## Data mining Concepts & Algorithms

Introduction to deep learning

Flavio Giobergia

(slides from Large Language Models course)

## The perceptron

- The perceptron is the simplest unit of neural networks
- It takes an input with *multiple features*, and does the following:
  - It weights each input feature with a given *weight*,
  - It produces a weighted sum of the inputs, and
  - It applies a *function* to the output
- $y = f(w_0 + w_1x_1 + w_2x_2 + \dots + w_nx_n)$

### The perceptron

Politecnico D<sup>B</sup>G



Or, in other words,  $y = f(\sum_{i=0}^{n} w_i x_i) = f(w^T x)$  and  $x_0 = 1$ 

- $x = (x_1, x_2, ..., x_n)$  is the input sample
- *y* represents the output of the perceptron.
- $f(\cdot)$  represents a non-linear "activation" function
- *w<sub>i</sub>* (and *w<sub>0</sub>*) are weights (and bias), which are
   "learned"

#### Note

With the exception of  $f(\cdot)$ , this looks like the classic *linear regression* And if  $f(\cdot) = \sigma(\cdot)$  (sigmoid function), this looks like the (just as classic) *logistic regression*  Politecnico D<sup>B</sup>G

## The perceptron, in 2D





The perceptron can be used to represent a family of functions,  $y = w_1 x_1 + w_2 x_2 + w_0$ 

Various values of  $w_0, w_1, w_2$  define the different functions that can be learned by the perceptron.

## Activation functions

- Activation functions are used for two *main* reasons:
  - 1. Enforce *properties* on perceptron's output
  - E.g., sigmoid → binds output to [0, 1] range
     Introduce *non-linearities* in the model
     + some others (faster convergence, sparsity, ...)
- Commonly adopted functions:
  - ReLU

Politecnico D<sup>B</sup>G

- Sigmoid
- Leaky ReLU
- Tanh
- Softmax
- Linear
- GeLU



### 1. Enforce properties on perceptron's output

- Binary classification problem
  - Separate positive ( $\bigstar$ ) and negative ( $\bigcirc$ ) samples
- For a point  $x \in \mathbb{R}^2$ , the perceptron can predict  $p(\mathbf{k} \mid x)$ 
  - For the binary case, this implies  $p(\bigcirc |x) = 1 p(\cancel{x} | x)$
- To get a valid probability, we must enforce  $p(\mathbf{x}|x) \in [0, 1]$ 
  - We already have  $p(\bigcirc | x) + p(\oiint | x) = 1$  by construction
- The Sigmoid maps any value in  $\mathbb R$  to the range [0,1]
  - i.e., the perceptron's output (in  $\mathbb{R}$ ) is squashed to [0, 1]

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



-2 -

Politecnico DBG

### Adding some perceptrons



### and adding other layers!

Politecnico DBG



### 2. Introduce non-linearities in the model

- if f(x) = x (i.e., no non-linearity is added), we get  $z = s^{T}W^{T}x$
- This implies:
  - 1. We could have used W' = Ws and get the same output
  - 2. We wouldn't have needed a second layer!
  - 3. But our model is still linear
- So, we use non-linear activation functions to model more complex functions



# Multi-layer perceptron models

- We can stack additional *layers* 
  - separated by *non-linearities* (activation functions) to prevent collapses
- Universal Approximation Theorem tells us that we can approximate "any" function with MLPs
  - "For any continuous function *g* defined on a compact subset of  $\mathbb{R}^n$  and for any  $\epsilon > 0$ , there exists a feedforward neural network with a single hidden layer and a finite number of neurons that can approximate g to within an arbitrary degree of accuracy  $\epsilon''$
  - A single-layer MLP works ... but no information on the number of neurons, or the weights' values!
  - Deeper, narrower networks are generally used







### Activation functions for classification models

- As argued, activation functions can be used to enforce properties on the model's output
- In classification problems, the output *before* the final activation is treated as *unnormalized probabilities* (*logits*)
- We still need a step to convert *logits* into *valid* probabilities
  - i.e., all probabilities should sum to 1, and be in [0, 1]



# Binary classification

- The model predicts the probability of a single class for point x
  - As a convention, the *positive* one P(pos|x)
- The model produces a logit z = model(x)
- We use the *sigmoid function* on the output logit z

• 
$$\sigma(z) = \frac{1}{1+e^{-z}}$$

Politecnico DBG

- This guarantees  $P(pos|x) \in [0, 1]$
- We work out the probability of the negative class
  - P(neg|x) = 1 P(pos|x)
  - We can easily show that  $P(neg|x) \in [0,1]$
- By construction, P(pos|x) + P(neg|x) = 1



# Multi-class classification

- The output class is one of many  $(c_1, c_2, \dots, c_n)$
- The model produces *n* logits for a point *x* 
  - (i.e., the last layer will have *n* perceptrons)
  - $z = (z_1, z_2, ..., z_n) = model(x)$
- We need to obtain, from the logits, valid probabilities
  - $P(c_1|x), P(c_2|x), \dots, P(c_n|x)$
- The *softmax* function is applied:

• 
$$P(c_i|x) = \frac{e^{z_i}}{\sum_j e^{z_j}}$$

Politecnico D<sup>B</sup>G

- It can be easily shown that:
  - $P(c_i|x) \in [0,1]$   $\sum_i P(c_i|x) = 1$



### Activation functions for regression models

- In regression, models generally predict real numbers
- Typically, there is no need to enforce properties
- Output activation function can be the identity function
  - f(x) = x

Politecnico DBG

• Generally the only situation where it makes sense to use it!

# Defining weights (parameters)

- So far, we assumed all weights and biases (let's call them  $\theta$ ) to be known
  - But, we still need to figure out how we find them!
- We pick a function (objective, or loss),  $\mathcal{L}(\theta)$ , that we want to minimize
  - e.g., in Linear Regression we minimize the Mean Squared Error

• 
$$\mathcal{L}(\theta) = MSE(\theta) = \frac{1}{n}\sum(y_i - \theta^T x_i)^2$$

• Then, we pick  $\theta$  that minimizes it

Note

 $\mathcal{L}$  also depends on the training points  $x_i$ ,  $y_i$ , so we should refer to it as  $\mathcal{L}(\theta, X, y)$ .

However, the training set X, y is generally fixed. Thus, we only have control over  $\theta$ , so we use the notation  $\mathcal{L}(\theta)$ .

# Linear regression

• For simple models, we can find the optimal weights in closed form

• 
$$\frac{\partial \mathcal{L}(\theta)}{\partial \theta} = \frac{\partial MSE(\theta)}{\partial \theta} = 0$$

- Quadratic in  $\theta$ , can be solved easily!
- Or, we can evaluate the loss function for a bunch of  $\theta$ 's, and find the "best" one

#### Note

For linear regression, we don't try a bunch of  $\theta$  since we can easily find the best value in closed form.

However, this provides the intuition for what we will do next with more complex loss functions/models.



## More complex losses/models

- For *more complex* loss functions/models, we may not be able to solve the problem in closed form
  - But we can evaluate  $\mathcal{L}(\theta)$  for various values of  $\theta$
- We can iteratively update  $\theta$  to reach a local minimum:
  - We start from a random value  $\theta$ , then
  - we "move around" according to "some policy"



# We "<mark>move around</mark>" according to "<mark>some policy</mark>"

- Move around = update  $\theta$  incrementally, based on its current value
  - The new value of  $\theta$  at any step depends on the previous step's value
  - $\theta_{t+1} := \theta_t + update$
- Some policy = we take a small step in the direction where the function decreases locally
  - i.e. in the *opposite* direction of the gradient
  - $\theta_{t+1} := \theta_t \alpha \nabla_{\theta} \mathcal{L}(\theta_t)$ 
    - for 1-dimensional  $\theta$ , we have  $\theta_{t+1} = \theta_t \alpha \frac{\partial \mathcal{L}(\theta)}{\partial \theta}$
    - $\alpha$ : learning rate, controls the "size" of the step
- Gradient Descent!



# Some limitations of GD

#### Note

Different initializations will lead to the global minimum for convex loss functions. However, that represents a trivial situation we typically do not encounter.



- Different initializations can lead to different solutions!
- GD can get stuck in local minima
- Various solutions to help prevent local minima:
  - Adding momentum
  - Adaptive learning rates
  - Learning rate schedules





## Backpropagation

Politecnico D<sup>B</sup>G

- So far, we assumed we were able to compute  $\nabla_{\theta} \mathcal{L}(\theta)$
- However, any loss/model combination would need a different gradient computation!
- We can use backpropagation to compute the gradient of the loss w.r.t. any weight!
  - Backpropagation is just a fancy word for "using the chain rule"

# Using the chain rule

- We use the chain rule from calculus,  $\frac{\partial f}{\partial x} = \frac{\partial f}{\partial g} \frac{\partial g}{\partial x}$ 
  - Sometimes known as  $f(g(x))' = f'(g(x)) \cdot g'(x)$
- And apply it from the end of the computational graph, backwards
  - (hence the name, backpropagation)

# Computational graph

- A computational graph is a directed graph
  - Each node corresponds to an operation
  - Each edge represents the flow of data between nodes
- For instance, we may want to compute y = wx + q
  - We start from three variables, w, x and y
  - The computational graph performs one operation at a time
    - First, compute the intermediate variable a = wx
    - Then, compute the output variable y = a + q = wx + q



# Backpropagation example

- Let's say:
  - Our dataset has one point, (x, y)
  - Our (weird) model has two parameters,  $\theta_1$  and  $\theta_2$ , and predicts  $\theta_1 \theta_2 x$
  - Our loss function will be  $\mathcal{L} = (\theta_1 \theta_2 x y)^2$
- We build a computational graph with all operations and intermediate variables
  - $a = \theta_1 \theta_2$
  - $b = ax = \theta_1 \theta_2 x$
  - $c = b y = ax y = \theta_1 \theta_2 x y$
  - $\mathcal{L} = c^2 = (b y)^2 = (ax y)^2 = (\theta_1 \theta_2 x y)^2$

\_\_\_\_ [ Data Mining Concepts & Algorithms ] \_\_\_\_\_ [ Introduction to deep learning ]

$$\frac{\partial \mathcal{L}}{\partial \theta_{1}} = \frac{\partial \mathcal{L}}{\partial a} \frac{\partial a}{\partial \theta_{1}} = 2cx \frac{\partial \theta_{1} \theta_{2}}{\partial \theta_{1}} = 2cx\theta_{2}$$

$$\theta_{1}$$

$$\frac{\partial \mathcal{L}}{\partial a} = \frac{\partial \mathcal{L}}{\partial b} \frac{\partial b}{\partial a} = 2c \frac{\partial ax}{\partial a} = 2cx$$

$$\theta_{1}$$

$$\frac{\partial \mathcal{L}}{\partial b} = \frac{\partial \mathcal{L}}{\partial c} \frac{\partial c}{\partial b} = 2c \frac{\partial (b - y)}{\partial b} = 2c$$

$$\theta_{2}$$

$$\frac{\partial \mathcal{L}}{\partial b} = \frac{\partial \mathcal{L}}{\partial c} \frac{\partial c}{\partial b} = 2c \frac{\partial \mathcal{L}}{\partial b} = 2c$$

$$\frac{\partial \mathcal{L}}{\partial c} = \frac{\partial \mathcal{L}}{\partial c} = 2c$$

$$\frac{\partial \mathcal{L}}{\partial c} = \frac{\partial \mathcal{L}}{\partial c} = 2c$$

$$\frac{\partial \mathcal{L}}{\partial \theta_{2}} = 2cx \frac{\partial \theta_{1} \theta_{2}}{\partial \theta_{2}} = 2cx\theta_{1}$$

$$x$$

$$y$$

$$y$$

#### Forward step

Politecnico DBG

• The loss  $\mathcal{L}$  is computed starting from the "inputs"  $\theta_1, \theta_2, x, y$ 

#### Backward step (backpropagation)

- The loss  $\mathcal{L}$  is used to compute the derivative w.r.t.  $c \rightarrow \frac{\partial \mathcal{L}}{\partial c}$
- The derivative  $\frac{\partial \mathcal{L}}{\partial c}$  is used to compute the derivative w.r.t.  $b \rightarrow \frac{\partial \mathcal{L}}{\partial b}$  The derivative  $\frac{\partial \mathcal{L}}{\partial b}$  is used to compute the derivative w.r.t.  $a \rightarrow \frac{\partial \mathcal{L}}{\partial a}$  The derivative  $\frac{\partial \mathcal{L}}{\partial a}$  is used to compute the derivative w.r.t.  $\theta_1, \theta_2 \rightarrow \frac{\partial \mathcal{L}}{\partial \theta_1}, \frac{\partial \mathcal{L}}{\partial \theta_2}$

$$\frac{\partial \mathcal{L}}{\partial \theta_1} = 2(\theta_1 \theta_2 x - y) x \theta_2$$
$$\frac{\partial \mathcal{L}}{\partial \theta_2} = 2(\theta_1 \theta_2 x - y) x \theta_1$$

24

#### 25

# Loss functions

- Regression
  - Mean Squared Error, Mean Absolute Error
- Binary

Politecnico D<sup>B</sup>G

- Binary Cross-Entropy (BCE)
- $y = \{0, 1\} \rightarrow$  ground truth
- $\hat{y} = model(x) \in [0,1] \Rightarrow$  predicted value

 $\mathcal{L} = -ylog(\hat{y}) - (1-y)log(1-\hat{y})$ 

 y (ground truth) acts as a "selector" of the loss term to be applied

• 
$$y = 1 \rightarrow \mathcal{L} = -log(\hat{y})$$

•  $y = 0 \Rightarrow \mathcal{L} = -log(1 - \hat{y})$ 



# Loss functions

- Multi-class classification
  - Cross-Entropy

- Generalization to multiple classes of BCE
- $y_i = 1$  when ground truth is the ith class, 0 otherwise
- $y_i$  plays the same "selector" mechanism as in BCE

$$\mathcal{L} = -\sum_{i} y_i \log(\widehat{y_i})$$