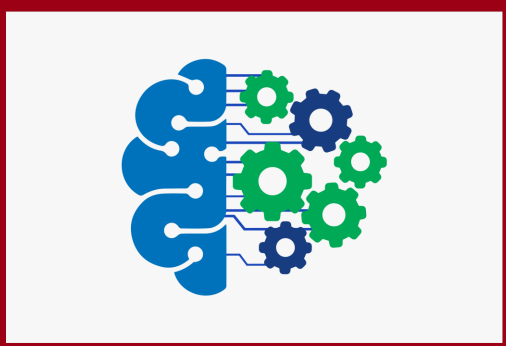




Università degli Studi di Padova



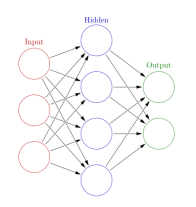
Neural Networks

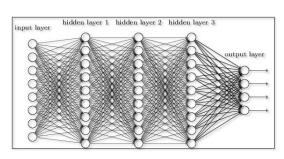
Machine Learning 2023-24
UML book chapter 20

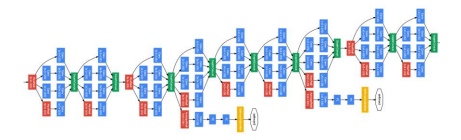
Slides P. Zanuttigh (some material from F. Vandin slides)



Artificial Neural Networks







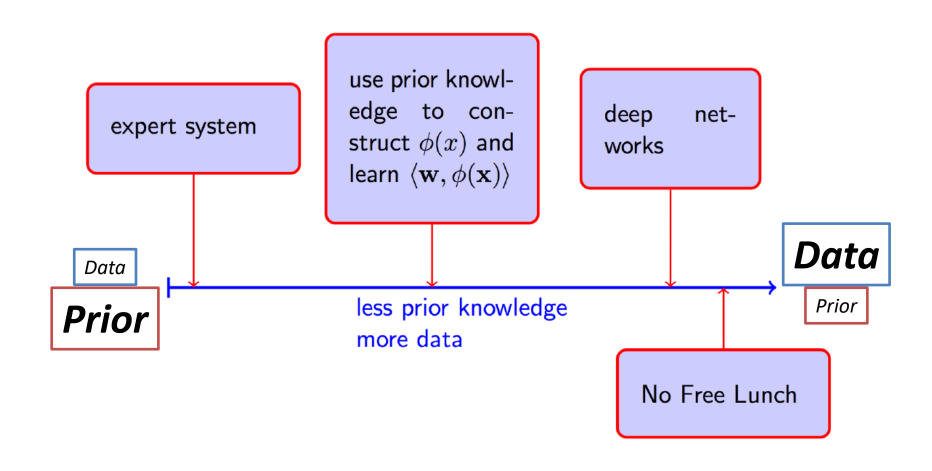
- Model of computation inspired by the structure of neural networks in the brain
- Large number of basic computing devices (neurons) connected to each other
- Neural Networks (NN) are represented with directed graphs where the nodes are the neurons, and the edges corresponds to the links between the neurons

Brief History:

- Firstly proposed in 1940-50
- First practical applications in the 80-90s but practical results were lower than SVM, Random Forests and other techniques
- From 2010 on deep architectures with impressive performances

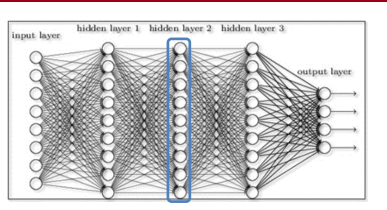


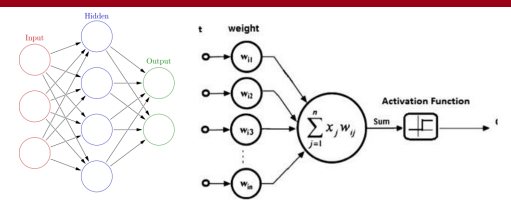
From Simple Predictors to Deep Learning





Feedforward Neural Networks





Feedforward network: the graph representing the network has no cycles (data flows only in one direction)

The network is typically organized into layers: each neuron takes in input only the output of neurons in the previous layer

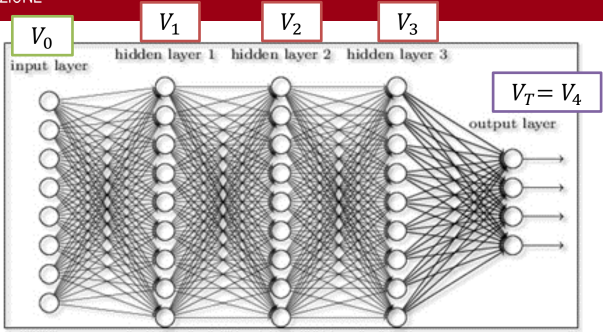
Notation (NN): Graph G=(V,E) and function $w:E\to\mathbb{R}$

- V: neurons (|V| is the size of the network)
- E: connections between neurons (directed edges)
- $w: E \to \mathbb{R}$ weight function over the edges (the weights w are the parameters to be learned)

Each neuron:

- 1. Takes in input the sum of the outputs of the connected neurons from previous layer weighted by the edge weights (w)
- 2. Applies to the result a simple scalar function (activation function, σ)

Notation (1)

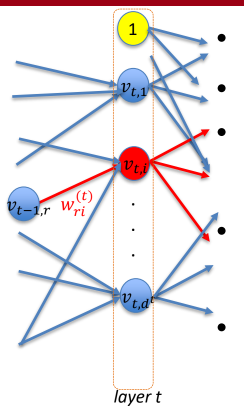


Represent a network as the union of a set of (disjoint) layers: $V = \bigcup_{t=0}^{T} V_t$

- V_t , t = 0, ..., T: t-th layer,
- $d^t + 1$ number of nodes of layer t
 - "+1": constant neuron (avoid bias, incorporate as in homogenous coord.)
- V_0 : input layer, V_T : output layer, $V_1, ..., V_{T-1}$ inner (hidden) layers
- T : depth of the network
 - T=2 in "classic" NN, T>>2 in deep networks



Notation (2)



 $v_{t,i}$: *i-th* neuron in the *t-th* layer

$$oldsymbol{v}^{(t)} = \left(1, v_{t,1}, \dots, v_{t,d^t}\right)^T$$
 : all neurons of layer $oldsymbol{t}$

Weights $w_{rj}^{(t+1)} = w(v_{t,r}, v_{t+1,j})$: weight of arc from neuron r of layer t to neuron j of layer t+1

$$\mathbf{w}_{j}^{(t)} = \left(w_{0j}^{(t)}, \dots, w_{d^{(t-1)}j}^{(t)}\right)^{T}$$
: all weights of arcs in input to neuron j of layer t (notice: from layer t -1 to t) $w^{(t)}$: matrix of weights of all arcs incoming to layer t

$$w^{(t)} = \begin{bmatrix} w_{01}^{(t)} & w_{02}^{(t)} & \cdots & w_{0d^t}^{(t)} \\ w_{11}^{(t)} & w_{12}^{(t)} & \cdots & w_{1d^t}^{(t)} \\ \vdots & \vdots & \ddots & \vdots \\ w_{d^{(t-1)}1}^{(t)} & w_{d^{(t-1)}2}^{(t)} & \cdots & w_{d^{(t-1)}d^{(t)}}^{(t)} \end{bmatrix}$$



How a Neuron Works

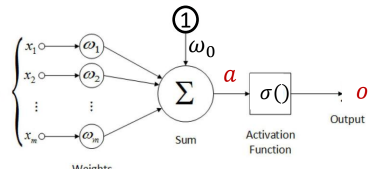
Compute output $o_{t,i}(x)$ of the *i-th* neuron in the *t-th* layer when x is fed to the network

- Compact notation: use $v_{t,i}$ also to represent the output of the neuron
- The output of a neuron is a non-linear (activation) function applied to the linear combination of the inputs coming from the previous layer
 - σ : non-linear activation function
 - $a_{t+1,j} = \langle w_i^{(t+1)}, v^{(t)} \rangle$ output of neuron before the activation function

$$o_{t+1,j}(x) = \sigma \left(\sum_{r: (v_{t,r}, v_{t+1,j}) \in E} w(v_{t,r}, v_{t+1,j}) o_{t,r}(x) \right) = \sigma \left(a_{t+1,j}(x) \right)$$

In vector notation:

In vector notation:
$$v_{t+1,j} = \sigma\left(< \boldsymbol{w}_{j}^{(t+1)}, \boldsymbol{v}^{(t)}>\right) = \sigma(a_{t+1,j})$$
 Inputs
$$v_{t+1,j} = \sigma\left(< \boldsymbol{w}_{j}^{(t+1)}, \boldsymbol{v}^{(t)}>\right) = \sigma(a_{t+1,j})$$
 Natisher.





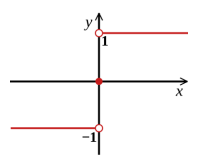
Activation Functions

Various activation functions $\sigma(a)$ can be exploited:

- Sign function
- Threshold function
- Sigmoid function
- 4. Hyperbolic Tangent
- 5. Rectified Linear Unit



Activation: Sign and Threshold

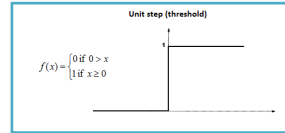


Outputs the sign of the input

$$\sigma(a) = sign(a)$$

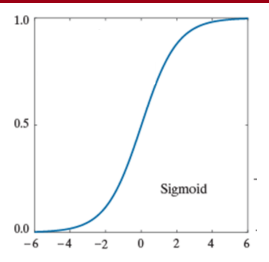
$$\mathbb{R} \to [-1; 1]$$

- + Simple/fast
- + Nice interpretation as the firing rate of a neuron
 - -1 = not firing
 - 1 = firing
- Output is not smooth/continuous
- saturate and kill gradients, thus NN will barely learn



Threshold function: similar behaviour

Activation: Sigmoid



Takes a real-valued number and "squashes" it into range between 0 and 1

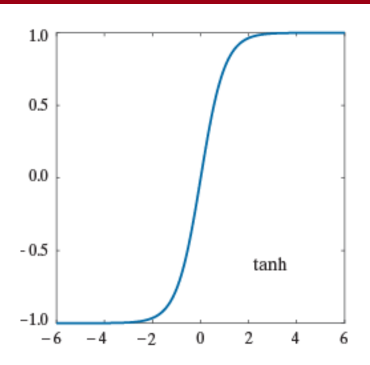
$$\sigma(a) = \frac{1}{1 + e^{-a}}$$

$$\sigma'(a) = \sigma(a)[1 - \sigma(a)]$$

$$\mathbb{R} \to [0,1]$$

- + Smooth output
- + Nice interpretation as the firing rate of a neuron
 - 0 = not firing at all
 - 1 = fully firing
- Sigmoid neurons saturate and kill gradients, thus NN will have issues in learning
 - when the neuron's activation are 0 or 1 (saturate)
 - (x) gradient at these regions almost zero
 - almost no signal will flow to its weights
 - if initial weights are too large then most neurons would saturate

Activation: Tanh



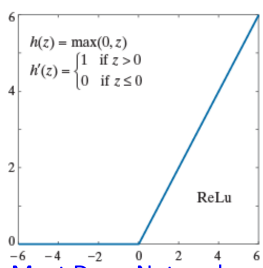
Takes a real-valued number and "squashes" it into range between -1 and 1

$$\sigma(a) = \tanh(a) = \frac{e^{a} - e^{-a}}{e^{a} + e^{-a}} = \frac{e^{2a} - 1}{e^{2a} + 1}$$
$$\sigma'(a) = 1 - [\tanh(a)^{2}]$$
$$\mathbb{R} \to [0,1]$$

Tanh is a scaled and shifted sigmoid: tanh(x) = 2sigm(2x) - 1

- + Like sigmoid, tanh neurons saturate
- Unlike sigmoid, output is zero-centered

Activation: ReLU



Takes a real-valued number and thresholds it at zero

$$\sigma(a) = \max(0, a) = \begin{cases} a & \text{if } a > 0 \\ 0 & \text{if } a \le 0 \end{cases}$$
$$\sigma'(a) = \begin{cases} 1 & \text{if } a > 0 \\ 0 & \text{if } a \le 0 \end{cases}$$
$$\mathbb{R} \to \mathbb{R}_+$$

Most Deep Networks use ReLU nowadays

- + Trains much faster
 - accelerates the convergence of SGD
 - due to linear, non-saturating form
- + Less expensive operations
 - compared to sigmoid/tanh (exponentials etc.)
 - implemented by simply thresholding a matrix at zero
- + More expressive
- + Prevents the gradient vanishing problem



Forward Propagation

$$v^{(0)} \rightarrow v^{(1)} \rightarrow v^{(2)} \rightarrow \cdots \rightarrow v^{(T)}$$

Take an input sample and compute the output of the network

Start from the input (layer 0)...compute the output of layer 1, send to layer 2 and get output....

.... through all the layers up to the output layer

```
Input: \mathbf{x} = (x_1, \dots, x_d)^T; NN with 1 output node

Output: prediction y of NN;

v^{(0)} \leftarrow (1, x_1, \dots, x_d)^T;

for t \leftarrow 1 to T do

From first to last layer

a^{(t)} \leftarrow (w^{(t)})^T v^{(t-1)};

v^{(t)} \leftarrow (1, \sigma(a^{(t)})^T)^T;

Non-linear activation

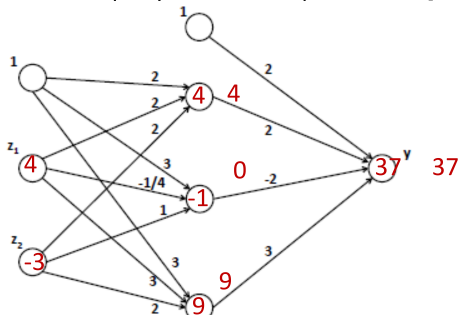
Function («1» for bias)
```

Example

Consider the neural network in the figure and assume the activation function $\sigma(x)$ is the Rectified Linear Unit (ReLU):

$$\sigma(a) = \begin{cases} a & a > 0 \\ 0 & a \le 0 \end{cases}$$

Compute the value of the output y when the input z is z = [4 -3]



Learning Neural Networks

Neural Network (NN): (V, E, σ, w)

- Corresponds to a function $h_{V,E,\sigma,w}\colon \mathbb{R}^{|V_0-1|} o \mathbb{R}^{|V_T|}$
- The hypothesis class of a network is defined by fixing its architecture:

$$\mathcal{H}_{V,E,\sigma} = \{h_{V,E,\sigma,w} : w \text{ is a mapping from } E \text{ to } \mathbb{R}\}$$

- V, E, σ defines the architecture of the network
- w contains the parameters that are going to be learned
- Training of the NN: finding the optimal set of weights w



Expressive Power of NN (boolean functions)

Theorem 1:

For every d, there exist a graph (V,E) of depth 2 such that $\mathcal{H}_{V,E,sign}$ contains all functions from $\{-1,1\}^d$ to $\{-1,1\}$

- NN can implement any boolean function
- Recall: Boolean functions include any function that can be implemented in a computer

But... the graph (V,E) is very big!

Theorem 2:

For every d, let s(d) be the minimal integer such that there exists a graph (V,E) with |V| = s(d) such that $\mathcal{H}_{V,E,sign}$ contains all functions from $\{-1,1\}^d$ to $\{-1,1\}$. Then s(d) is an exponential function of d

Note: for the sigmoid activation function similar results



Expressive Power of NN (demonstration)

Consider sign activation $\mathcal{H}_{V,E,sign}$

1. Use this 3 layers NN:
$$\begin{cases} INPUT\colon |V_0|=n+1\\ HIDDEN\colon |V_1|=2^n+1\\ OUTPUT\colon |V_2|=1 \end{cases}$$

- 2. Define: $u_1, \dots, u_k \in \{\pm 1\}^n$: all input vectors leading to an output of 1
- 3. Notice: $\langle x, u_i \rangle = \begin{cases} n & \text{if } x = u_i \\ \leq n 2 \text{ if } x \neq u_i \end{cases}$ (all bits match)
- 4. Define $g_i = sign(\langle \mathbf{x}, \mathbf{u_i} \rangle n + 1) = \begin{cases} 1 & if \mathbf{x} = \mathbf{u_i} \\ -1 & otherwise \end{cases}$
- 5. Adapt weights w to get g_i in the hidden layer \rightarrow hidden layer neuron $v_{1,i}$ looks if the input is u_i
- 6. Output layer: $f(x) = sign(\sum_{i=1}^{k} g_i(x) + k 1)$ (sign is 1 if at least one is true)

Notice: the network is exponentially large, it works but it is a «brute-force» solution probably leading to overfitting



Expressive Power of NN (real valued functions)

Proposition

For every fixed $\varepsilon > 0$ and every Lipschitz function $f: [-1,1]^d \to [-1,1]$ it is possible to construct a neural network such that for every input $\mathbf{x} \in [-1,1]^d$ the output of the neural network is in $[f(\mathbf{x}) - \varepsilon, f(\mathbf{x}) + \varepsilon]$.

Not part of the course **Note:** first result proved by Cybenko (1989) for sigmoid activation function, requires only 1 hidden layer!

NNs are universal approximators!

But again...

Proposition

Fix some $\varepsilon \in (0,1)$. For every d, let s(d) be the minimal integer such that there exists a graph (V, E) with |V| = s(d) such that $\mathcal{H}_{V,E,\sigma}$, with $\sigma =$ sigmoid, can approximate, with precision ε , every 1-Lipschitz function $f: [-1,1]^d \to [-1,1]$. Then s(d) is exponential in d.



Implement Conjunction and Disjunction with NN

- Neural Networks can implement boolean AND / OR
- \Box Consider sign activation and k inputs with values ± 1

Conjunction (AND)

$$f(\mathbf{x}) = sign\left(1 - k + \sum_{i=1}^{k} \mathbf{x}_i\right)$$

(positive if all positive, AND)

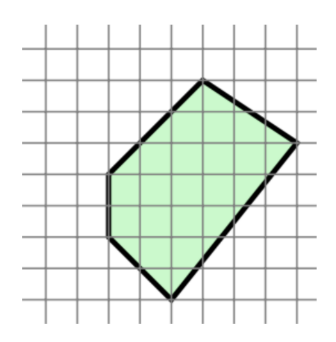
Disjunction (OR)

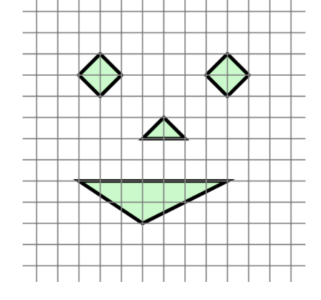
$$f(\mathbf{x}) = sign\left(k - 1 + \sum_{i=1}^{k} \mathbf{x_i}\right)$$

(positive if at least one positive, OR)



Expressive Power of NN (example)





- Input in \mathbb{R}^2 , $extstyle{2}$ -layers NN
- k neurons, sign activation
- Each neuron: an halfspace
- Intersection of halfspaces
- Convex polytopes with k-1 faces

- Input in \mathbb{R}^2 , 3-layers NN
- k neurons, sign activation
- Each neuron: an halfspace
- Intersection and unions of halfspaces
- Union of polytopes

VC dimension of NN

- With sign activation VC dimension of $\mathcal{H}_{V,E,sign} = O(|E|\log|E|)$ (no demonstration)
- With sigmoid (σ) activation VC dimension of $\mathcal{H}_{V.E.\sigma} = O(|V|^2|E|^2)$ (no demonstration)
- →Large NNs require a lot of data!

If we have enough data, what about the computation time?



Runtime of NN

Applying the ERM rule to a NN (V,E,σ,w) is computationally difficult, even for Not part of the course relatively small NN...

Theorem:

Hypothesis: Let $k \geq 3$. For every d, let (V,E) be a layered graph with d input nodes, k+1 nodes at the (only) hidden layer (where one of them is the constant neuron), and a single output node.

Thesis: It is NP-hard to implement the ERM rule with respect to $\mathcal{H}_{V.E.sign}$ (no demonstration)

Even approximations of ERM rule are infeasible Also by changing the activation things do not get better

Need a different strategy....

SGD and backpropagation algorithm!



Neural Networks Optimization

Target of ERM: given training data $(x_1, y_1), ..., (x_m, y_m)$ find the weights that minimize the training error:

$$L_s(h) = \frac{1}{m} \sum_{i=1}^{m} \ell(h, (\boldsymbol{x_i}, y_i))$$

The problem is challenging!

Idea:

- 1. Forward propagate the training data and compute the loss
- 2. Consider the loss as a function of the weights and compute the gradient of the loss w.r.t. the weights
- 3. Update the weights with SGD

Good Idea! But we need the gradient of the loss w.r.t. the weights



SGD fror Neural Networks: Algorithm

SGD for Neural Networks parameters: Number of iterations τ Step size sequence $\eta_1, \eta_2, ..., \eta_{\tau}$ Regularization parameter $\lambda > 0$ Input: Network : layered graph G=(V,E)differentiable activation function $\sigma: \mathbb{R} \to \mathbb{R}$ adaptive learning rate Algorithm: chose $\mathbf{w}^{[1]} \in \mathbb{R}^{|E|}$ at random (from a distribution s.t. $w^{[1]}$ is close enough to 0) for $s = 1, 2, ..., \tau$ sample $(x, y) \sim \mathcal{D}$ calculate gradient $v_s = backpropagation(x, y, w, (V, E), \sigma)$ update $w^{[s+1]} = w^{[s]} - \eta_s(v_s + \lambda w^{[s]})$ regularization output: $\overline{\boldsymbol{w}}$ is the best performing $\boldsymbol{w}^{[s]}$ on a validation set



SGD for NN:

How to Compute the Gradient

Update Rule (baseline version)
$$w_{ij}^{(t)[s+1]} = w_{ij}^{(t)[s]} - \eta_s \frac{\partial L_s}{\partial w_{ij}^{(t)[s]}}$$

learning rate

gradient of the loss w.r.t each single weight

$$\frac{\partial L_S}{\partial w} = \frac{\partial}{\partial w} \left(\frac{1}{m} \sum_{i=1}^m \ell(h, (\boldsymbol{x}_i, y_i)) \right) = \frac{1}{m} \sum_{i=1}^m \frac{\partial L(h, (\boldsymbol{x}_i, y_i))}{\partial w}$$

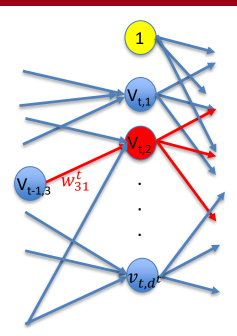
- We need the gradient w.r.t. each single weight in the network
- But we can compute the loss only on the output (i.e., after the last layer)
- Recall that each neuron contains also the non-linear activation function

$$\boldsymbol{\delta^{(t)}} = \frac{\partial L}{\partial \boldsymbol{a}^{(t)}} = \begin{bmatrix} \frac{\partial L}{\partial a_{t,1}} \\ \vdots \\ \frac{\partial L}{\partial a_{t,d^t}} \end{bmatrix}$$

 $\boldsymbol{\delta^{(t)}}$: change in error w.r.t. to the weighted average before the non-linear transformation



BackPropagation (1)

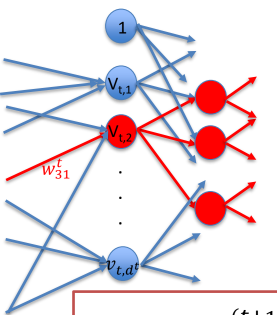


- Decompose the gradient with the chain rule
- Recall $a_{t,j} = \sum_{k=0}^{d^{(t-1)}} w_{kj}^{(t)} v_{t-1,k}$
- \circ Each weight $w_{ij}^{(t)}$ impacts only on $a_{t,j}$
- We need $\delta^{(t)} = \frac{\partial L}{\partial a^{(t)}}$ to compute the gradient
- \circ σ' depends on the selected activation function

$$\begin{split} \frac{\partial L}{\partial w_{ij}^{(t)}} &= \frac{\partial L}{\partial a_{t,j}} \frac{\partial a_{t,j}}{\partial w_{ij}^{(t)}} = \delta_j^{(t)} \frac{\partial}{\partial w_{ij}^{(t)}} \left(\sum_{k=0}^{d^{(t-1)}} w_{kj}^{(t)} v_{t-1,k} \right) = \delta_j^{(t)} v_{t-1,i} \\ \delta_j^{(t)} &= \frac{\partial L}{\partial v_{t,j}} \frac{\partial v_{t,j}}{\partial a_{t,j}} = \frac{\partial L}{\partial v_{t,j}} \sigma'(a_{t,j}) \end{split}$$



BackPropagation (2)



- \circ Understand how the loss changes w.r.t. $v_{t,j}$
- Change in layer t affects only neurons in layer t+1 (and then each following layer up to the loss at the end)
- Each neuron can affect all the neurons in next layer
- Need to sum contributions to all the neurons in layer t+1
- o To compute $\delta^{(t)}$ we need $\delta^{(t+1)}$ (solution of the next layer)

$$\frac{\partial L}{\partial v_{t,j}} = \sum_{k=1}^{d^{(t+1)}} \frac{\partial L}{\partial a_{t+1,k}} \frac{\partial a_{t+1,k}}{\partial v_{t,j}} = \sum_{k=1}^{d^{(t+1)}} w_{jk}^{(t+1)} \delta_k^{(t+1)}$$

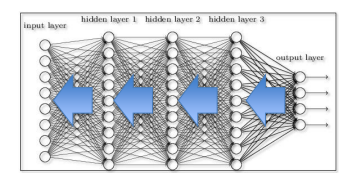
$$\delta_j^{(t)} = \frac{\partial L}{\partial v_{t,j}} \frac{\partial v_{t,j}}{\partial a_{t,j}} = \sigma'(a_{t,j}) \sum_{k=1}^{d^{(t+1)}} w_{jk}^{(t+1)} \delta_k^{(t+1)}$$

BackPropagation (3)

$$\delta_j^{(t)} = \sigma'(a_{t,j}) \sum_{k=1}^{d^{(t+1)}} w_{jk}^{(t+1)} \delta_k^{(t+1)}$$

- The solution for each layer need the solution of the following one
- o Start from the last layer ($\delta^{(L)}$ can be computed from the loss on the output)
- Backpropagate the gradients through all the layers up to the first

$$v^{(0)} \leftarrow v^{(1)} \leftarrow v^{(2)} \leftarrow \cdots \leftarrow v^{(T)}$$



BackPropagation: Algorithm

```
Input: data point (\mathbf{x}_i, y_i), NN (with weights w_{ij}^{(t)}, for 1 \leq t \leq T)

Output: \delta^{(t)} for t = 1, \ldots, T

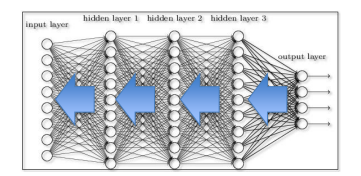
compute a^{(t)} and v^{(t)} for t = 1, \ldots, T;

\delta^{(T)} \leftarrow \frac{\partial L}{\partial a^{(T)}}

for t = T - 1 downto 1 do

\begin{bmatrix} \delta_j^{(\ell)} \leftarrow \sigma'(a_{t,j}) \cdot \sum_{k=1}^{d^{(\ell+1)}} w_{jk}^{(t+1)} \delta_k^{(t+1)} \text{ for all } j = 1, \ldots, d^{(t)}; \\ \text{return } \delta^{(1)}, \ldots, \delta^{(T)}; \end{bmatrix}
```

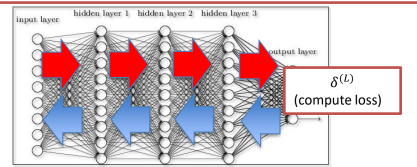
$$v^{(0)} \leftarrow v^{(1)} \leftarrow v^{(2)} \leftarrow \cdots \leftarrow v^{(T)}$$





NN Training: complete algorithm

```
BackPropagation algorithm with SGD
                            Input: training data (x_1, y_1), ..., (x_m, y_m)
                                      Output: NN weights w_{ij}^{(t)}
Initialize w_{i,i}^{(t)} \quad \forall i, j, t;
for s \leftarrow 0,1,2,... do
                                                                       // until convergence
            pick (x_k, y_k) at random from training data; // SGD: 1 sample at random
            compute v_{t,j} \forall j, t;
                                                                       // forward propagation
           compute \delta_i^{(t)} \quad \forall j, t;
                                                                       // backward propagation
           w_{ij}^{(t)[s+1]} = w_{ij}^{(t)[s]} - \eta v_{t-1,i} \delta_j^{(t)} \quad \forall i, j, t; \quad // \text{ update weights}
if converged then return w_{ij}^{(t)[s]} \forall i, j, t;
```





NN Training Details:

Pre-processing and Initialization

Pre-processing:

- Typically, all inputs are normalized and centred around 0
- Both local or global normalization strategies

Initialization of the weights

- All to 0 does not work
- Random values around 0 (regime where model is roughly linear)
- Uniform or normal (Gaussian) distribution can be used
- Sometimes multiple initializations and trainings, then select best result (smallest training error)
- In deep NN "Glorot" initialization: normal distribution with variance inversely proportional to the sum of the number of incoming and outcoming connections of the neuron



NN Training details: Convergence

When to stop?

- Small training error
- Small marginal improvement in error at each step
- Upper bound on number of iterations

Loss function usually has multiple local minima

- With highly dimensional spaces the risk is smaller than in low dimensional ones, but no guarantee
- Run stochastic gradient descent (SGD) from different (random) initial weights

Regularization

- Minimize weighted sum of the loss with the sum of all the weights
- Avoid too large weights and make optimization more stable
- Regularization parameter λ
- L1 or L2 regularization can be used

$$L_s(h) + \frac{\lambda}{m} \sum_{i,j,t} \left| w_{ij}^{(t)} \right|$$

$$L_s(h) + \frac{\lambda}{m} \sum_{i,j,t} \left(w_{ij}^{(t)} \right)^2$$