### Machine learning lecture slides

COMS 4771 Fall 2020

### **Optimization II: Neural networks**

# Outline

- Architecture of (layered) feedforward neural networks
- Universal approximation
- Backpropagation
- Practical issues

### Parametric featurizations

- So far: data features (x or  $\varphi(x)$ ) are fixed during training
  - Consider a (small) collection of feature transformations  $\varphi$
  - ▶ Select  $\varphi$  via cross-validation outside of normal training
- "Deep learning" approach:
  - Use  $\varphi$  with many tunable parameters
  - Optimize parameters of  $\varphi$  during normal training process

• <u>Neural network</u>: parameterization for function  $f : \mathbb{R}^d \to \mathbb{R}$ 

 $\blacktriangleright \ f(x) = \varphi(x)^{\mathsf{T}} w$ 

 $\blacktriangleright$  Parameters include both w and parameters of  $\varphi$ 

- Varying parameters of  $\varphi$  allows f to be essentially any function!
- Major challenge: optimization



Figure 1: Neural networks as feature maps

#### Feedforward neural network

Architecture of a <u>feedforward neural network</u>

- Directed acyclic graph G = (V, E)
- One <u>source</u> node (vertex) per input to the function  $(x_1, \ldots, x_d)$
- One <u>sink</u> node per output of the function
- Internal nodes are called <u>hidden units</u>
- ▶ Each edge  $(u, v) \in E$  has a weight parameter  $w_{u,v} \in \mathbb{R}$
- ► <u>Value</u>  $h_v$  of node v given values of parents  $\pi_G(v) = \{u \in V : (u, v) \in E\}$  is

$$h_v := \sigma_v(z_v), \quad z_v := \sum_{u \in \pi_G(v)} w_{u,v} \cdot h_u.$$

•  $\sigma_v \colon \mathbb{R} \to \mathbb{R}$  is the <u>activation function</u> (a.k.a. <u>link function</u>) • E.g., sigmoid function  $\sigma_v(z) = 1/(1 + e^{-z})$ 

Inspired by neurons in the brain



Figure 2: Computation DAG of a feedforward neural network

### Standard layered architectures

- Standard feedforward architecture arranges nodes into *layers* 
  - Initial layer (layer zero): source nodes (input)
  - ► Final layer (layer *L*): sink nodes (output)
  - (Layer counting is confusing; usually don't count input)
- Edges only go from one layer to the next
  - (Non-standard feedforward architectures (e.g., ResNets) break this rule)
- Can write function using matrices of weight parameters

$$f(x) = \sigma_L(W_L \sigma_{L-1}(\cdots \sigma_1(W_1 x) \cdots))$$

- Layer  $\ell$  has  $d_{\ell}$  nodes
- $W_{\ell} \in \mathbb{R}^{d_{\ell} \times d_{\ell-1}}$  are the weight parameters for layer  $\ell$
- Scalar-valued activation function σ<sub>ℓ</sub>: ℝ → ℝ (e.g., sigmoid) is applied coordinate-wise to input
- Often also include "bias" parameters  $b_\ell \in \mathbb{R}^{d_\ell}$

$$f(x) = \sigma_L(b_L + W_L \sigma_{L-1}(\cdots \sigma_1(b_1 + W_1 x) \cdots))$$

• The tunable parameters:  $\theta = (W_1, b_1, \dots, W_L, b_L)$ 



input hidden units output

Figure 3: Layered feedforward neural network

### Well-known activation functions

• <u>Heaviside</u>:  $\sigma(z) = \mathbf{1}_{\{z > 0\}}$ Popular in the 1940s; also called step function • Sigmoid (from logistic regression):  $\sigma(z) = 1/(1 + e^{-z})$ Popular since 1970s • Hyperbolic tangent:  $\sigma(z) = \tanh(z)$ Similar to sigmoid, but range is (-1,1) rather than (0,1)• <u>Rectified Linear Unit</u> (<u>ReLU</u>):  $\sigma(z) = \max\{0, z\}$ Popular since 2012 • Identity:  $\sigma(z) = z$ Popular with luddites

### Power of non-linear activations

- What happens if every activation function is linear/affine?
  - Overall function is affine
  - An unusual way to parameterize an affine function
- ► Therefore, use non-linear/non-affine activation functions

# Necessity of multiple layers (1)

• Suppose only have input and output layers, so function f is

$$f(x) = \sigma(b + w^{\mathsf{T}}x)$$

where  $b \in \mathbb{R}$  and  $w \in \mathbb{R}^d$  (so  $w^{\mathrm{\scriptscriptstyle T}} \in \mathbb{R}^{1 \times d})$ 

X

 If σ is monotone (e.g., Heaviside, sigmoid, hyperbolic tangent, ReLU, identity), then f has same limitations as a linear/affine classifier

0



Figure 4: XOR data set

# Necessity of multiple layers (2)

# XOR problem • Let $x^{(1)} = (+1, +1), x^{(2)} = (+1, -1), x^{(3)} = (-1, +1),$ $x^{(4)} = (-1, -1).$ • $y^{(i)} = +1$ iff coordinates of $x^{(i)}$ are the same. (XNOR) Suppose $(w, b) \in \mathbb{R}^2 \times \mathbb{R}$ satisfies $-b - w^{\mathsf{T}} r^{(1)} < 0$ $b + w^{\mathsf{T}} x^{(2)} < 0.$ $b + w^{\mathsf{T}} r^{(3)} < 0$ Add up the equations: $b + w^{\mathsf{T}}(x^{(2)} + x^{(3)} - x^{(1)}) < 0.$ • But $x^{(2)} + x^{(3)} - x^{(1)} = x^{(4)}$ , so

 $b + w^{\mathsf{T}} x^{(4)} < 0.$ 

In other words, cannot correctly label  $x^{(4)}$ .

### Neural network approximation theorems

Theorem (Cybenko, 1989; Hornik, Stinchcombe, & White, 1989): Let σ<sub>1</sub> be any continuous non-linear activation function from above. For any continuous function f: ℝ<sup>d</sup> → ℝ and any ε > 0, there is a two-layer neural network (with parameters θ = (W<sub>1</sub>, b<sub>1</sub>, w<sub>2</sub>)) s.t.

$$\max_{x \in [0,1]^d} |f(x) - w_2^{\mathsf{T}} \sigma_1 (b_1 + W_1 x)| < \varepsilon.$$

- This property of such families of neural networks is called universal approximation.
- Many caveats
  - "Width" (number of hidden units) may need to be very large
  - Does not tell us how to find the network
  - Does not justify deeper networks

## Stone-Weierstrass theorem (polynomial version)

**Theorem** (Weierstrass, 1885): For any continuous function  $f: [a, b] \to \mathbb{R}$ , and any  $\epsilon > 0$ , there exists a polynomial p such that

$$\sup_{x \in [a,b]} |f(x) - p(x)| < \epsilon.$$

# Stone-Weierstrass theorem (general version)

**Theorem** (Stone, 1937): Let  $K \subset \mathbb{R}^d$  be any bounded set. Let A be a set of continuous functions on K such that the following hold.

- (1) A is an <u>algebra</u> (i.e., A is closed under addition, multiplication, and scalar multiplication).
- (2) A does not vanish on K (i.e., for all  $x \in K$ , there exists  $h \in A$  such that  $h(x) \neq 0$ ).
- (3) A separates points in K (i.e., for all distinct  $x, y \in K$ , there exists  $h \in A$  such that  $h(x) \neq h(y)$ ).

For any continuous function  $f\colon K\to\mathbb{R},$  and any  $\epsilon>0,$  there exists  $h\in A$  such that

$$\sup_{x \in K} |f(x) - h(x)| < \epsilon.$$

#### Two-layer neural networks with cosine activation functions

Let 
$$K = [0, 1]^d$$
, and let  

$$A = \left\{ x \mapsto \sum_{k=1}^m a_k \cos(x^{\mathsf{T}} w_k + b_k) : m \in \mathbb{N}, a_k, b_k \in \mathbb{R}, w_k \in \mathbb{R}^d \text{ for } k = 1, \dots, m \right\}.$$

(Check that A satisfies properties of Stone-Weierstrass theorem.)

#### Two-layer neural networks with exp activation functions

Let 
$$K = [0, 1]^d$$
, and let  

$$A = \left\{ x \mapsto \sum_{k=1}^m a_k \exp(x^\mathsf{T} w_k + b_k) : m \in \mathbb{N}, a_k, b_k \in \mathbb{R}, w_k \in \mathbb{R}^d \text{ for } k = 1, \dots, m \right\}.$$

(Check that A satisfies properties of Stone-Weierstrass theorem.)

#### Fitting neural networks to data

- ▶ Training data  $(x_1, y_1), \dots, (x_n, y_n) \in \mathbb{R}^d \times \mathbb{R}$
- Fix architecture: G = (V, E) and activation functions
- ERM: find parameters  $\theta$  of neural network  $f_{\theta}$  to minimize empirical risk (possibly with a surrogate loss)

• Regression  $y_i \in \mathbb{R}$ 

$$\widehat{\mathcal{R}}(\theta) = \frac{1}{n} \sum_{i=1}^{n} (f_{\theta}(x_i) - y_i)^2$$

• Binary classification  $y_i \in \{-1, +1\}$ 

$$\widehat{\mathcal{R}}(\theta) = \frac{1}{n} \sum_{i=1}^{n} \ln(1 + \exp(-y_i f_{\theta}(x_i)))$$

(Could use other surrogate loss functions ...)

Can also add regularization terms, but also common to use algorithmic regularization

• Typically objective is not convex in parameters  $\theta$ 

Nevertheless, local search (e.g., gradient descent, SGD) often works well!

#### Backpropagation

Backpropagation (backprop): Algorithm for computing partial derivatives wrt weights in a feedforward neural network

- Clever organization of partial derivative computations with <u>chain rule</u>
- Use in combination with gradient descent, SGD, etc.
- Consider loss on a single example (x, y), written as

$$J := \ell(y, f_{\theta}(x))$$

- ▶ Goal: compute  $\frac{\partial J}{\partial w_{u,v}}$  for every edge  $(u,v) \in E$
- Initial step of backprop: forward propagation
  - Compute  $z_v$ 's and  $h_v$ 's for every node  $v \in V$
  - Running time: linear in size of network
- We'll see that rest of backprop also just requires time linear in size of network

#### Derivative of loss with respect to weights

- Let  $\hat{y}_1, \hat{y}_2, \ldots$  denote the values at the output nodes.
- Then by chain rule,

$$\frac{\partial J}{w_{u,v}} = \sum_{i} \frac{\partial J}{\partial \hat{y}_i} \cdot \frac{\partial \hat{y}_i}{w_{u,v}}.$$

∂J/∂ŷ<sub>i</sub> is just determined by the loss function (e.g., squared loss)

 So just have to focus on ∂ŷ<sub>i</sub>/w<sub>u,v</sub>
 Assume for simplicity there is just a single output, ŷ

#### Derivative of output with respect to weights

Chain rule, again:

$$\frac{\partial \hat{y}}{\partial w_{u,v}} = \frac{\partial \hat{y}}{\partial h_v} \cdot \frac{\partial h_v}{\partial w_{u,v}}$$

- First term: trickier; we'll handle later
- Second term:

▶ 
$$h_v = \sigma_v(z_v)$$
  
▶  $z_v = w_{u,v} \cdot h_u + (\text{terms not involving } w_{u,v})$   
▶ Therefore

$$\frac{\partial h_v}{\partial w_{u,v}} = \frac{\partial h_v}{\partial z_v} \cdot \frac{\partial z_v}{\partial w_{u,v}} = \sigma'(z_v) \cdot h_u.$$

 $\blacktriangleright \ z_v$  and  $h_u$  were computed during forward propagation



Figure 5: Derivative of a node's output with respect to an incoming weight

#### Derivative of output with respect to hidden units

- If v is not the output node, then by chain rule (yet again),

$$\frac{\partial \hat{y}}{\partial h_v} = \sum_{v': (v,v') \in E} \frac{\partial \hat{y}}{\partial h_{v'}} \cdot \frac{\partial h_{v'}}{\partial h_v}$$

- $\frac{\partial \hat{y}}{\partial h_{v'}}
   was already computed since v' is in higher layer than v
   <math>h_{v'} = \sigma_{v'}(z_{v'})$   $2_{v'} = w_{v,v'} \cdot h_v + (\text{terms not involving } h_v)$  Therefore
    $\frac{\partial h_{v'}}{\partial h_v} = \frac{\partial h_{v'}}{\partial z_{v'}} \cdot \frac{\partial z_{v'}}{\partial h_v}
   = \sigma'(z_{v'}) \cdot w_{v,v'}.$
- z<sub>v</sub>'s were computed during forward propagation
   w<sub>v,v'</sub>'s are the values of the weight parameters at which we want to compute the gradient



Figure 6: Derivative of the network output with respect to hidden unit values

# Example: chain graph (1)

- Function  $f_{\theta} \colon \mathbb{R} \to \mathbb{R}$
- Architecture
  - $\blacktriangleright \mathsf{DAG:} \ 0 \longrightarrow 1 \longrightarrow 2 \longrightarrow \cdots \longrightarrow L$
  - Same activation  $\sigma$  in every layer
- Parameters  $\theta = (w_{0,1}, w_{1,2}, \dots, w_{L-1,L})$ 
  - ▶ Input is at vertex 0, and output is at vertex L
- ▶ Fix input value  $x \in \mathbb{R}$ ; what is  $\frac{\partial h_L}{\partial w_{i-1,i}}$  for i = 1, ..., L? ▶ Forward propagation:

• 
$$h_0 := x$$
  
• For  $i = 1, 2, ..., L$ :

$$z_i := w_{i-1,i}h_{i-1}$$
$$h_i := \sigma(z_i)$$

Example: chain graph (2)

#### Backprop:

• For i = L, L - 1, ..., 1:

$$\begin{split} \frac{\partial h_L}{\partial h_i} &:= \begin{cases} 1 & \text{if } i = L\\ \frac{\partial h_L}{\partial h_{i+1}} \cdot \sigma'(z_{i+1}) w_{i,i+1} & \text{if } i < L \end{cases}\\ \frac{\partial h_L}{\partial w_{i-1,i}} &:= \frac{\partial h_L}{\partial h_i} \cdot \sigma'(z_i) h_{i-1} \end{split}$$



Figure 7: Neural network with a chain computation graph

### Practical issues I: Initialization

- Ensure inputs are <u>standardized</u>: every feature has zero mean and unit variance (wrt training data)
  - Even better: different features have zero covariance (again, on training data)
  - But this can be expensive
- Initialize weights randomly for gradient descent / SGD
  - Standard normal random variables (or similar)
  - What should variance be?
  - Heuristic: ensure  $h_v$  have similar statistics as inputs
  - E.g., using tanh-activation, if v has in-degree k, use variance 1/k for all weights  $w_{u,v}$
  - Many initialization schemes for other activations (e.g., ReLU), dealing with bias parameters, ...

### Practical issues II: Architecture choice

- Architecture can be regarded as a "hyperparameter"
  - Could use cross-validation to select, but ...
  - Many "good" architectures are known for popular problems (e.g., image classification)
  - Unclear what to do for completely new problems
- Optimization-inspired architecture choice
  - With wide enough network, can get zero training error
  - Use the smallest network that lets you get zero training error
  - Then add regularization term to objective (e.g., sum of squares of weights), and optimize the regularized ERM objective
- Entire research communities are trying to figure out good architectures for their problems

### Multi-class

Vector-valued activation:  $\sigma \colon \mathbb{R}^{d_{\ell}} \to \mathbb{R}^{d_{\ell}}$ 

- <u>Softmax</u> activation:  $\sigma(v)_i = \exp(v_i) / \sum_j \exp(v_j)$
- Common to use this in final layer

# Convolutional nets

- Neural networks with convolutional layers
  - Useful when inputs have locality structure
  - Sequential structure (e.g., speech waveform)
  - 2D grid structure (e.g., image)
  - ▶ ...
- Weight matrix  $W_{\ell}$  is highly-structured
  - Determined by a small <u>filter</u>
  - Time to compute  $W_{\ell}h_{\ell-1}$  is typically  $\ll d_{\ell} \times d_{\ell-1}$  (e.g., closer to max $\{d_{\ell}, d_{\ell-1}\}$ )

### Convolutions I

 $\blacktriangleright$  Convolution of two continuous functions:  $h:=f\ast g$ 

$$h(x) = \int_{-\infty}^{+\infty} f(y)g(x-y) \,\mathrm{d}y$$

 $\blacktriangleright \ \, {\rm If} \ \, f(x)=0 \ {\rm for} \ \, x\notin [-w,+w] {\rm , \ then}$ 

$$h(x) = \int_{-w}^{+w} f(y)g(x-y) \,\mathrm{d}y$$

• Replaces g(x) with weighted combination of g at nearby points

### Convolutions II

▶ For functions on discrete domain, replace integral with sum

$$h(i) = \sum_{j=-\infty}^{\infty} f(j)g(i-j)$$

E.g., suppose only f(0),f(1),f(2) are non-zero. Then:

$$\begin{bmatrix} h(1) \\ h(2) \\ h(3) \\ h(4) \\ h(5) \\ h(6) \\ h(7) \end{bmatrix} = \begin{bmatrix} f(0) & 0 & 0 & 0 & 0 \\ f(1) & f(0) & 0 & 0 & 0 \\ f(2) & f(1) & f(0) & 0 & 0 \\ 0 & f(2) & f(1) & f(0) & 0 \\ 0 & 0 & f(2) & f(1) & f(0) \\ 0 & 0 & 0 & f(2) & f(1) \\ 0 & 0 & 0 & 0 & f(2) \end{bmatrix} \begin{bmatrix} g(1) \\ g(2) \\ g(3) \\ g(3) \\ g(4) \\ g(5) \end{bmatrix}$$

(Here, we pretend g(i) = 0 for i < 1 and i > 5.)



Figure 8: Convolutional layer

# Convolutions III

- Similar for 2D inputs (e.g., images), except now sum over two indices
  - ▶ g is the input image
  - ► *f* is the filter
  - Lots of variations (e.g., padding, strides, multiple "channels")
- Use additional layers/activations to down-sample after convolution
  - E.g., max-pooling



Figure 9: 2D convolution



Figure 10: 2D convolution



Figure 11: 2D convolution



Figure 12: 2D convolution