

#### 10-301/601 Introduction to Machine Learning

Machine Learning Department School of Computer Science Carnegie Mellon University

# Regularization + Neural Networks

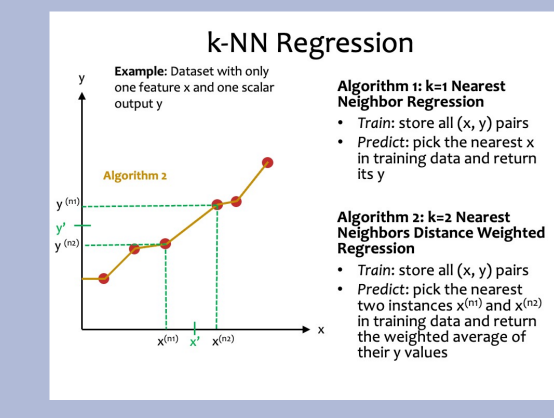
Matt Gormley Lecture 11 Oct. 3, 2022

#### Reminders

- Practice Problems: Exam 1
- Exam 1
  - Tue, Oct 4, 6:30pm 8:30pm
  - see Piazza for exam location
- Homework 4: Logistic Regression
  - Out: Tue, Oct 4
  - Due: Thu, Oct 13 at 11:59pm
- Be careful to submit to the correct submission slot on Gradescope!

#### Q&A

#### **Q:** Is this correct?

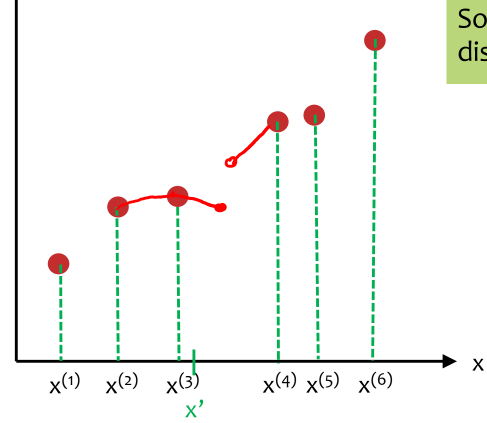


#### A: Oops! No.

3

### k-NN Regression

**Example:** Dataset with only one feature x and one scalar output y



Q: What are the k=2 nearest neighbors of the test point x'?

A: x<sup>(2)</sup> and x<sup>(3)</sup>. So the output curve is actually discontinuous!

#### Algorithm 2: k=2 Nearest Neighbors Distance Weighted Regression

- Train: store all (x, y) pairs
- Predict: pick the nearest two instances x<sup>(n1)</sup> and x<sup>(n2)</sup> in training data and return the weighted average of their y values

#### REGULARIZATION

# Overfitting

**Definition:** The problem of **overfitting** is when the model captures the noise in the training data instead of the underlying structure

Overfitting can occur in all the models we've seen so far:

- Decision Trees (e.g. when tree is too deep)
- KNN (e.g. when k is small)
- Perceptron (e.g. when sample isn't representative)
- Linear Regression (e.g. with nonlinear features)
- Logistic Regression (e.g. with many rare features)

### Motivation: Regularization

- Occam's Razor: prefer the simplest hypothesis
- What does it mean for a hypothesis (or model) to be simple?
  - 1. small number of features (model selection)
  - small number of "important" features (shrinkage)

- **Given** objective function:  $J(\theta)$
- **Goal** is to find:  $\hat{\theta} = \operatorname{argmin} J(\theta) + \lambda r(\theta)$ θ
- Key idea: Define regularizer r( $\theta$ ) s.t. we tradeoff between fitting the data and keeping the model simple
- Choose form of r(θ):

- Example: q-norm (usually p-norm):  $\|\boldsymbol{\theta}\|_q = \left(\sum_{m=1}^{M} |\theta_m|^q\right)^{\overline{q}}$ 

$\overline{q}$	$r(oldsymbol{ heta})$	yields parame- ters that are	name	optimization notes
0	$  \boldsymbol{\theta}  _0 = \sum \mathbb{1}(\theta_m \neq 0)$	zero values	Lo reg.	no good computa- tional solutions
$rac{1}{2}$	$egin{aligned}   oldsymbol{ heta}  _1 &= \sum   heta_m  \ (  oldsymbol{ heta}  _2)^2 &= \sum  heta_m^2 \end{aligned}$	zero values small values	L1 reg. L2 reg.	subdifferentiable differentiable

#### **Regularization Examples**

Add an L2 regularizer to Linear Regression (aka. Ridge Regression)

$$J_{\mathsf{RR}}(\boldsymbol{\theta}) = J(\boldsymbol{\theta}) + \lambda ||\boldsymbol{\theta}||_2^2$$
$$= \frac{1}{N} \sum_{i=1}^N \frac{1}{2} (\boldsymbol{\theta}^T \mathbf{x}^{(i)} - y^{(i)})^2 + \lambda \sum_{m=1}^M \theta_m^2$$

Add an L1 regularizer to Linear Regression (aka. LASSO)

$$J_{\text{LASSO}}(\boldsymbol{\theta}) = J(\boldsymbol{\theta}) + \lambda ||\boldsymbol{\theta}||_{1}$$
$$= \frac{1}{N} \sum_{i=1}^{N} \frac{1}{2} (\boldsymbol{\theta}^{T} \mathbf{x}^{(i)} - y^{(i)})^{2} + \lambda \sum_{m=1}^{M} |\boldsymbol{\theta}_{m}|$$

#### **Regularization Examples**

Add an L2 regularizer to Logistic Regression

$$J'(\boldsymbol{\theta}) = J(\boldsymbol{\theta}) + \lambda ||\boldsymbol{\theta}||_2^2$$
$$= \frac{1}{N} \sum_{i=1}^N -\log p(y^{(i)} \mid \mathbf{x}^{(i)}, \boldsymbol{\theta}) + \lambda \sum_{m=1}^M \theta_m^2$$

Add an L1 regularizer to Logistic Regression

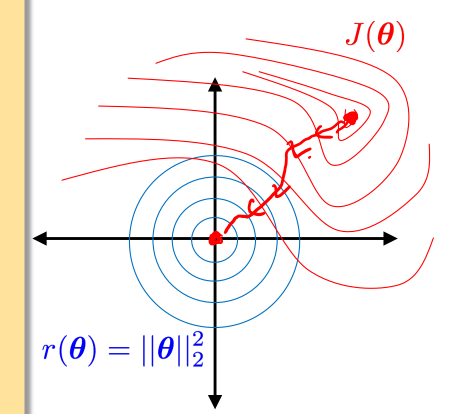
$$J'(\boldsymbol{\theta}) = J(\boldsymbol{\theta}) + \lambda ||\boldsymbol{\theta}||_{1}$$
$$= \frac{1}{N} \sum_{i=1}^{N} -\log p(y^{(i)} \mid \mathbf{x}^{(i)}, \boldsymbol{\theta}) + \lambda \sum_{m=1}^{M} |\boldsymbol{\theta}_{m}|$$

#### Question: (

Suppose we are minimizing J'( $\theta$ ) where  $J'(\theta) = J(\theta) + \lambda r(\theta) - 22$ As  $\lambda$  increases, the minimum of J'( $\theta$ ) will...

- A. ... move towards the midpoint between  $J(\theta)$  and  $r(\theta)$
- B. ... move towards the minimum of  $J(\theta)$
- C. ... move towards the minimum of  $r(\theta)$
- D. ... move towards a theta vector of positive infinities
- E. ... move towards a theta vector of negative infinities

F. ... stay the same



#### Whiteboard

– Why does L2 regularization lead to small numbers and L1 regularization lead to zeros?

#### Don't Regularize the Bias (Intercept) Parameter!

- In our models so far, the bias / intercept parameter is usually denoted by  $\theta_0$  -- that is, the parameter for which we fixed  $x_0 = 1$
- Regularizers always avoid penalizing this bias / intercept parameter
- Why? Because otherwise the learning algorithms wouldn't be invariant to a shift in the y-values

#### **Whitening Data**

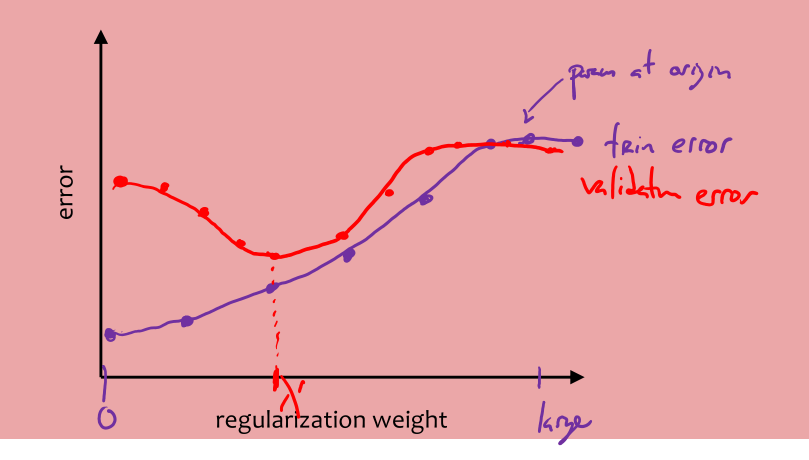
- It's common to whiten each feature by subtracting its mean and dividing by its variance
- For regularization, this helps all the features be penalized in the same units (e.g. convert both centimeters and kilometers to z-scores)

#### **Regularization Exercise**

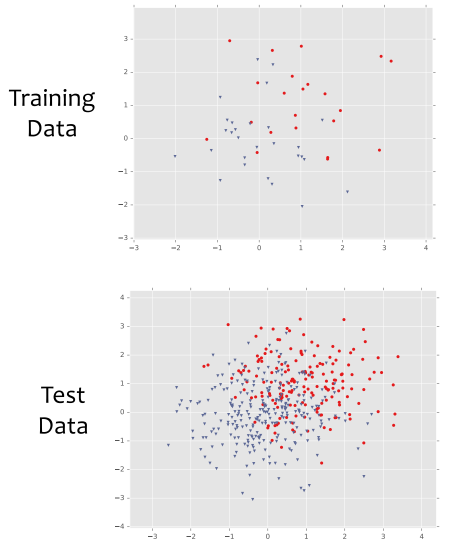
J'(0) = (1 - 3)J(0) + 3r(0)

#### In-class Exercise

- 1. Plot train error vs. regularization weight (cartoon)
- 2. Plot validation error vs. regularization weight (cartoon)

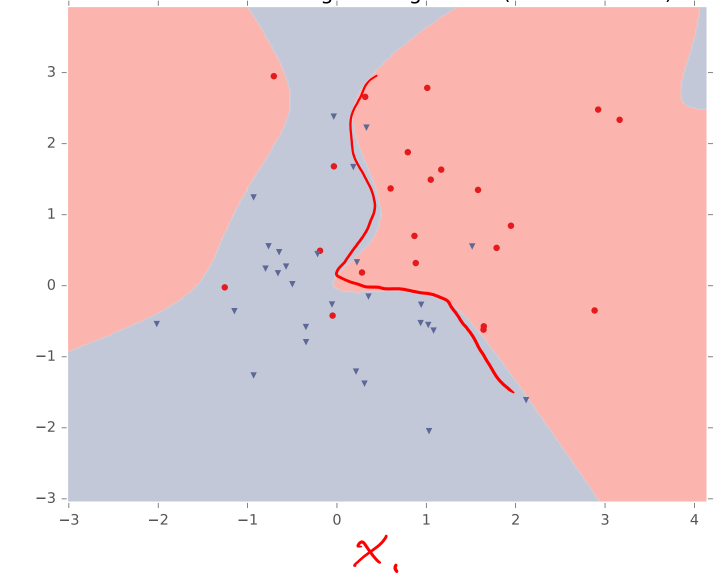


### **REGULARIZATION EXAMPLE:** LOGISTIC REGRESSION

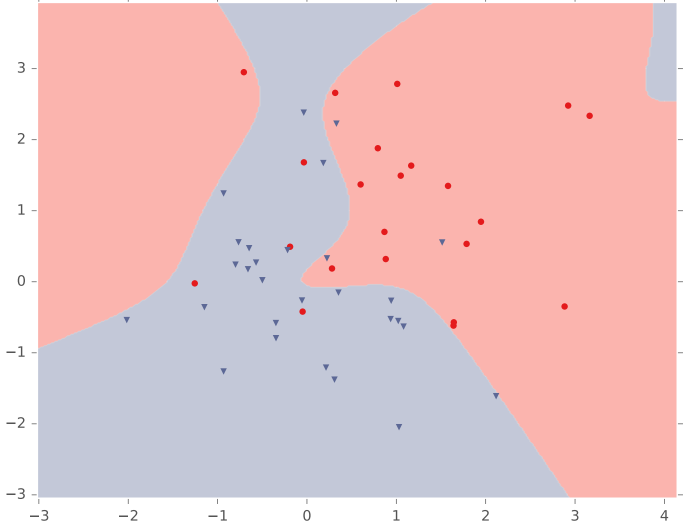


- For this example, we construct **nonlinear features** (i.e. feature engineering)
- Specifically, we add polynomials up to order 9 of the two original features x<sub>1</sub> and x<sub>2</sub>
- Thus our classifier is linear in the high-dimensional feature space, but the decision boundary is nonlinear when visualized in low-dimensions (i.e. the original two dimensions)

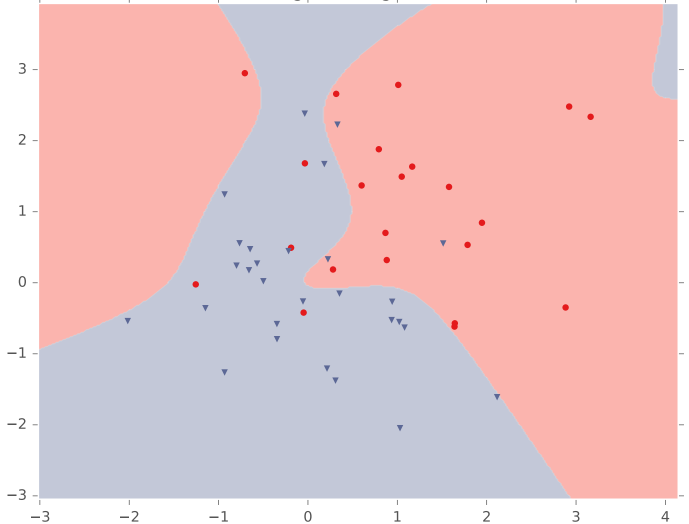
Classification with Logistic Regression (lambda=1e-05)



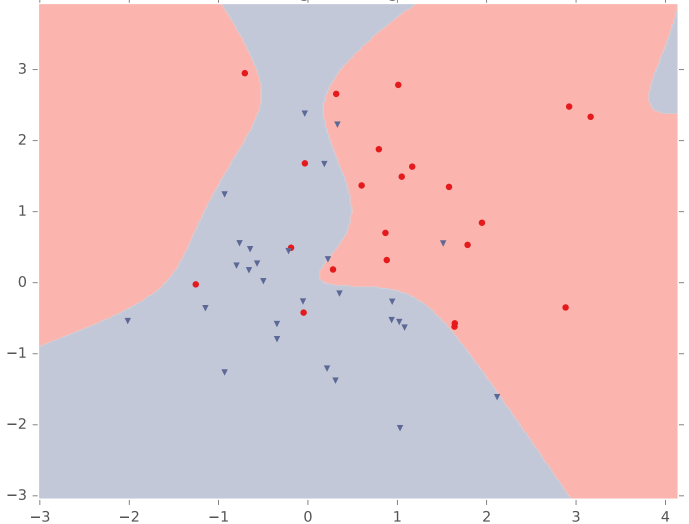
Classification with Logistic Regression (lambda=0.0001)



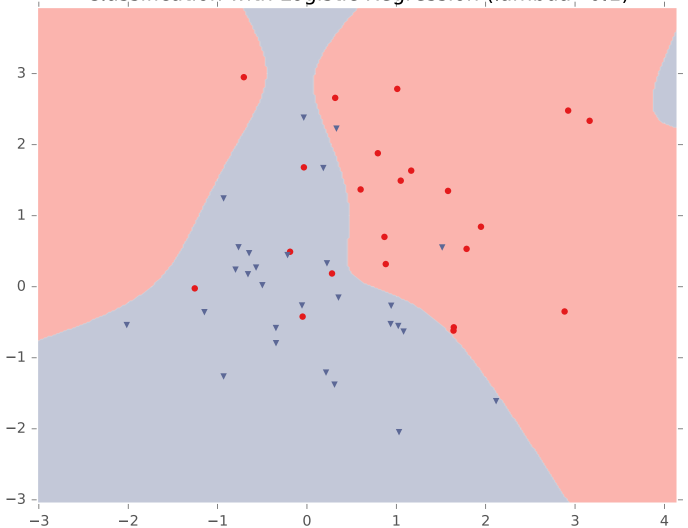
Classification with Logistic Regression (lambda=0.001)



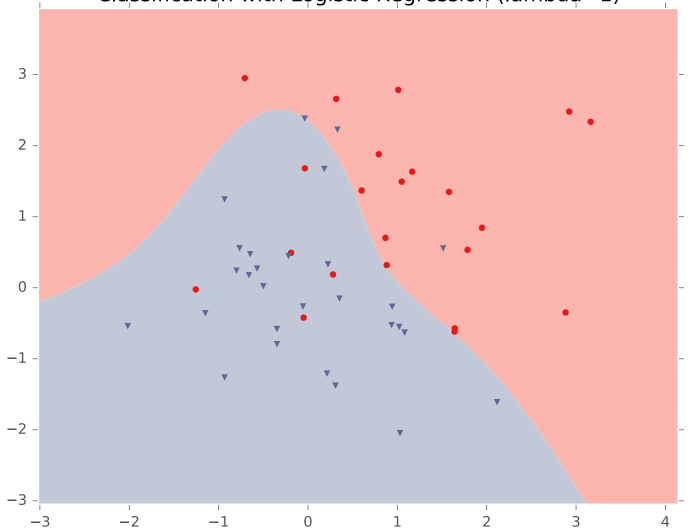
Classification with Logistic Regression (lambda=0.01)



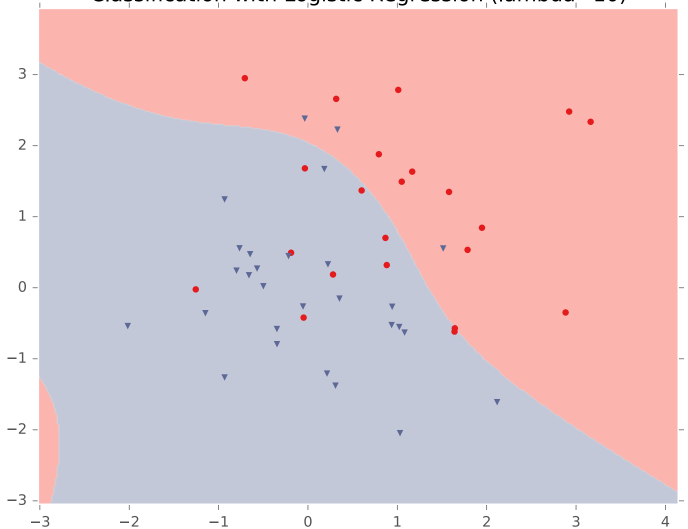
Classification with Logistic Regression (lambda=0.1)



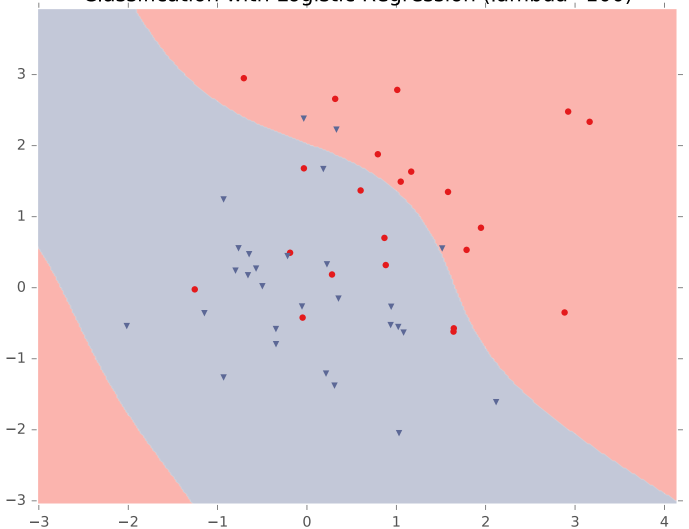
Classification with Logistic Regression (lambda=1)



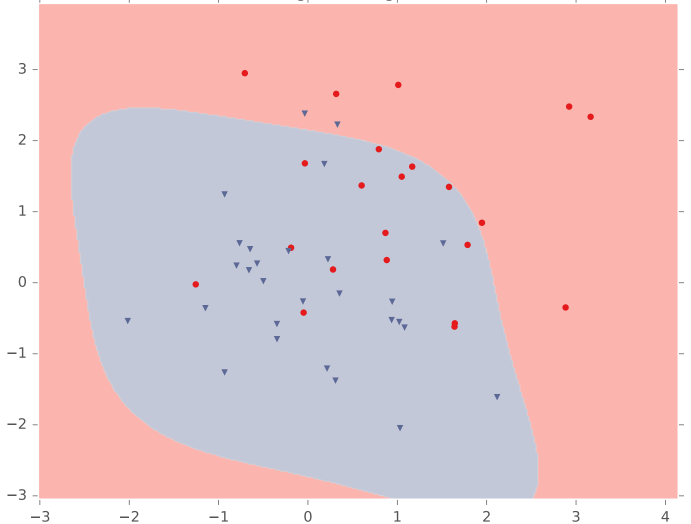
Classification with Logistic Regression (lambda=10)



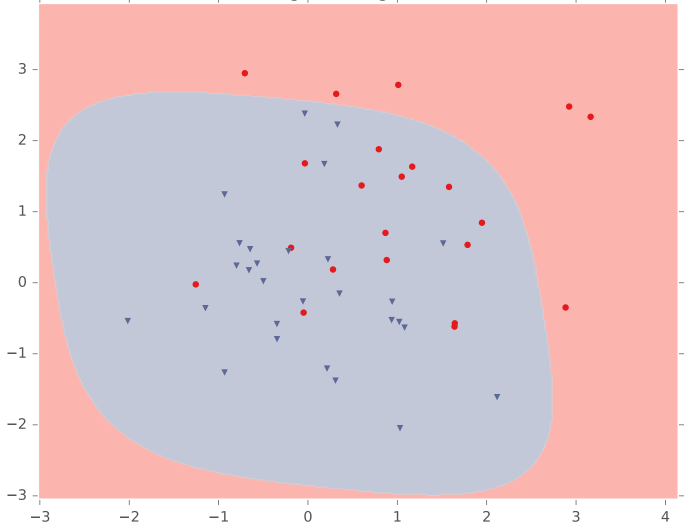
Classification with Logistic Regression (lambda=100)



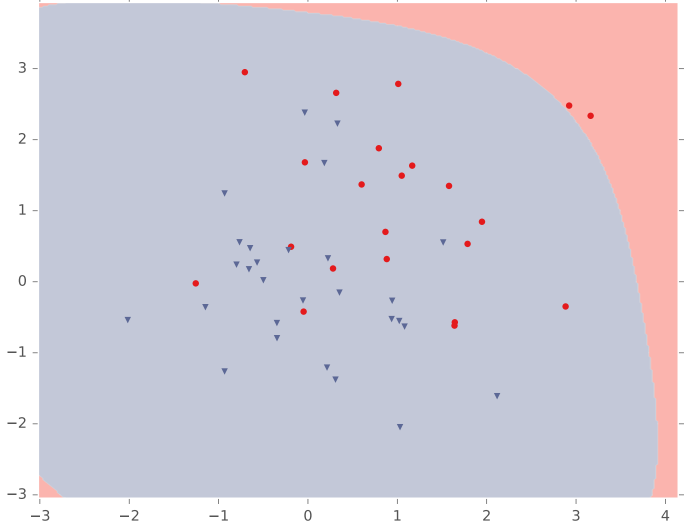
Classification with Logistic Regression (lambda=1000)



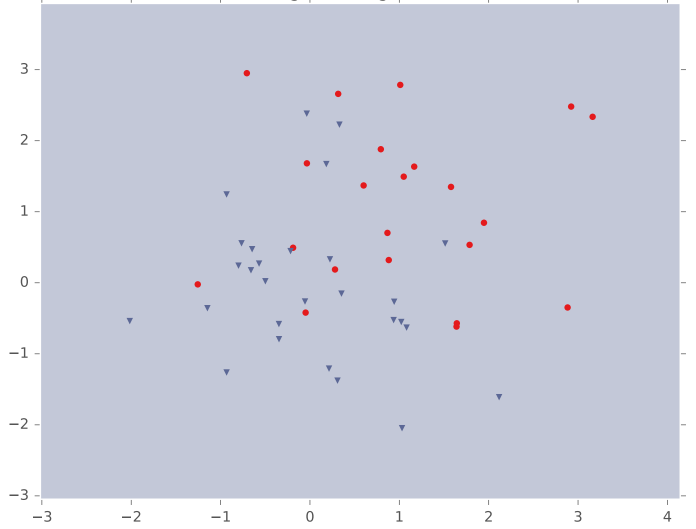
Classification with Logistic Regression (lambda=10000)



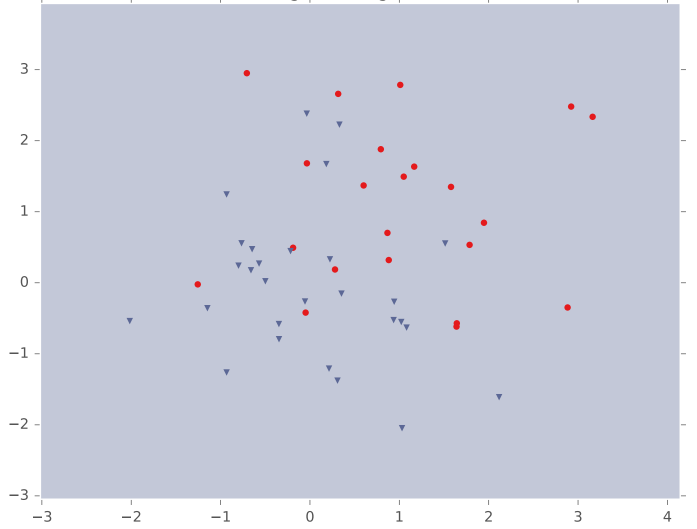
Classification with Logistic Regression (lambda=100000)

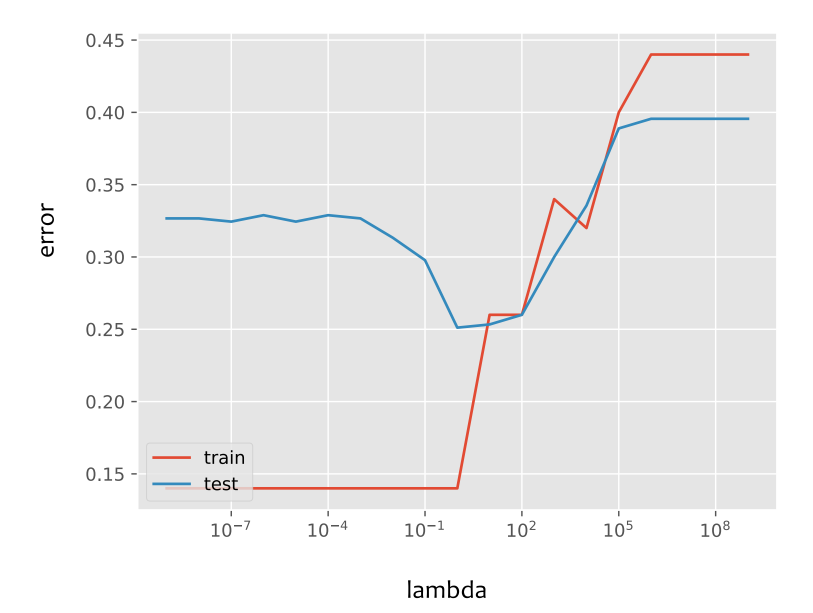


Classification with Logistic Regression (lambda=1e+06)



Classification with Logistic Regression (lambda=1e+07)





### OPTIMIZATION FOR L1 REGULARIZATION

#### **Optimization for L1 Regularization**

Can we apply SGD to the LASSO learning problem? argmin  $J_{\rm LASSO}(\theta)$ 

$$J_{\text{LASSO}}(\boldsymbol{\theta}) = J(\boldsymbol{\theta}) + \lambda ||\boldsymbol{\theta}||_{1}$$
$$= \frac{1}{2} \sum_{i=1}^{N} (\boldsymbol{\theta}^{T} \mathbf{x}^{(i)} - y^{(i)})^{2} + \lambda \sum_{k=1}^{K} |\boldsymbol{\theta}_{k}|$$

### Optimization for L1 Regularization

• Consider the absolute value function:

$$r(\boldsymbol{\theta}) = \lambda \sum_{k=1}^{K} |\theta_k|$$

• The L1 penalty is subdifferentiable (i.e. not differentiable at 0)

Def: A vector  $g \in \mathbb{R}^M$  is called a **subgradient** of a function  $f(\mathbf{x})$ :  $\mathbb{R}^M \to \mathbb{R}$  at the point  $\mathbf{x}$  if, for all  $\mathbf{x}' \in \mathbb{R}^M$ , we have:

$$f(\mathbf{x}') \ge f(\mathbf{x}) + \mathbf{g}^T(\mathbf{x}' - \mathbf{x})$$

# Optimization for L1 Regularization

- The L1 penalty is subdifferentiable (i.e. not differentiable at 0)
- An array of optimization algorithms exist to handle this issue:
   Basically the same as GD
  - Subgradient descent
  - Stochastic subgradient descent
  - Coordinate Descent
  - Othant-Wise Limited memory Quasi-Newton (OWL-QN) (Andrew & Gao, 2007) and provably convergent variants
  - Block coordinate Descent (Tseng & Yun, 2009)
  - Sparse Reconstruction by Separable Approximation (SpaRSA) (Wright et al., 2009)
  - Fast Iterative Shrinkage Thresholding Algorithm (FISTA) (Beck & Teboulle, 2009)

and SGD, but you use

one of the subgradients

when necessary

### **Regularization as MAP**

- L1 and L2 regularization can be interpreted as maximum a-posteriori (MAP) estimation of the parameters
- To be discussed later in the course...

#### Takeaways

- Nonlinear basis functions allow linear models (e.g. Linear Regression, Logistic Regression) to capture nonlinear aspects of the original input
- Nonlinear features are require no changes to the model (i.e. just preprocessing)
- 3. Regularization helps to avoid overfitting
- **4. Regularization** and **MAP estimation** are equivalent for appropriately chosen priors

# Feature Engineering / Regularization Objectives

You should be able to...

- Engineer appropriate features for a new task
- Use feature selection techniques to identify and remove irrelevant features
- Identify when a model is overfitting
- Add a regularizer to an existing objective in order to combat overfitting
- Explain why we should **not** regularize the bias term
- Convert linearly inseparable dataset to a linearly separable dataset in higher dimensions
- Describe feature engineering in common application areas

## **NEURAL NETWORKS**

#### Background

# A Recipe for Machine Learning

- 1. Given training data: $\{m{x}_i,m{y}_i\}_{i=1}^N$
- 2. Choose each of these:
  - Decision function
    - $\hat{\boldsymbol{y}} = f_{\boldsymbol{\theta}}(\boldsymbol{x}_i)$
  - Loss function
    - $\ell(\hat{oldsymbol{y}},oldsymbol{y}_i)\in\mathbb{R}$



**Examples:** Linear regression, Logistic regression, Neural Network

**Examples:** Mean-squared error, Cross Entropy

#### Background

# A Recipe for Machine Learning

- 1. Given training data: $\{m{x}_i,m{y}_i\}_{i=1}^N$
- 2. Choose each of these:
  - Decision function
    - $\hat{\boldsymbol{y}} = f_{\boldsymbol{\theta}}(\boldsymbol{x}_i)$
  - Loss function

 $\ell(\hat{oldsymbol{y}},oldsymbol{y}_i)\in\mathbb{R}$ 

3. Define goal:  $\underset{\theta}{\mathsf{risk}}^{N}$ 

4. Train with SGD:(take small steps opposite the gradient)

 $\boldsymbol{\theta}^{(t+1)} = \boldsymbol{\theta}^{(t)} - \eta_t \nabla \ell(f_{\boldsymbol{\theta}}(\boldsymbol{x}_i), \boldsymbol{y}_i)$ 

Background

## A Recipe for Gradients

1. Given training dat  $\{oldsymbol{x}_i,oldsymbol{y}_i\}_{i=1}^N$ 

2. Choose each of the

Decision function

$$\hat{\boldsymbol{y}} = f_{\boldsymbol{\theta}}(\boldsymbol{x}_i)$$

Loss function

 $\ell(\hat{oldsymbol{y}},oldsymbol{y}_i)\in\mathbb{R}$ 

**Backpropagation** can compute this gradient!

And it's a **special case of a more general algorithm** called reversemode automatic differentiation that can compute the gradient of any differentiable function efficiently!

opposite the gradient)  
$$\boldsymbol{\theta}^{(t)} = -\eta_t \nabla \ell(f_{\boldsymbol{\theta}}(\boldsymbol{x}_i), \boldsymbol{y}_i)$$

## A Recipe for

# Goals for Today's Lecture

- Explore a new class of decision functions (Neural Networks)
  - 2. Consider variants of this recipe for training

#### 2. choose each of these:

Decision function

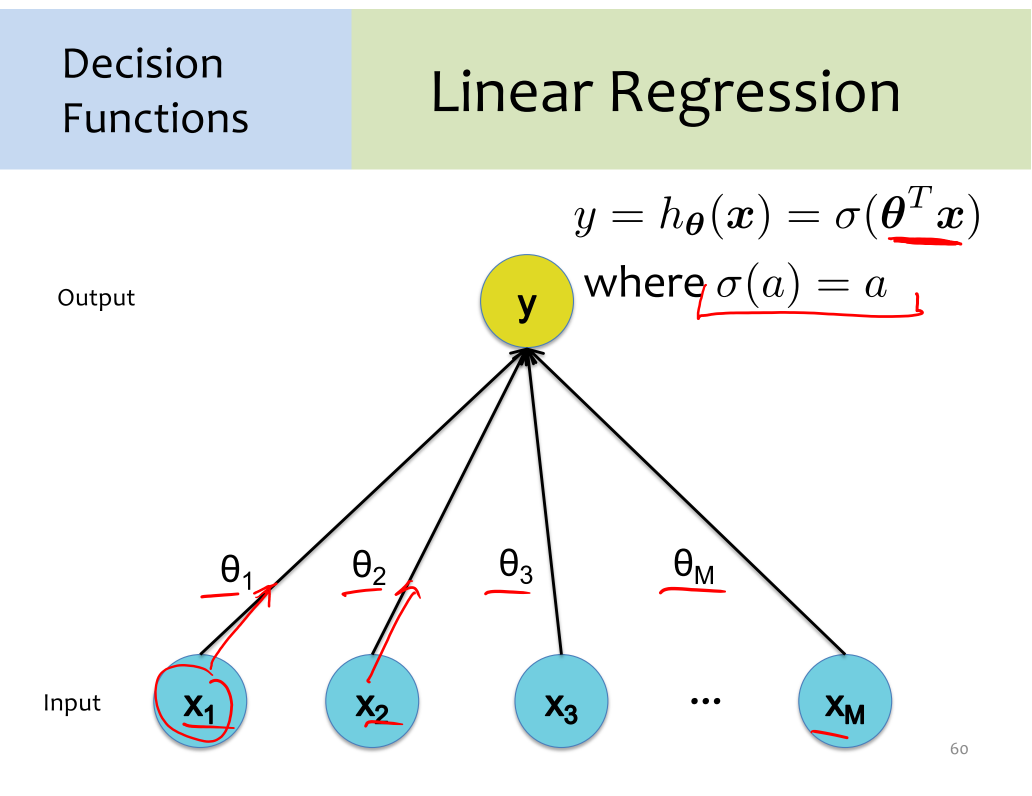
$$\hat{\boldsymbol{y}} = f_{\boldsymbol{\theta}}(\boldsymbol{x}_i)$$

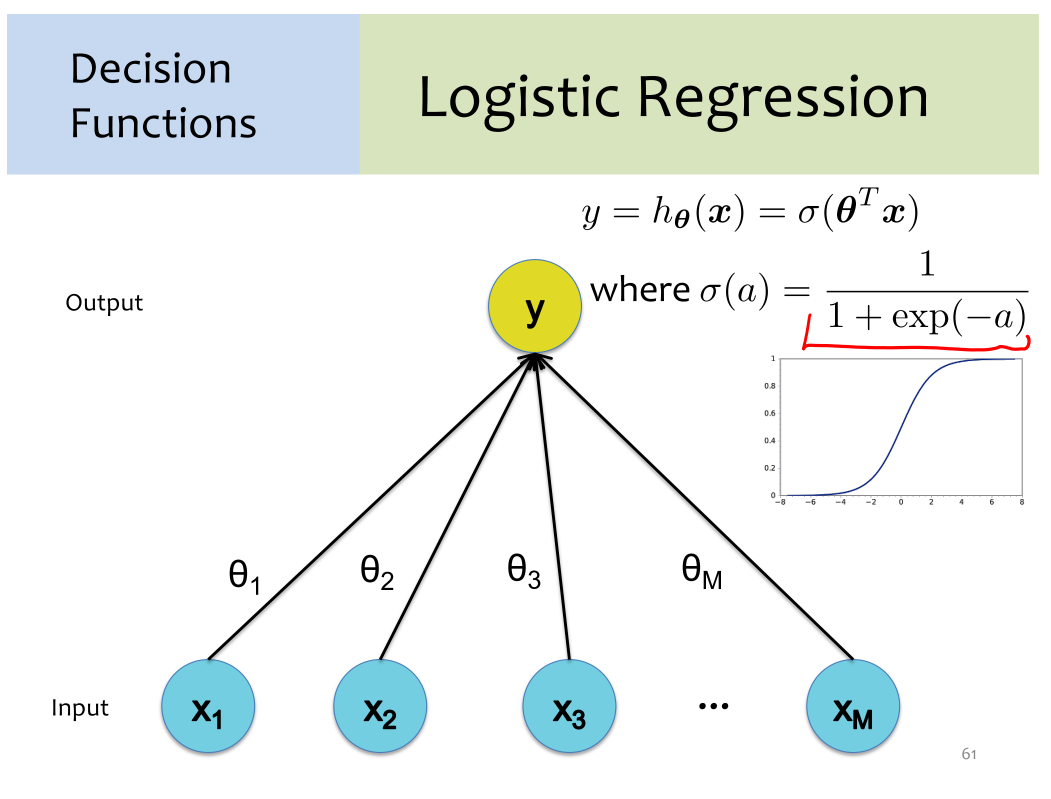
Loss function

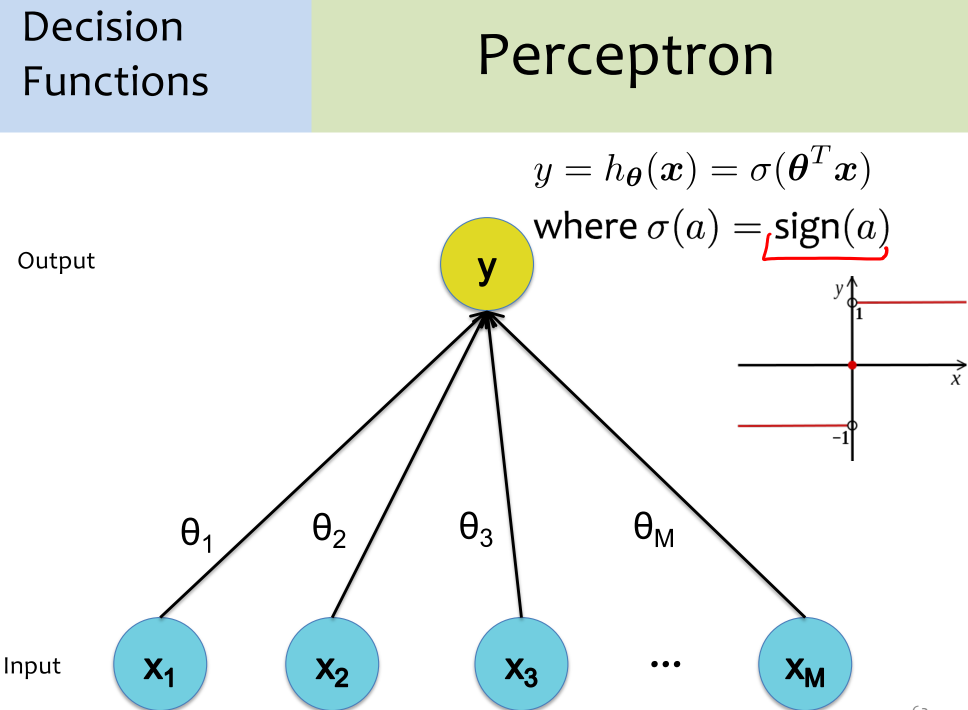
 $\ell(\hat{oldsymbol{y}},oldsymbol{y}_i)\in\mathbb{R}$ 

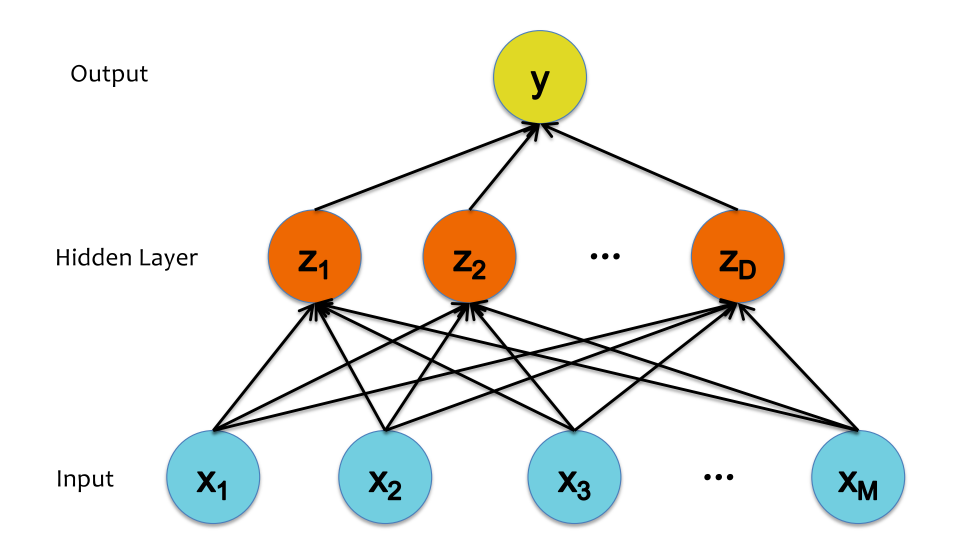
Train with SGD:
ke small steps
opposite the gradient)

 $\boldsymbol{\theta}^{(t+1)} = \boldsymbol{\theta}^{(t)} - \eta_t \nabla \ell(f_{\boldsymbol{\theta}}(\boldsymbol{x}_i), \boldsymbol{y}_i)$ 



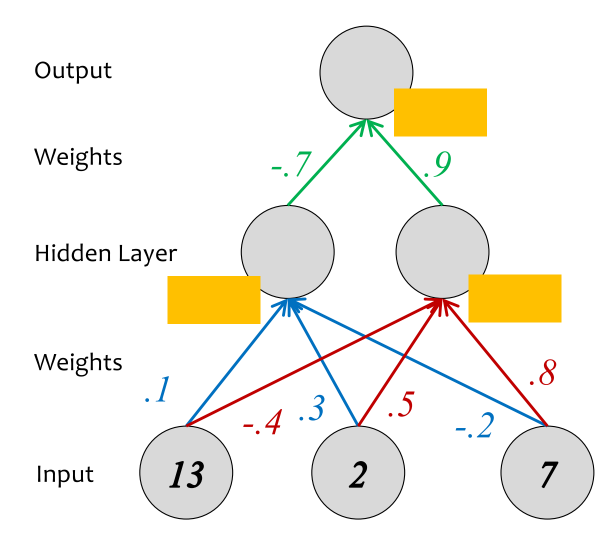






# COMPONENTS OF A NEURAL NETWORK

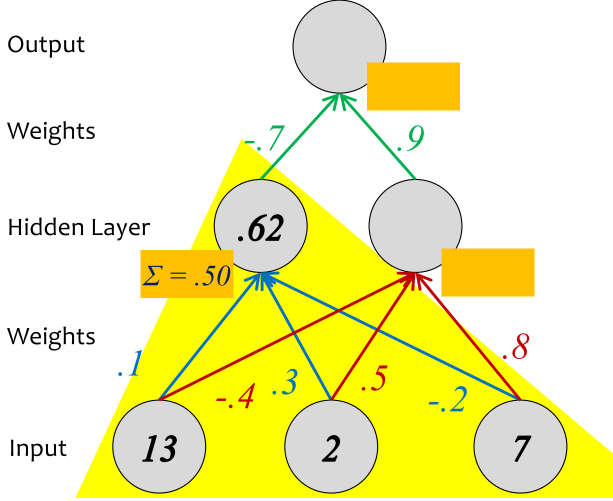
# Neural Network



Suppose we already learned the weights of the neural network.

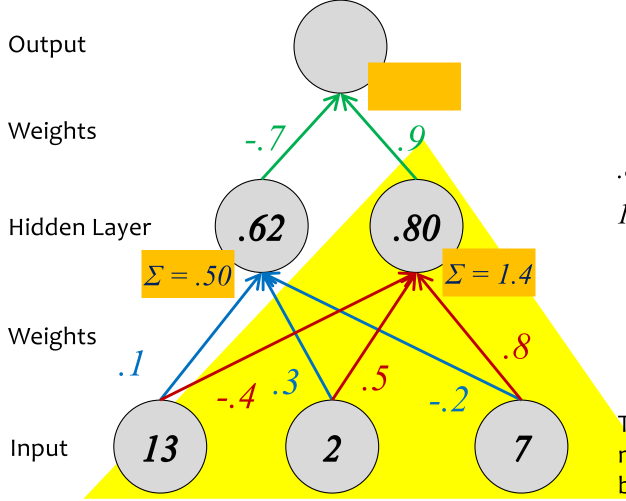
To make a new prediction, we take in some new features (aka. the input layer) and perform the feed-forward computation.

# Neural Network

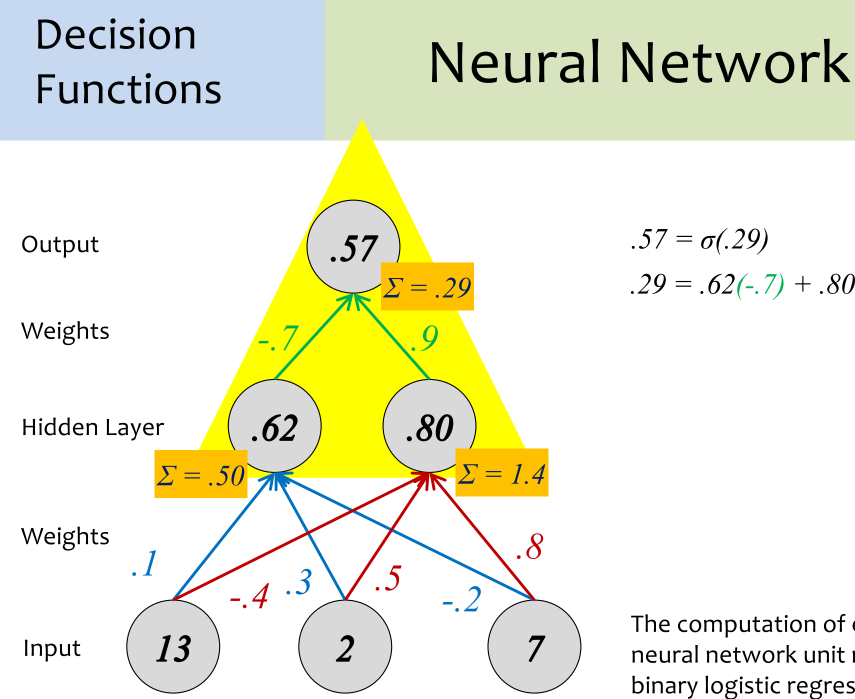


 $.62 = \sigma(.50)$ .50 = 13(.1) + 2(.3) + 7(-.2)

# Neural Network

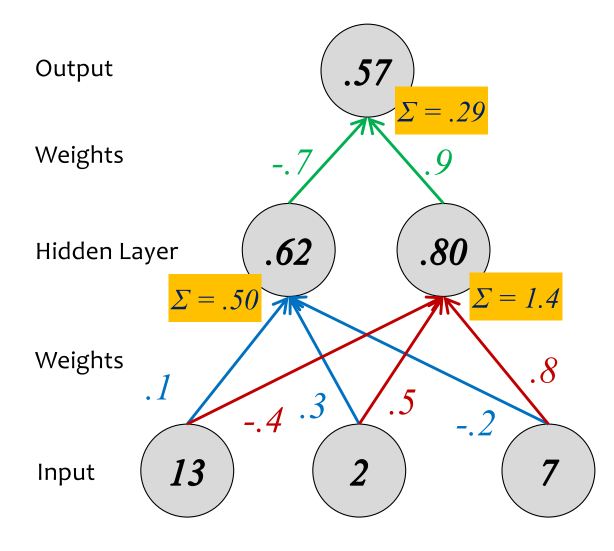


 $.80 = \sigma(1.4)$ 1.4 = 13(-.4) + 2(.5) + 7(.8)



 $.57 = \sigma(.29)$ .29 = .62(-.7) + .80(.9)

# Neural Network

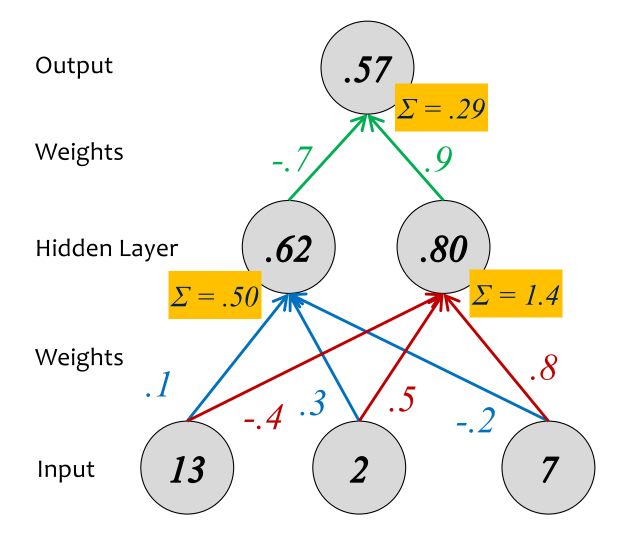


 $.57 = \sigma(.29)$ .29 = .62(-.7) + .80(.9)

 $.80 = \sigma(1.4)$ 1.4 = 13(-.4) + 2(.5) + 7(.8)

$$.62 = \sigma(.50)$$
  
 $.50 = 13(.1) + 2(.3) + 7(-.2)$ 

# Neural Network



Except we only have the target value for y at training time! We have to learn to create "useful" values of z<sub>1</sub> and z<sub>2</sub> in

the hidden layer.



# From Biological to Artificial

The motivation for Artificial Neural Networks comes from biology...

#### **Biological "Model"**

- Neuron: an excitable cell
- **Synapse:** connection between neurons
- A neuron sends an electrochemical pulse along its synapses when a sufficient voltage change occurs
- **Biological Neural Network:** collection of neurons along some pathway through the brain

#### **Biological "Computation"**

- Neuron switching time : ~ 0.001 sec
- Number of neurons:  $\sim 10^{10}$
- Connections per neuron: ~ 10<sup>4-5</sup>
- Scene recognition time: ~ 0.1 sec

#### **Artificial Model**

- Neuron: node in a directed acyclic graph (DAG)
- Weight: multiplier on each edge
- Activation Function: nonlinear thresholding function, which allows a neuron to "fire" when the input value is sufficiently high

Synapses

Dendrites

Axon

• Artificial Neural Network: collection of neurons into a DAG, which define some differentiable function

#### **Artificial Computation**

- Many neuron-like threshold switching units
- Many weighted interconnections among units
- Highly parallel, distributed processes

Nodes

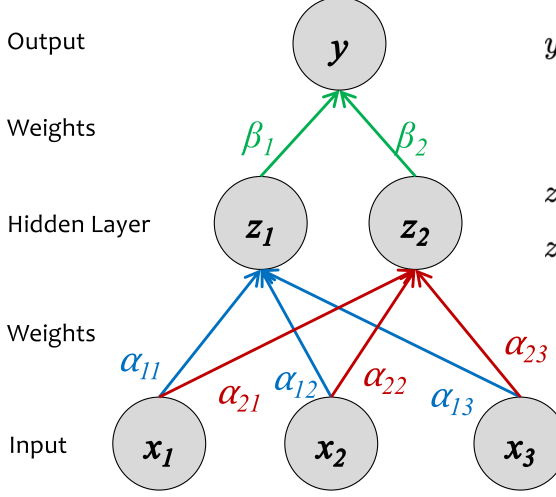
Synapses (weights)

# DEFINING A 1-HIDDEN LAYER NEURAL NETWORK

## Neural Networks

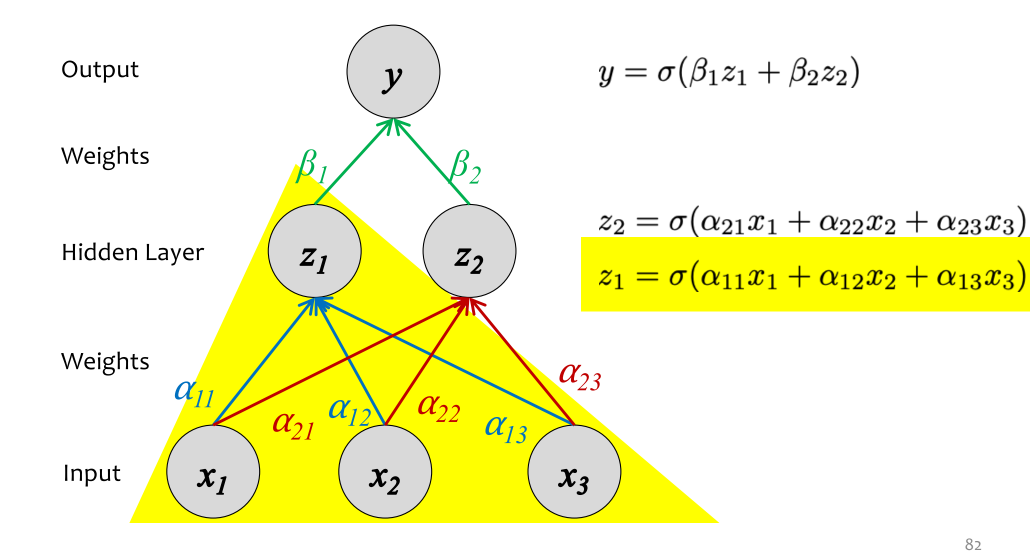
Chalkboard

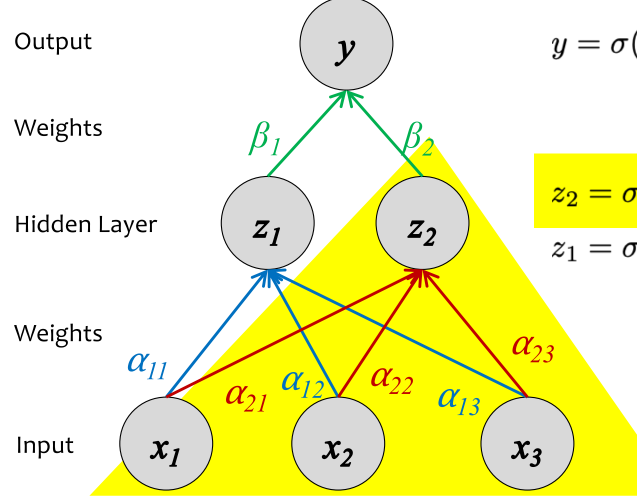
– Example: Neural Network w/1 Hidden Layer



$$y = \sigma(\beta_1 z_1 + \beta_2 z_2)$$

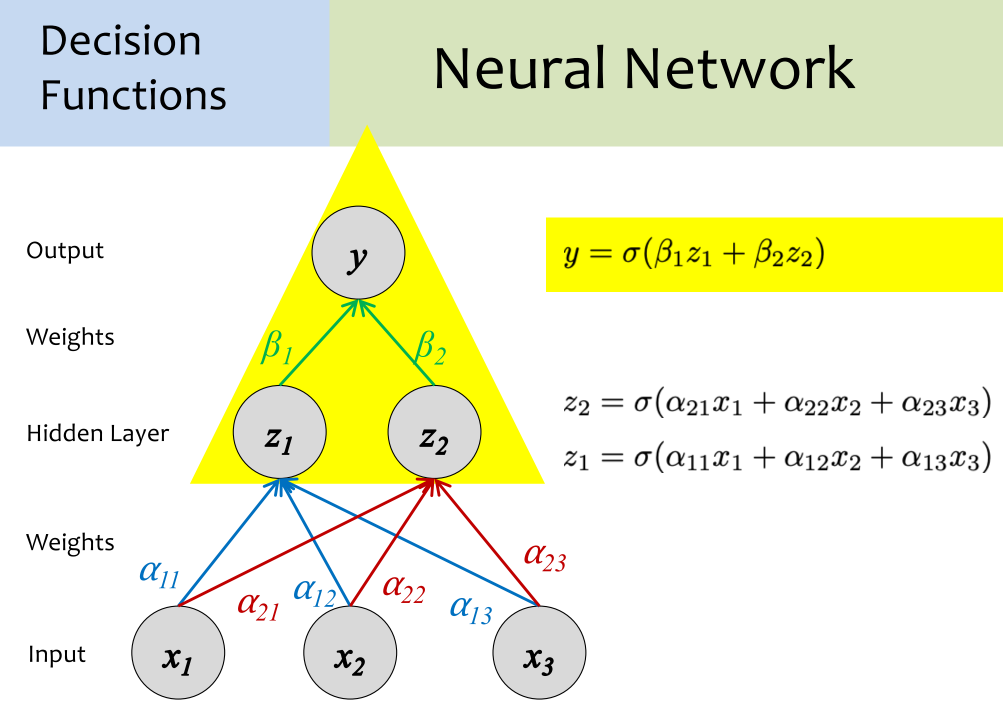
$$egin{aligned} &z_2 = \sigma(lpha_{21}x_1 + lpha_{22}x_2 + lpha_{23}x_3) \ &z_1 = \sigma(lpha_{11}x_1 + lpha_{12}x_2 + lpha_{13}x_3) \end{aligned}$$

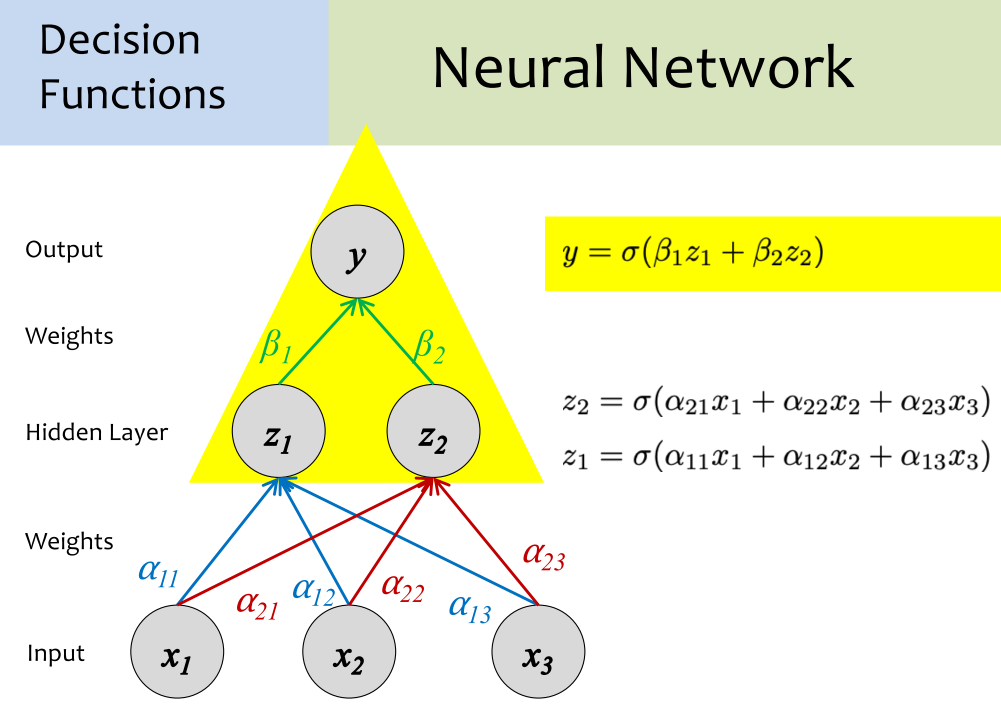


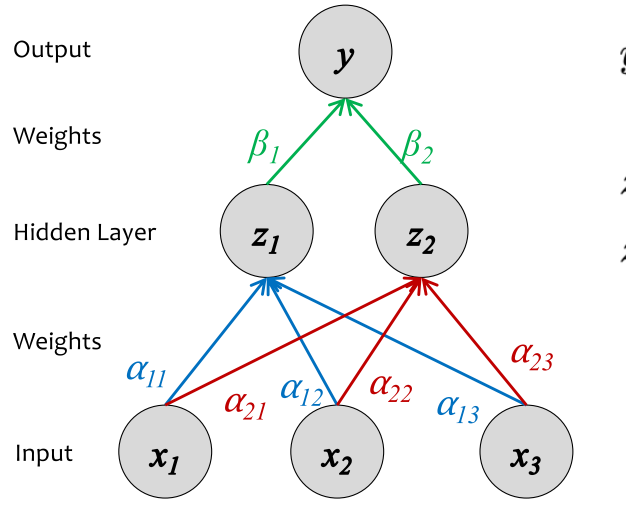


$$y = \sigma(\beta_1 z_1 + \beta_2 z_2)$$

$$egin{split} & z_2 = \sigma(lpha_{21}x_1 + lpha_{22}x_2 + lpha_{23}x_3) \ & z_1 = \sigma(lpha_{11}x_1 + lpha_{12}x_2 + lpha_{13}x_3) \end{split}$$



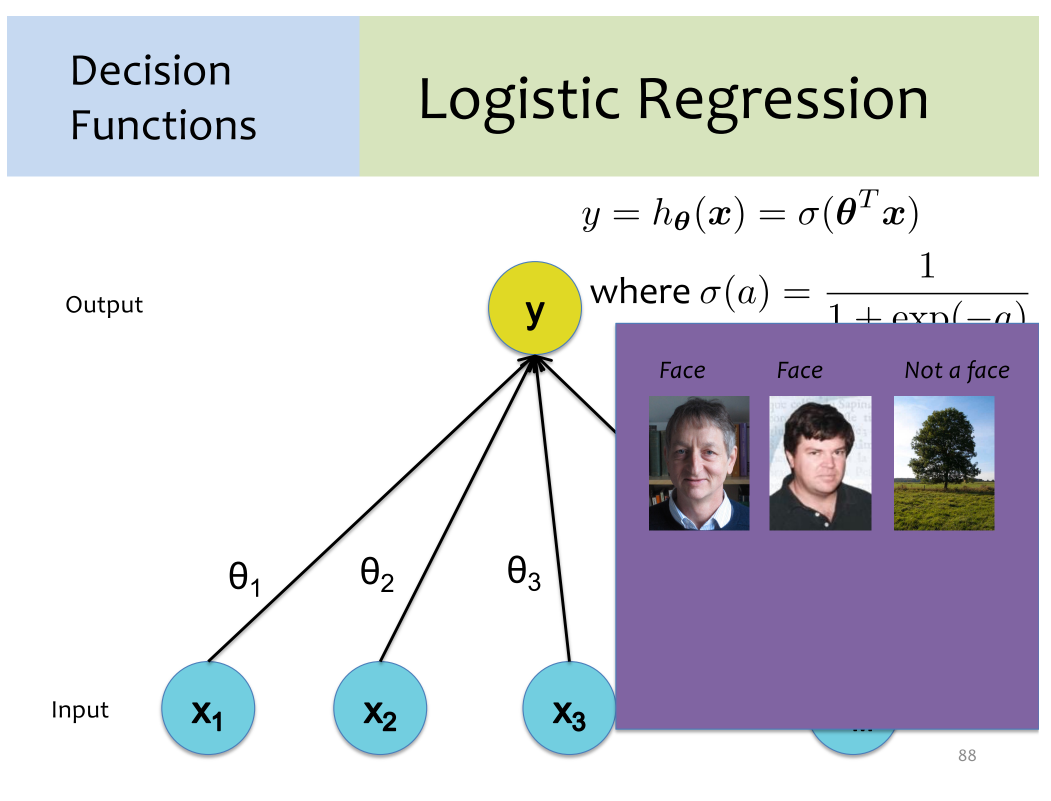


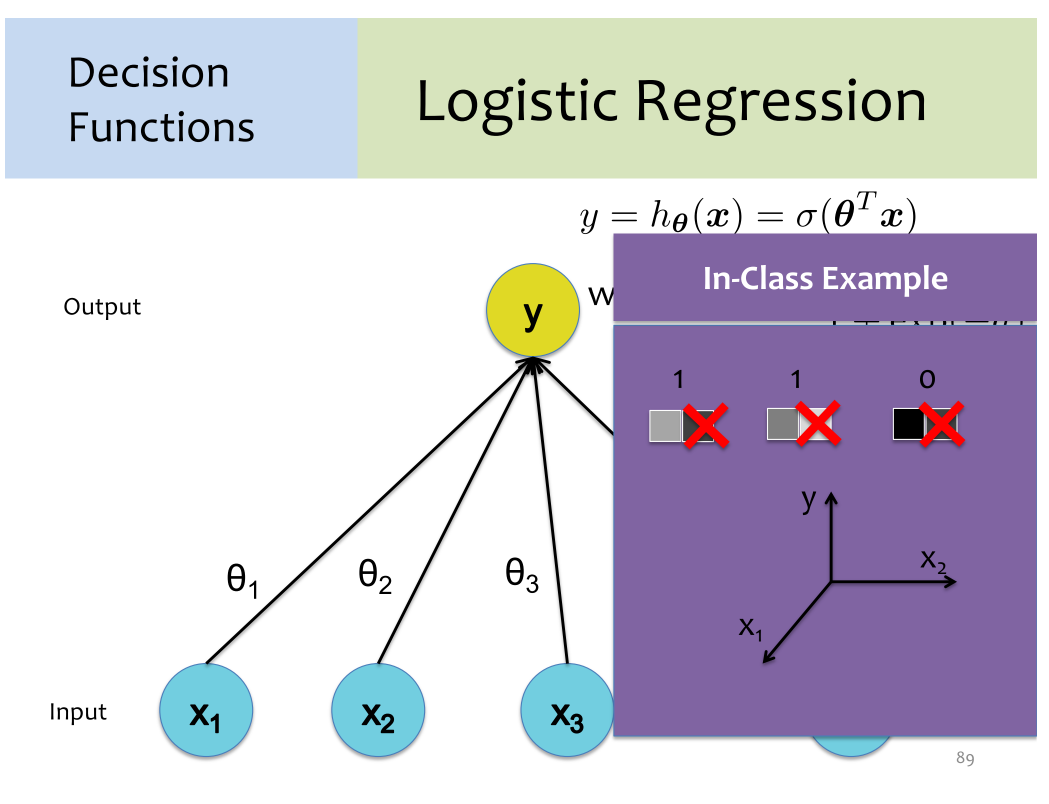


$$y = \sigma(\boldsymbol{\beta}^T \mathbf{z})$$

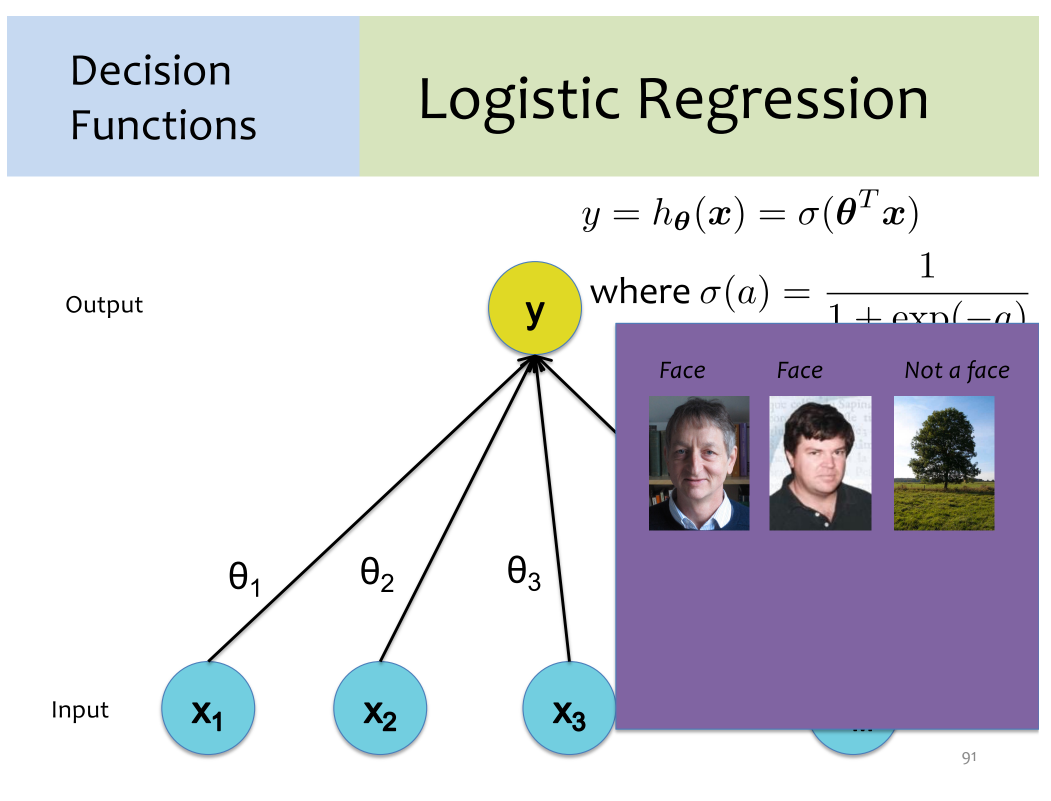
$$egin{aligned} &z_2 = \sigma(oldsymbol{lpha}_{2,\cdot}^T \mathbf{x}) \ &z_1 = \sigma(oldsymbol{lpha}_{1,\cdot}^T \mathbf{x}) \end{aligned}$$

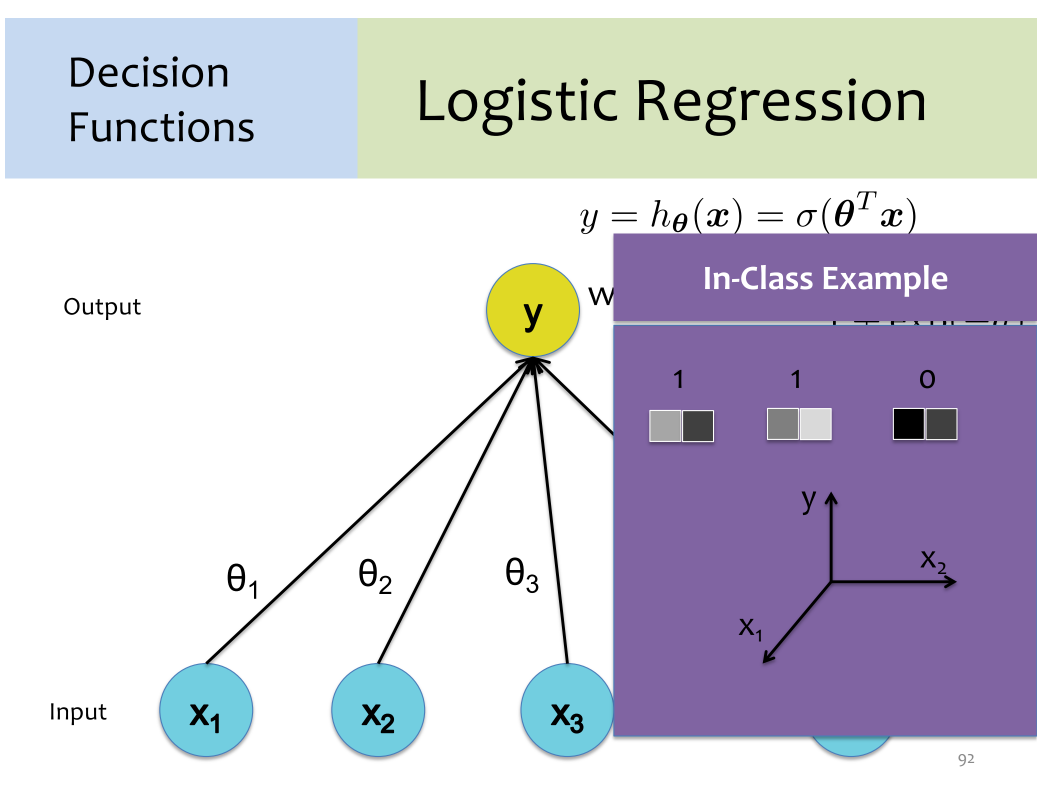
## NONLINEAR DECISION BOUNDARIES AND NEURAL NETWORKS





- Chalkboard
  - 1D Example from linear regression to logistic regression
  - 1D Example from logistic regression to a neural network





## Neural Network Parameters

## Question: 02

Suppose you are training a one-hidden layer neural network with sigmoid activations for binary classification.

True or False: There is a unique set of parameters that maximize the likelihood of the dataset above.

