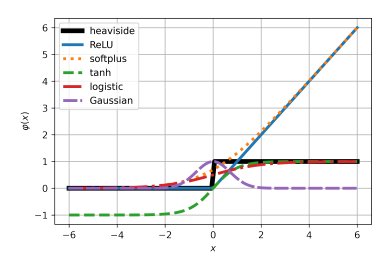
name	$\varphi(x)$	$\varphi'(x)$	range	continuity
heaviside	$\begin{cases} 0, & x \le 0 \\ 1, & x > 0 \end{cases}$	$\int 0, \qquad x \neq 0$	{0,1}	C-1
	$\int 1,  x > 0$	$\begin{cases} undefined, & x = 0 \end{cases}$	(0, 1)	C
ReLU	(0 ×<0	$\int 0, \qquad x < 0$		
	$\begin{cases} 0, & x \le 0 \\ x, & x > 0 \end{cases}$	$\begin{cases} x, & x > 0 \end{cases}$	$[0,\infty)$	$C^0$
	`	undefined, $x = 0$		
Gaussian	$e^{-x^2}$	$-2xe^{-x^2}$	(0, 1]	$C^{\infty}$
tanh	$\frac{e^x - e^{-x}}{e^x + e^{-x}}$	$1 - \tanh^2(x)$	(-1,1)	$C^{\infty}$
logistic	$\frac{1}{1+e^{-x}}$	$\frac{e^{-x}}{1+e^{-x}}$	(0, 1)	$C^{\infty}$
softplus	$\log(1+e^x)$	$\frac{1}{1+e^{-x}}$	$(0,\infty)$	$C^{\infty}$

See wiki

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### **Activation Functions**



## Layer dependent activations

- For classification problems it is chosen such that  $\sum_{k=1}^{q} \varphi_k^{(2)}(z_k^{(2)}) = 1$
- In order to ensure that output *probabilities* indeed sum to one, activations are chosen such that they are not a collection of scalar-valued mappings of neurons but vector-valued functions depending on the whole layer  $\varphi^{(n)}: \mathbb{R}^{p_n} \to \mathbb{R}^{p_n}$

name	$\varphi_i(x)$	$\partial_j arphi_i(x)$	range	cont.
softmax	$\frac{e^{x_i}}{\sum_{i=1}^{p_n} e^{x_j}}$	$\varphi_i(x)(\delta_{ij}-\varphi_j(x))$	(0,1)	$C^{\infty}$
maxout	$\max_{i=1,\dots,p_n} x_i$	$\begin{cases} 1, & j = \arg\max_i x_i \\ 0, & j \neq \arg\max_i x_i \end{cases}$	$(-\infty,\infty)$	C <sup>0</sup>

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Deep Neural Networks

# Fully-Connected Feedforward Deep Neural Networks

- So far: neural network = input + hidden + output layer
- These are called **shallow** feedforward ANNs

$$\Phi(x|\Theta) := \varphi^{(2)} \circ z^{(2)}(\cdot|\theta^{(2)}) \circ \varphi^{(1)} \circ z^{(1)}(x|\theta^{(1)})$$
(4)

- One can extend the model to allow for L-many hidden layers in the composition, with corresponding widths  $p_l$ ,  $l = 1 \dots, L$
- The resulting, deep neural network mapping reads as follows

$$\Psi(x|\Theta) := \varphi^{(L+1)} \circ z^{(L+1)} (\cdot |\theta^{(L+1)}) \circ \cdots \circ \varphi^{(1)} \circ z^{(1)} (x|\theta^{(1)}), \quad (5)$$
 where  $\Theta := (\theta^{(1)}, \dots, \theta^{(L+1)}) \in \mathbb{R}^p$ 

 The total number of parameters in the statistical model rapidly increases

$$p = d \times (p_1 + 1) + \sum_{l=1}^{L-1} p_l \times (p_{l+1} + 1) + p_L \times (q+1) \quad (6)$$

# Fully-Connected Feedforward Deep Neural Networks

$$\Psi(x|\Theta) := \varphi^{(L+1)} \circ z^{(L+1)} (\cdot | \theta^{(L+1)}) \circ \cdots \circ \varphi^{(1)} \circ z^{(1)} (x | \theta^{(1)}), \quad (7)$$

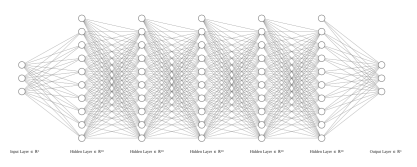


Figure: Illustration DNN, source: NN-SVG

$$p = d \times (p_1 + 1) + \sum_{l=1}^{L-1} p_l \times (p_{l+1} + 1) + p_L \times (q+1)$$

### Seminar Series Neural Networks for Finance

Universal Approximation Theorem

# Most commonly used carpet

#### **Topic**

- We have seen: (deep) neural networks provide *one way* to parametrize non-linear mappings
- We have not seen: is that a "good" way? What class of functions can actually be approximated by them?

### A classical analogy

### Theorem (Weierstrass approximation theorem)

Let  $f \in C([a,b];\mathbb{R})$ . For any  $\epsilon > 0$  there exists a polynomial  $p: x \mapsto a_0 + a_1x + a_2x^2 + \dots$  such that  $\sup_{x \in [a,b]} |f(x) - p(x)| < \epsilon$ .

- Weierstrass approximation theorem: any continuous function can be uniformly approximated by a polynomial functions over closed intervals, with arbitrary accuracy
- Terminology the set of polynomials is **dense** in  $C([a,b];\mathbb{R})$
- Conclusion: polynomials may be a good parametrization for continuous functions
- Notice: not implementable
  - No bound on the order of the polynomial
  - 2 No way to find the "right" polynomial within the class

# Universal Approximation Theorem

- In order to show that neural networks are "good" family of statistical models, we need a similar result → Universal Approximation Theorem (UAT)
- First: Weierstrass for shallow neural networks
- Definition: an activation  $\varphi: \mathbb{R} \to \mathbb{R}$  is called **sigmoidal**, if it is continuous and  $\lim_{x \to -\infty} \varphi(x) = 0$ ,  $\lim_{x \to \infty} \varphi(x) = 1$

### Theorem (UAT - continuous functions, Cybenko, 1989)

Let  $\mathcal{G}_{\varphi} := \{ \Phi(x|\Theta) : \Theta \in \mathbb{R}^p, p \in \mathbb{N}, \varphi \text{ fixed, sigmoidal} \}$  be the family of shallow neural networks of the form (3) with d = q = 1. Then, for any  $f \in C([0,1];\mathbb{R})$  and  $\epsilon > 0$  there exists a shallow neural network  $\Phi(\cdot|\Theta^*) \in \mathcal{G}_{\varphi}$  such that

$$\sup_{x\in[0,1]}|\Phi(x|\Theta^*)-f(x)|<\epsilon.$$

#### Restrictions and extensions

- Similar restrictions as in Weierstrass: arbitrary width, no insight on  $\Theta^*$
- Natural extension from shallow to deep neural networks of the form (7)
- Several results exist, very actively researched area since the early 90s
  - lacktriangledown sigmoidal o non polynomial
  - **2**  $[a,b] \to \mathbb{R} \longrightarrow$  compact subsets in general vector spaces to other vector spaces
  - **3** dense in  $C([a, b]; \mathbb{R})$  → dense in the space of Lebesgue integrable functions ( $L^p$  spaces)
  - approximating functions with arbitrary accuracy → approximating functions and their derivatives with arbitrary accuracy (Sobolev spaces) see Hornik et al., 1990
  - ⑤ arbitrary width/depth → fixed width/depth with dominated error term (curse of dimensionality)

**6** ...

### Seminar Series Neural Networks for Finance

Stochastic Gradient Descent

# Optimization

#### Question

- UAT → neural networks may be an appropriate statistical model for a wide range of phenomena
- UAT: in the space of neural networks there lives a neural network  $\Phi(x|\Theta^*)$  which approximates a given function f(x) with arbitrary accuracy
- We do not know: how do we find ⊕\*, or at least a good approximation of it?

#### Iterative schemes

• Recall: thanks to the **convexity** of  $\beta \mapsto \|y - X\beta\|^2$ , OLS has a closed form expression mapping measurements X, y to the **unique**, **global**, **minimizer** of MSE

$$\beta^* = (X^T X)^{-1} X^T y \tag{8}$$

- In neural network regression the loss is no longer convex  $L(\Theta|D=(x,y))=\|y-\Phi(x|\Theta)\|^2$  due to the non-linearity of  $\Phi(x|\Theta)$  in  $\Theta$
- No closed-form expression for a minimizer 
   iterative schemes needed
- No unique solution to  $\nabla_{\Theta} L(\Theta|D) = 0$
- A solution to  $\nabla_{\Theta} L(\Theta|D) = 0$  may not be an extremum at all (saddle points)
- **local minima** may not be global  $\implies$  solving  $\nabla_{\Theta} L(\Theta|D) = 0$  does not guarantee global minimum

## Iterative schemes – general formulation

- An iterative scheme is a sequence of approximations  $\Theta^n$  such that the resulting approximations  $\Phi^n := \Phi(\cdot|\Theta^n)$  converge to a desired solution H in an appropriate sense (e.g. MSE)
- An iterative scheme needs an appropriate *update/iterative rule* which determines  $\Theta^{n+1}$  given  $\Theta^n$
- How do we define a sensible update rule? → (Stochastic)
   Gradient Descent ((S)GD)

#### Gradient Descent - intuition

- Assume we want to minimize some  $f: \mathbb{R}^p \to \mathbb{R}_+$  non-linear real-valued function with some iterative scheme. Our current guess for the minimum is  $x^k$ . What should our next guess be?  $\longrightarrow$  steepest gradient
- It is always nice to gather an intuition in one dimension

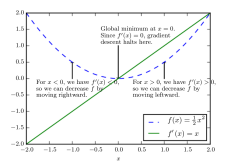


Figure: Goodfellow et al.: Deep Learning (2016), MIT Press [Figure 4.1, pg. 81]

### Gradient Descent – general

- Back to the general setting: given a point  $x^k \in \mathbb{R}^p$  we want to find the **direction**  $e \in \mathbb{R}^p$ , ||e|| = 1 in which the value of f decreases the fastest
- Formally we are concerned with the following minimization problem

$$\begin{split} \underset{e \in \mathbb{R}^p, \|e\| = 1}{\arg\min} \left\langle e \middle| \nabla_x f(x^k) \right\rangle &= \underset{e \in \mathbb{R}^p, \|e\| = 1}{\arg\min} \|e\| \left\| \nabla_x f(x^k) \right\| \cos \left(\vartheta_{e, x^k}\right) \\ &= \underset{e \in \mathbb{R}^p, \|e\| = 1}{\arg\min} \left\| \nabla_x f(x^k) \right\| \cos \left(\vartheta_{e, x^k}\right) \\ &= \underset{e \in \mathbb{R}^p, \|e\| = 1}{\arg\min} \cos \left(\vartheta_{e, x^k}\right), \end{split}$$

where  $\vartheta_{e,x^k}$  is the angle between e and  $\nabla_x f(x^k)$ 

• This is minimized when  $\vartheta_{e,x^k} = \pi$  leading to  $e = -\nabla_x f(x^k) / \|\nabla_x f(x^k)\| \implies$  steepest descent or gradient descent

#### **Gradient Descent**

### Gradient Descent (GD)

The iterative scheme of a gradient descent optimization then reads as

$$x^{k+1} = x_k - \eta^k \nabla_x f(x^k), \tag{9}$$

where the parameter  $\eta^k$  is called a **learning rate** and determines the size of each iteration step

Several strategies for the choice of  $k \to \eta^k$ : constant, adaptive (decay, validation loss linked, etc.), line search, . . .

# Back to machine learning (statistics)

Our goal is to minimize a performance measure (loss)

$$L(\Theta) := \mathbb{E}_{(x,y) \sim \mathbb{P}_{\text{data}}} [d(\Phi(x|\Theta), y)]$$
 (10)

indirectly by minimizing an empirical loss defined by

$$L(\Theta|D) := \frac{1}{M} \sum_{m=1}^{M} d(\Phi(x_m|\Theta), y_m)$$
 (11)

given a finite number of M measurements drawn from the same data generating distribution

capacity, generalization, overfitting, ...

#### From Gradient Descent to Stochastic Gradient Descent

• The GD iteration in (9) on (11) would read as

$$\Theta^{k+1} = \Theta^k - \eta^k \frac{1}{M} \sum_{m=1}^M \nabla_{\Theta} d(\Phi(x_m | \Theta^k), y_m)$$
 (12)

- This iteration scheme is sometimes called batch gradient descent (personal opinion: wrongly)
- Drawbacks: computationally intensive (gradient for every sample), large memory requirements, cannot update model on the fly, . . .
- Idea: random partition of the dataset and obtain gradient steps on these random subsets → Stochastic Gradient Descent (SGD)

# Stochastic Gradient Descent

The steepest descent is decomposed into steps. Choose B < M. Choose a random permutation of the dataset  $\pi: \{1, \dots, M\} \to \{1, \dots, M\}$ .

- **1** Random permutation of the dataset into  $\pi^k$
- **2** Loop over each resulting subset  $i = 1, ..., \lceil M/B \rceil$  of size B:  $\{x_{\pi_m}, y_{\pi_m}\}_{m=(i-1)B}^{\min(iB,M)}$
- **1** Update the parameters over each subset  $i = 1, ..., \lceil M/B \rceil$

$$\Theta^{k,i+1} = \Theta^{k,i} - \eta^k \frac{1}{B} \sum_{m=(i-1)B}^{\min(iB,M)} d(\Phi(x_m | \Theta^{k,i}), y_m)$$
 (13)

**1** The k+1'th iteration step receives  $\Theta^{k+1} = \Theta^{k+1,1} := \Theta^{k,\lceil M/B \rceil}$ 

Terminology: k: epoch, i: batch, B: batch size

#### Batch size

- B = 1 is often called **online learning**
- Larger Bs provide more accurate estimates of the gradient  $\nabla_{\Theta} L(\Theta|D)$  but with diminishing returns
- Small batches often have a *regularization effect* and thus generalize better
- Small batches require more iterations per epoch and are thus slower
- Eventually: trial and error...

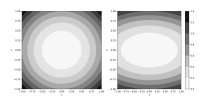
# Initialization and convergence criteria

Any implementable iterative optimization scheme requires two more ingredients

- An **initialization**  $\Theta^0$  determining the sequence  $\Theta^k \leftarrow \Theta^{k-1}, k > 1$ 
  - biases are usually initialized with an arbitrarily chosen constant value – most often 0
  - weights are initialized randomly, several options exist:  $W_{ij}^{(I)} \sim U\left(-\sqrt{\frac{6}{p_{l-1}+p_l}},\sqrt{\frac{6}{p_{l-1}+p_l}}\right),~W_{ij}^{(I)} \sim \mathcal{N}\left(0,\sqrt{\frac{2}{p_{l-1}+p_l}}\right)$
  - usually designed in a way to fix the variances of each layer's output activations, but also the variances of the gradients
  - crucial step for any successful iteration (no local optima, ...), actively researched topic
- A **convergence criterion** which stops the iteration at a given step K concluding in approximations  $\widehat{\Theta} := \Theta^K$  based on validation loss, maximum iteration number, . . .

## Learning rate

- The size of an iteration step in (9)–(13) in the *steepest* direction depends on the **learning rate**
- Closer to the minimum, gradients tend to decrease; smaller steps can thus lead to a better approximation of the minimum
- Several strategies  $k \to \eta^k$ : constant, adaptive (decay, validation loss linked, etc.), line search, . . .
- Eventually: trial and error
- Momentum based estimates: momentum SGD, Nesterov, ...
- Alternative adaptive optimization methods: Adam, AdaGrad, RMSProp, . . .



#### Seminar Series Neural Networks for Finance

Automatic differentiation: descending into a computer

# **Topic**

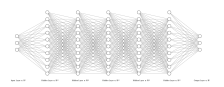
- We have seen: gradient based iterative optimization methods to minimize empirical loss
- However, they by definition rely on being able to calculate the **gradient** of the loss with respect to the parameters in the model  $\nabla_{\Theta} L(\Theta|D)$
- If d(f,g) in (11) is the mean-squared error we, e.g., have  $\nabla_{\Theta} L(\Theta|D) = 2(\nabla_{\Theta} \Phi(x|\Theta))^T (\Phi(x|\Theta) y)$
- $\Theta \in \mathbb{R}^p$  is a very high-dimensional vector (huge parameter spaces) (6)

#### Question

How can such a large dimensional gradient can be computed in a **precise**, **efficient**, and **robust** way?

Finite differences: high errors, slow, bad scaling; symbolic expressions: memory consumption, very long and redundant expressions — automatic differentiation

### Recall: neural networks' hierarchical structure



- Hierarchical sequence of compositions of the form (7)
- Complete parameter space  $\Theta := (W^{(1)}, b^{(1)}, \dots, W^{(L+1)}, b^{(L+1)}) \in \mathbb{R}^p$  with  $W^{(I)} \in \mathbb{R}^{p_I \times p_{I-1}}, b^{(I)} \in \mathbb{R}^{p_I}$  and  $p_0 = d$
- Activations by non-linear mappings  $\varphi^{(l)}$  and their affine combinations

$$z^{(l+1)} = W^{(l+1)}z^{(l)} + b^{(l+1)}, \quad a^{(l+1)} = \varphi^{(l+1)}(z^{(l+1)}),$$

$$l = 0, \dots, L$$

# Computational graphs

This structure

$$x \stackrel{W^{(1)},b^{(1)}}{\longmapsto} z^{(1)} \stackrel{\varphi^{(1)}}{\longmapsto} a^{(1)} \mapsto \dots \stackrel{\varphi^{(L+1)}}{\longmapsto} a^{(L+1)}$$

can be represented as **directed**, **acyclic graph**  $\longrightarrow$  **computational graph** 

- Each node in the graph corresponds to an operation and *data* flows through the vertices
- An operation is characterized by three characteristics: a compute function which determines the node's output; set of input nodes; and set of output nodes
- Computational graphs: complex calculations decomposed into a sequence of elementary operations; far more general than machine learning

# Forward propagation

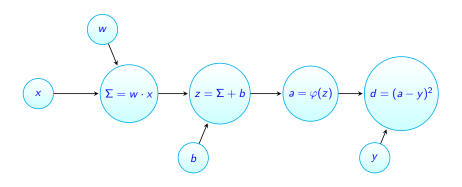


Figure: Forward Propagation - with mean-squared loss

### **Gradients**

• The input nodes of  $z^{(l)}$  are  $a^{(l-1)}$ ,  $W^{(l)}$ ,  $b^{(l)}$ 

$$z_i^{(l)} = \sum_{j=1}^{p_{l-1}} W_{ij}^{(l)} a_j^{(l-1)} + b_i^{(l)}$$
 (14)

With respect to these inputs are easily obtained

$$\frac{\partial z_i^{(l)}}{\partial a_j^{(l-1)}} = W_{ij}^{(l)}, \quad \frac{\partial z_i^{(l)}}{\partial b_j^{(l)}} = \delta_{ij}, \quad \frac{\partial z_i^{(l)}}{\partial W_{kj}^{(l)}} = \delta_{ki} a_j^{(l-1)}$$
(15)

• The input nodes of  $a^{(I)}$  are in  $z^{(I)}$ :  $a_i^{(I)} = \varphi_i^{(I)}(z^{(I)})$ 

$$\frac{\partial a_i^{(l)}}{\partial z_i^{(l)}} = (\nabla_{z^{(l)}} \varphi^{(l)}(z^{(l)}))_{ij}$$
(16)

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#### Derivation

With Jacobian matrices

Forward 
$$\begin{cases} a^{(0)} &= x, \\ z^{(1)} &= W^{(1)} \cdot a^{(0)} + \mathbf{b}^{(1)}, \\ a^{(1)} &= \varphi^{(1)}(z^{(1)}), \\ \dots &\\ z(L+1) &= W^{(L+1)} \cdot a^{(L)} + \mathbf{b}^{(L+1)}, \\ a^{(L+1)} &= \varphi^{(L+1)}(z^{(L+1)}) \end{cases}$$

$$\begin{cases} \nabla_x a^{(0)} &= I_d, \\ \nabla_{a^{(0)}} z^{(1)} &= W^{(1)}, \\ \nabla_{z^{(1)}} a^{(1)} &= \nabla_{z^{(1)}} \varphi^{(1)}(z^{(1)}), \\ \dots &\\ \nabla_{a^{(L)}} z^{(L+1)} &= W^{(L+1)}, \\ \nabla_{z^{(L+1)}} a^{(L+1)} &= \nabla_{z^{(L+1)}} \varphi^{(L+1)}(z^{(L+1)}) \end{cases}$$

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## **Back-propagation**

- Suppose there is one node in the last layer, matrix  $W^{(L+1)} = [w_{1j}^{(L+1)}, j = 1, \dots, p_L]$  is a vector then
- We compute the sensitivity of the mean-squared distance function d with respect to weight  $w_{1i}^{(L+1)}$  by the **chain rule**

$$\frac{\partial d}{\partial W_{1j}^{(L+1)}} = \frac{\partial d}{\partial a_1^{(L+1)}} \frac{\partial a_1^{(L+1)}}{\partial W_{1j}^{(L+1)}}$$

$$= \frac{\partial d}{\partial a_1^{(L+1)}} \frac{\partial a_1^{(L+1)}}{\partial z_1^{(L+1)}} \frac{\partial z_1^{(L+1)}}{\partial W_{1j}^{(L+1)}}$$

$$= \frac{\partial d}{\partial a_1^{(L+1)}} \varphi^{(L+1)'}(z_1^{(L+1)}) a_j^{(L)}$$

from Eq. (15)

• If d equals the mean-squared loss we obtain  $\frac{\partial d}{\partial W_{1i}^{(L+1)}} = 2(a_1^{(L+1)} - y)\varphi^{(L+1)'}(z_1^{(L+1)})a_j^{(L)}$ 

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#### Gradient

 Similarly, the chain rule gives an expression for the derivative of L wrt the inputs x

$$(\nabla_{x}d)^{T} = (\nabla_{a^{(L+1)}}d)^{T}\nabla_{z^{(L+1)}}a^{(L+1)}\nabla_{a^{(L)}}z^{(L+1)}\dots$$

$$\dots \nabla_{z^{(1)}}a^{(1)}\nabla_{a^{(0)}}z^{(1)}\nabla_{x}a^{(0)}$$

$$= (\nabla_{a^{(L+1)}}d)^{T}\nabla_{z^{(L+1)}}\varphi^{(L+1)}(z^{(L+1)})W^{(L+1)}\dots$$

$$\dots \nabla_{z^{(1)}}\varphi^{(1)}(z^{(1)})W^{(1)}I$$
(18)

where each element is a Jacobian matrix

• Three types of terms:, the gradient of the distance with respect to the outputs  $\nabla_{a^{(L+1)}}d$ ; Jacobian matrix of each layer's output activations  $\nabla_{z^{(l)}}\varphi^{(l)}(z^{(l)})$ ; and the derivative of the affine transformation function with respect to input activations  $\nabla_{a^{(l-1)}}z^{(l)}$ 

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# Backpropagation

To derive the closed-form expressions of backward propagation, we define the auxiliary error vector

$$[\delta^{(l)}]^{T} := [\nabla_{a^{(l+1)}} d]^{T} \nabla_{z^{(l+1)}} \varphi^{(l+1)} (z^{(l+1)}) W^{(l+1)} \dots$$

$$\dots W^{(l+1)} \nabla_{z^{(l)}} \varphi^{(l)} (z^{(l)}),$$
(19)

which is a vector of length  $\mathbb{R}^{p_l}$ 

### Backpropagation

The backpropagation scheme, implementing the derivative of the loss function is then defined by the following recursive expressions

$$\nabla_{b^{(l)}} L = \delta^{(l)},$$

$$\nabla_{W^{(l)}} L = \delta^{(l)} (z^{(l-1)})^{T},$$

$$[\delta^{(l)}]^{T} = [\delta^{(l+1)}]^{T} (W^{(l+1)}) \nabla_{z^{(l)}} \varphi^{(l)} (z^{(l)}).$$
(20)

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## Backward propagation

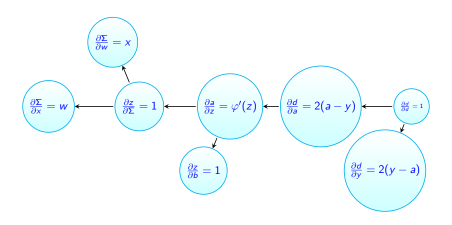


Figure: Backpropagation - with mean-squared loss

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Gradient based optimization problems

# Challenges: local minima, saddle points, etc.

Together with backpropagation/automatic differentiation, SGD is a fully-implementable iterative optimization algorithm – their combination is called **training**However, not without challenges

- local minima
- saddle points
- cliffs and plateaus: exploding and vanishing gradients
- ill-conditioned problems
- inaccurate gradient estimates over batches
- regularization: overfitting
- ...

### Local minima

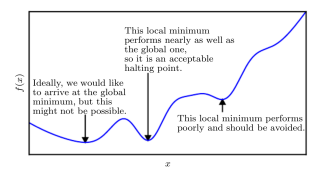


Figure: Goodfellow et al.: Deep Learning (2016), MIT Press [Figure 4.3, pg. 83]

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# Saddle points

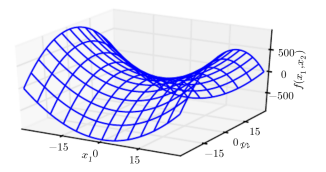


Figure: Goodfellow et al.: Deep Learning (2016), MIT Press [Figure 4.5, pg. 88]

# Vanishing gradients, plateaus, cliffs, etc.

- The gradient in the current layer is equal to a product of terms of all the later layers
- Vanishing gradient problem when one term is close to zero; gradient explosion when the product approaches infinity (numerical instabilities, rounding errors)

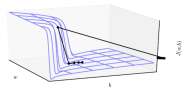


Figure: Goodfellow et al.: Deep Learning (2016), MIT Press [Figure 8.3, pg. 285]

Remedies: **ReLU** – unbounded from above; renormalization; gradient clipping, etc.

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# Advanced training techniques

#### Without the sake of completeness

- Input normalization
- Batch normalization: reparametrization, shifts and renormalizes activations within a layer across batches, parametrizes appropriate the appropriate means and variances
- Dropout is technique, which randomly turns off certain neurons (i.e., the gradient becomes zero for those neurons) to force the ANN to learn more features rather than saturate
- Constrained optimization: regularization with  $\mathcal{L}^1 \mathcal{L}^2$  penalties, etc.

• . . .

# Curvature, adaptive and second-order optimization

 $\bullet$  Momentum: descent modeled as a particle's momentum with unit mass; previous updates' gradients decay with some hyperparameter  $\alpha$ 

$$v^{k+1} = \alpha v^k - \eta^k \nabla_{\Theta} L(\Theta^k | D)$$

$$\Theta^{k+1} = \Theta^k + v^{k+1}$$
(21)

• Second-order optimization schemes: **Newton's method** to approximate the root of a multivariate function  $g: \mathbb{R}^n \to \mathbb{R}^n$  by

$$z^{0} \in \mathbb{R}^{n}, \quad z^{k+1} = z^{k} - [\nabla_{z}g(z^{k})]^{-1}g(z^{k}),$$
 (22)

where  $\nabla_z g(z)$  is the Jacobian matrix of the vector-valued mapping g defined by  $[\nabla_z g(z)]_{ii} := \partial_{z_i} g_i(z)$ 

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# Curvature, adaptive and second-order optimization

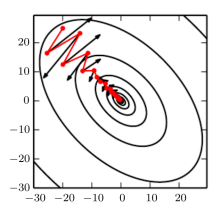


Figure: Goodfellow et al.: Deep Learning (2016), MIT Press [Figure 8.5, pg. 293]

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## Seminar Series Neural Networks for Finance

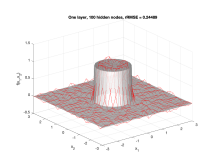
Deep versus Shallow

## Deep versus Shallow

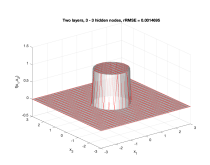
#### Questions

- When does it make sense to go deep?
- Why does it make sense to go deep?
- Poggio, T., Mhaskar, H., Rosasco, L., Miranda, B., & Liao, Q. (2017). Why and when can deep – but not shallow – networks avoid the curse of dimensionality: A review. arXiv:1611.00740 [cs]
- Mhaskar, H., Liao, Q., & Poggio, T. (2017). When and why are deep networks better than shallow ones? [Number: 1].
   Proceedings of the AAAI Conference on Artificial Intelligence, 31(1)
- Mhaskar, H., & Poggio, T. (2016). Deep vs. shallow networks
   : An approximation theory perspective. arXiv:1608.03287 [cs, math]

# Empirical difference



RMSE = 0.2449 **1 hidden layer** with 302 neurons
Total number of neurons: 302



RMSE = 0.0014 **3 hidden layers** with 3 neurons
Total number of neurons: 25

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# Degree of Approximation

- ullet Complexity: number of parameters  $\sim$  of neurons/units in a network
- Let  $\mathcal{V}_N \coloneqq \{\Phi(\cdot|\Theta): \Theta \in \mathbb{R}^N\} \subseteq \mathcal{G}$  be the set of networks with complexity N,  $\mathcal{V}_N \subseteq \mathcal{V}_{N+1}$
- Degree of Approximation is defined by

$$\mathsf{dist}(f,\mathcal{V}_N) \coloneqq \inf_{\Phi \in \mathcal{V}_N} ||f - \Phi||_X,$$

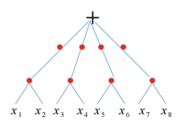
for  $f \in \mathcal{X}$ 

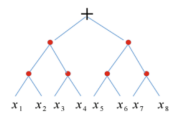
• dist $(f, \mathcal{V}_N) = \mathcal{O}(N^{-\gamma}) \implies$  a network with complexity  $N = \mathcal{O}(\epsilon^{-\frac{1}{\gamma}})$  is sufficient to guarantee accuracy at least  $\epsilon$ 

# **Compositional Functions**

We will consider hierarchical compositions of functions, such as

$$f(x_1,...,x_8) = h_3 \Big( h_{21}(h_{11}(x_1,x_2),h_{12}(x_3,x_4)), h_{22}(h_{13}(x_5,x_6),h_{14}(x_7,x_8)) \Big)$$





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# Compositional Functions' Class

• Let  $I^n = [-1, 1]^n$  and  $C(I^n)$  be the space of continuous functions on  $I^n$  with norm

$$||f|| = \max_{x \in I^n} |f(x)|$$

• Then consider the subspace  $W_r^n \subset C(I^n)$  consisting of r times continuously differentiable functions on  $I^n$ , where

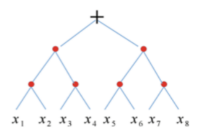
$$||f|| + \sum_{1 \le |k|_1 \le r} ||D^k f|| \le 1,$$

where k is a multiindex and  $D^k$  denotes the partial derivative indicated by k, and  $|k|_1$  is the sum of the multiindex

# Compositional Functions - Function Class

- Let  $W_r^{n,2}$  be the class of functions with a hierarchical structure with each constituent function,  $h_{ij} \in W_r^2$ 
  - Binary tree functions
- Note  $\mathcal{W}_r^{n,2} \subset \mathcal{W}_r^n$
- Example of a  $W_r^{4,2}$  function:

$$f(x_1, x_2, x_3, x_4) = \underbrace{h_2}_{\in \mathcal{W}_r^2} \underbrace{(h_{11}}_{\in \mathcal{W}_r^2} (x_1, x_2), \underbrace{h_{12}}_{\in \mathcal{W}_r^2} (x_3, x_4))$$



### Shallow Networks

- Shallow network: One hidden layer
  - Let  $S_{N,n}$  be the class of shallow networks with N neurons and n-dimensional input of the form

$$x\mapsto \sum_{k=1}^N a_k\sigma(w_k\cdot x+b_k),\quad w_k\in\mathbb{R}^n,\quad a_k,b_k\in\mathbb{R},$$

- where  $\sigma: \mathbb{R} \to \mathbb{R}$  is the (nonlinear) activation function.
- Number of trainable parameters:  $p = (n+2)N \sim N$
- Deep neural networks: at least two hidden layers
  - Let  $\mathcal{D}_{N,2}$  be the class of Deep networks that take n-dimensional input with N neurons and binary tree structure
  - ullet Each constituent function in the deep network is a  $\mathcal{S}_{Q,2}$  network
  - Complexity: (n-1)Q = N

# Comparison theorems

#### Theorem - Shallow neural networks

Under some assumptions on the non-linear activations, for any  $f \in \mathcal{W}_r^n$  the complexity of shallow networks that provide accuracy at least  $\epsilon$  is  $N = \mathcal{O}(\epsilon^{-n/r})$  and is the best possible  $\Longrightarrow$  complexity increases **exponentially** with the input dimension - curse of dimensionality

### Theorem – Deep neural networks

Under some assumptions on the non-linear activations, for any  $f \in \mathcal{W}_r^{n,2}$  the complexity of a deep network (with the same compositional achitecture) that provide accuracy at least  $\epsilon$  is  $N = \mathcal{O}((n-1)\epsilon^{-2/r})$  and is the best possible  $\implies$  The complexity does NOT increase exponentially with the input dimension!

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# Why is this an interesting result?

Many functions have an underlying hierarchical composition.
 Example:

$$f(x_1, x_2, x_3, x_4) = ac^2 x_4^4 x_1^3 + 2acx_1^3 x_4^2 x_3^3 + ax_1^3 x_3^6 + bc^2 x_1 x_2 x_4^4 + 2bcx_1 x_2 x_4^4 x_3^3 + bx_1 x_2 x_3^6 = (ax_1^3 + bx_2 x_1)(x_3^3 + cx_4^2)^2 = h(h_{11}(x_1, x_2), h_{12}(x_3, x_4))$$

where

$$h(x,y) = xy^2$$
,  $h_{11}(x,y) = ax^3 + byx$ ,  $h_{12}(x,y) = x^3 + cy^2$ 

instead of approximating a 9th degree polynomial, a deep network approximates 3 polynomials of 3rd degree

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