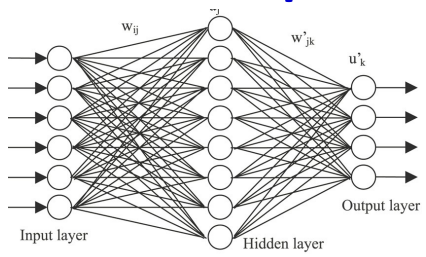
# Neural Networks Learning the network: Part 1

11-785, Fall 2021 Lecture 3

#### **Topics for the day**

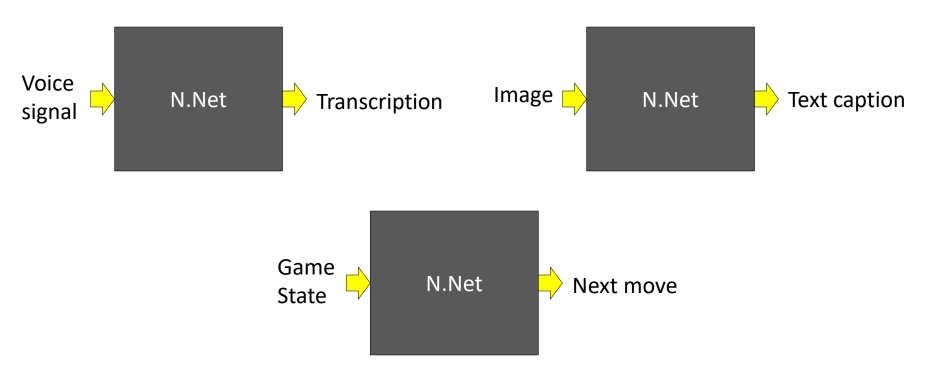
- The problem of learning
- The perceptron rule for perceptrons
  - And its inapplicability to multi-layer perceptrons
- Greedy solutions for classification networks:
   ADALINE and MADALINE
- Learning through Empirical Risk Minimization
- Intro to function optimization and gradient descent

#### Recap



- Neural networks are universal function approximators
  - Can model any Boolean function
  - Can model any classification boundary
  - Can model any continuous valued function
- Provided the network satisfies minimal architecture constraints
  - Networks with fewer than the required number of parameters can be very poor approximators

#### These boxes are functions



- Take an input
- Produce an output
- Can be modeled by a neural network!

#### Questions



- Preliminaries:
  - How do we represent the input?
  - How do we represent the output?
- How do we compose the network that performs the requisite function?

#### Questions



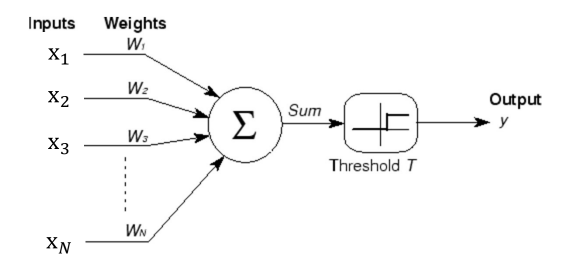
- Preliminaries:

  - How do we regin the program

    How do Abit later in the program

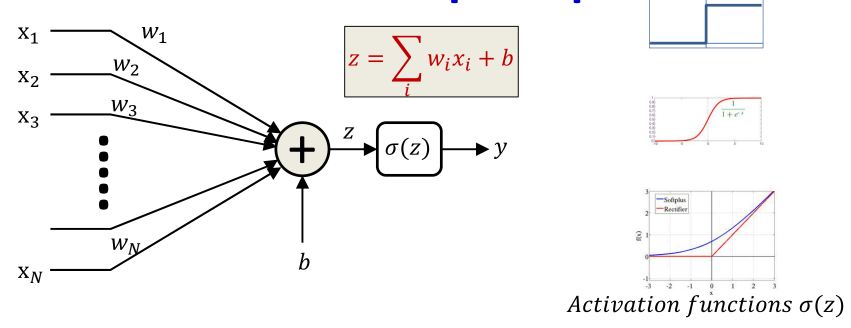
    How do Abit
- How do we compose the network that performs the requisite function?

## The original perceptron



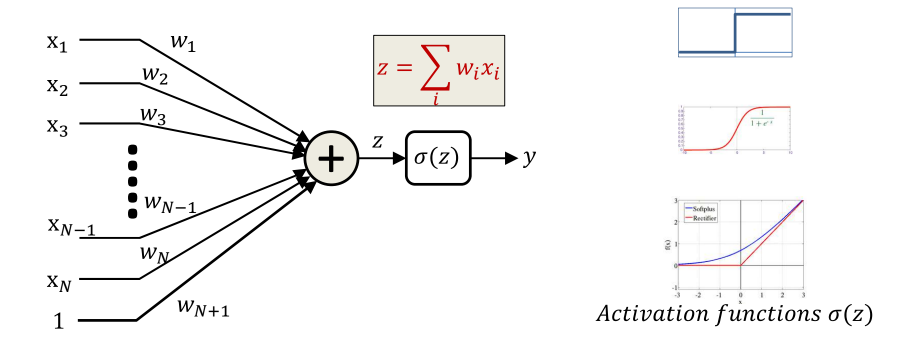
- Simple threshold unit
  - Unit comprises a set of weights and a threshold

# Preliminaries: The units in the network – the perceptron



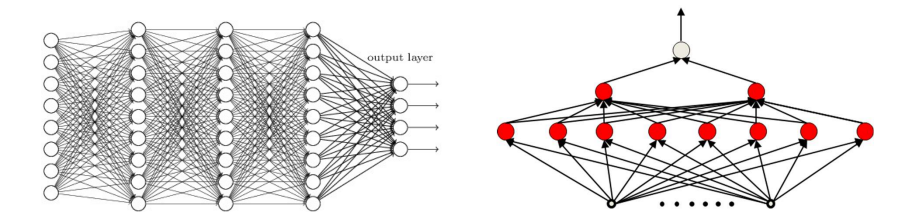
- Perceptron
  - General setting, inputs are real valued
  - A bias b representing a threshold to trigger the perceptron
  - Activation functions are not necessarily threshold functions
- The parameters of the perceptron (which determine how it behaves) are its weights and bias

#### **Preliminaries: Redrawing the neuron**



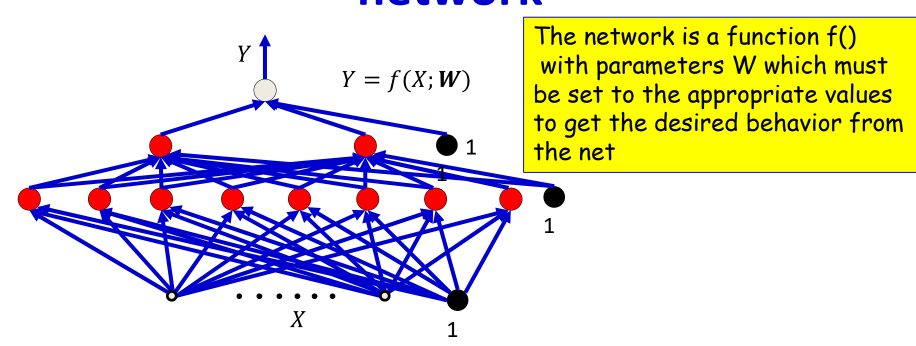
- The bias can also be viewed as the weight of another input component that is always set to 1
  - If the bias is not explicitly mentioned, we will implicitly be assuming that every perceptron has an additional input that is always fixed at 1

#### First: the structure of the network



- We will assume a *feed-forward* network
  - No loops: Neuron outputs do not feed back to their inputs directly or indirectly
  - Loopy networks are a future topic
- Part of the design of a network: The architecture
  - How many layers/neurons, which neuron connects to which and how, etc.
- For now, assume the architecture of the network is capable of representing the needed function

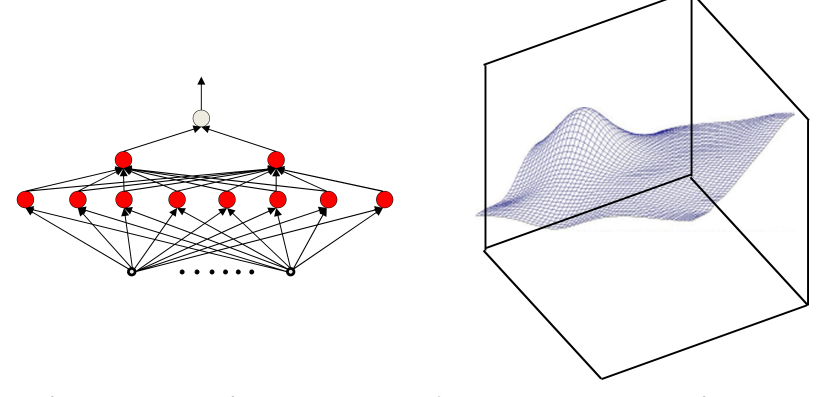
# What we learn: The parameters of the network



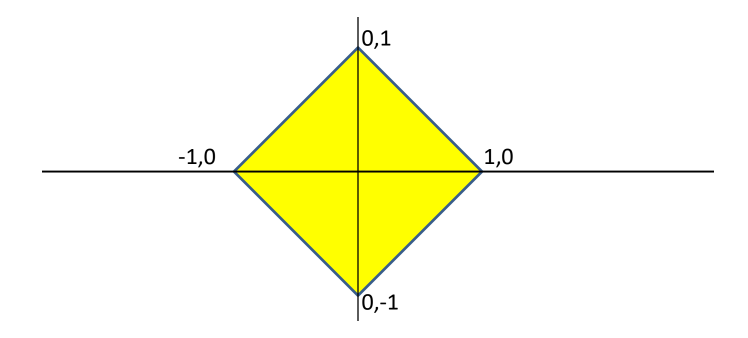
- **Given:** the architecture of the network
- The parameters of the network: The weights and biases
  - The weights associated with the blue arrows in the picture
- Learning the network: Determining the values of these parameters such that the network computes the desired function

• Moving on..

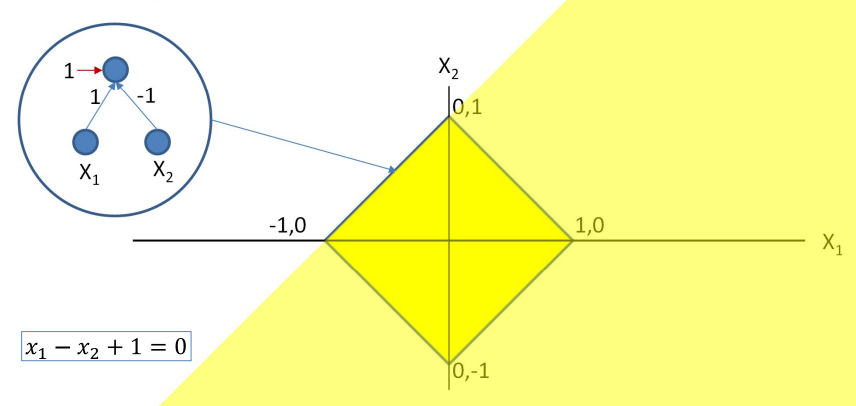
## The MLP can represent anything

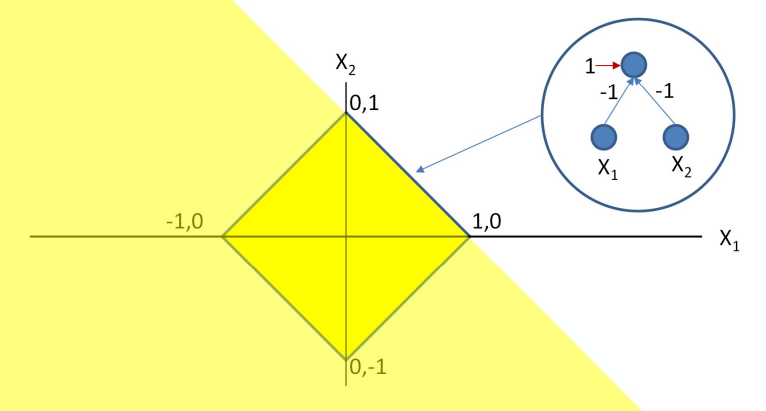


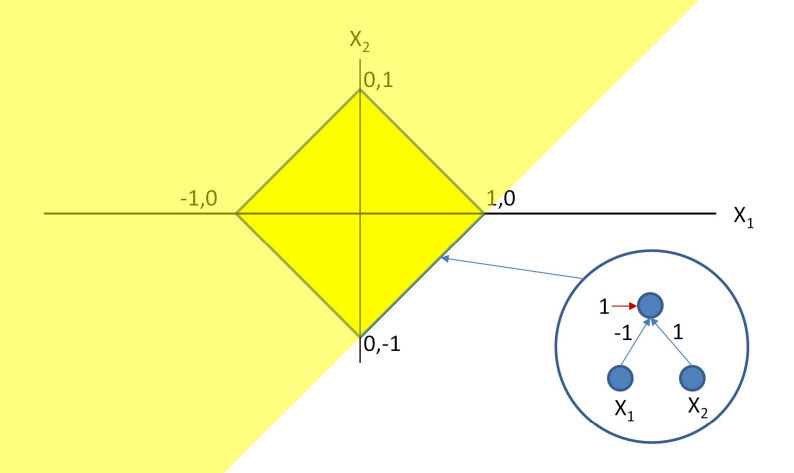
- The MLP can be constructed to represent anything
- But how do we construct it?



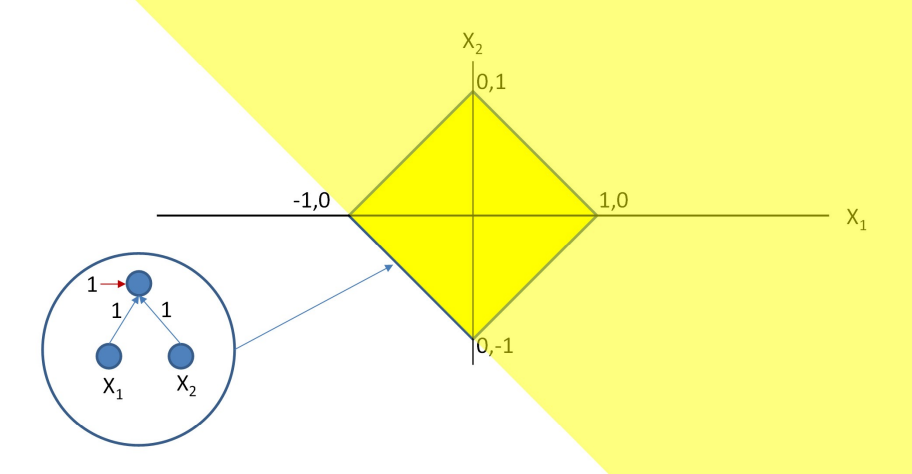
- Given a function, handcraft a network to satisfy it
- E.g.: Build an MLP to classify this decision boundary



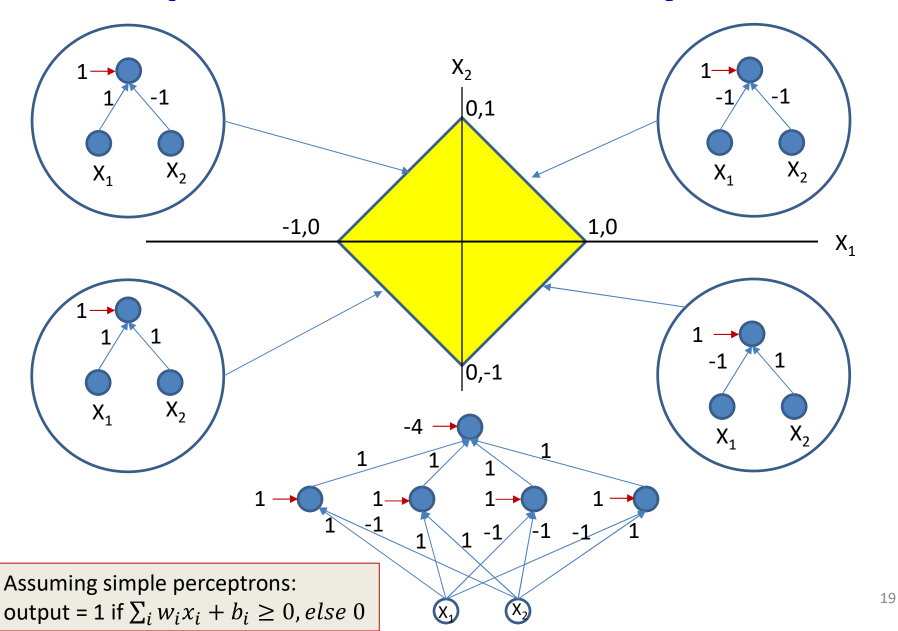


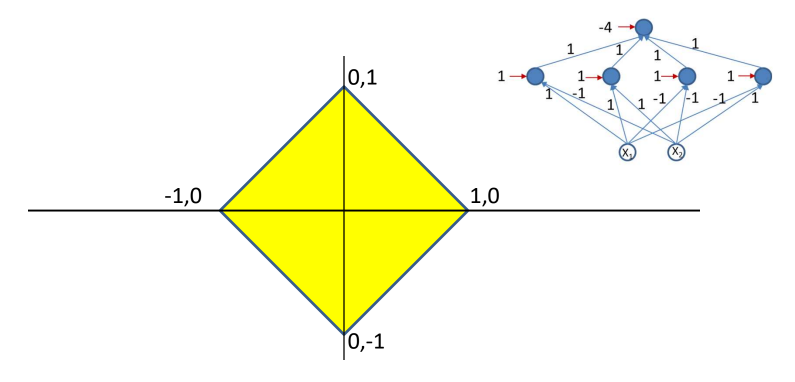


Assuming simple perceptrons: output = 1 if  $\sum_i w_i x_i + b_i \ge 0$ , else 0



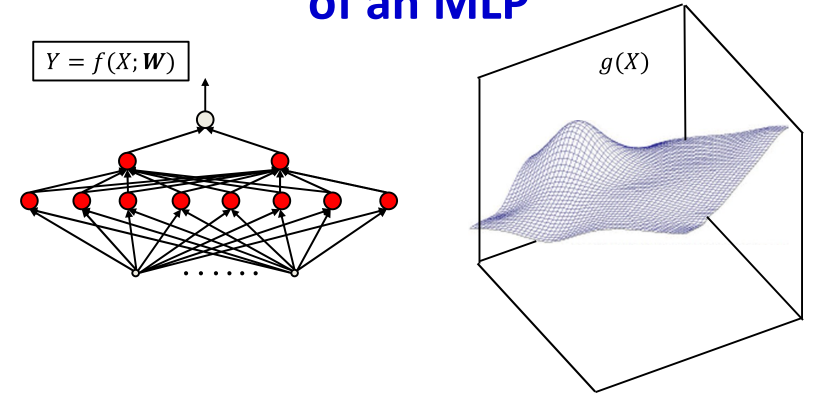
Assuming simple perceptrons: output = 1 if  $\sum_i w_i x_i + b_i \ge 0$ , else 0





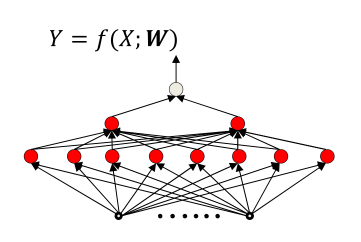
- Given a function, handcraft a network to satisfy it
- E.g.: Build an MLP to classify this decision boundary
- Not possible for all but the simplest problems..

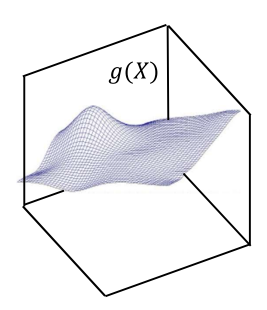
# Option 2: Automatic estimation of an MLP



• More generally, given the function g(X) to model, we can derive the parameters of the network to model it, through computation

#### How to learn a network?



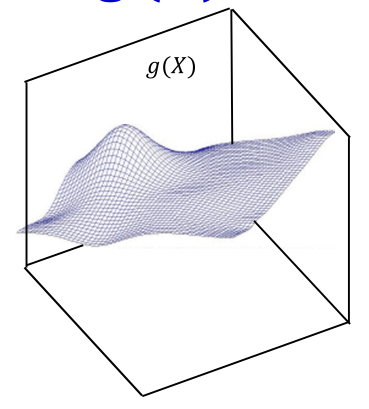


• When f(X; W) has the capacity to exactly represent g(X)

$$\widehat{\boldsymbol{W}} = \underset{W}{\operatorname{argmin}} \int_{X} div(f(X; W), g(X))dX$$

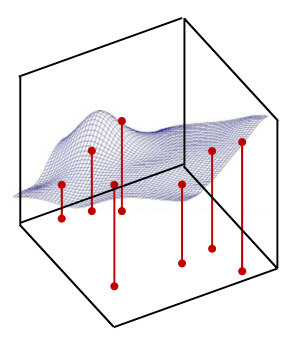
• div() is a divergence function that goes to zero when f(X; W) = g(X)

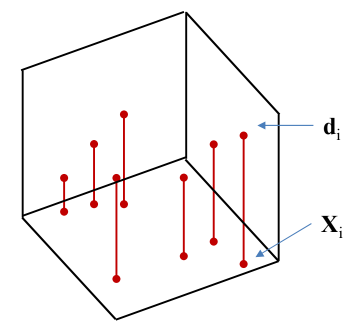
Problem g(X) is unknown



- Function g(X) must be fully specified
  - Known everywhere, i.e. for every input X
- In practice we will not have such specification

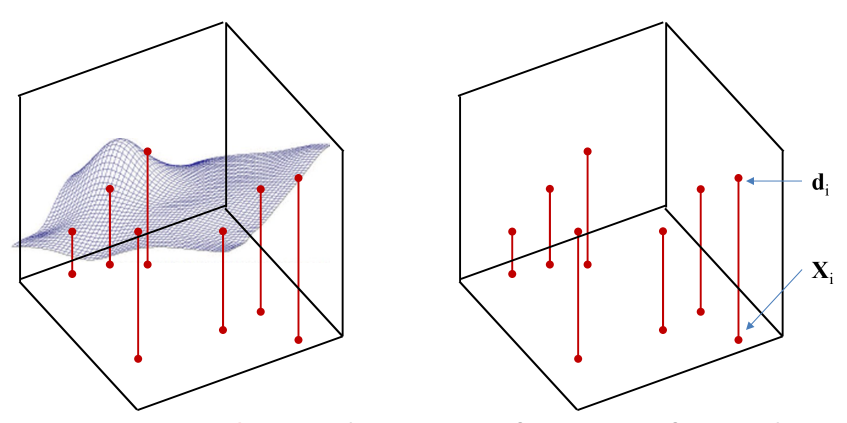
## Sampling the function





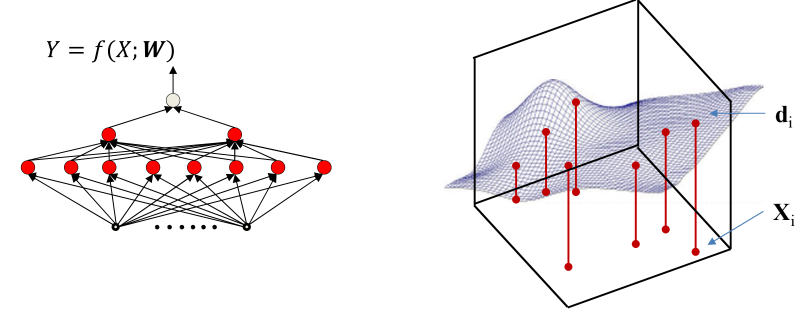
- Sample g(X)
  - Basically, get input-output pairs for a number of samples of input  $X_i$ 
    - Many samples  $(X_i, d_i)$ , where  $d_i = g(X_i) + noise$
- Very easy to do in most problems: just gather training data
  - E.g. set of images and their class labels
  - E.g. speech recordings and their transcription

## **Drawing samples**



- We must *learn* the *entire* function from these few examples
  - The "training" samples

## **Learning the function**



- Estimate the network parameters to "fit" the training points exactly
  - Assuming network architecture is sufficient for such a fit
  - Assuming unique output d at any  $\mathbf{X}$ 
    - And hopefully the resulting function is also correct where we don't have training samples

#### Story so far

- "Learning" a neural network == determining the parameters of the network (weights and biases) required for it to model a desired function
  - The network must have sufficient capacity to model the function
- Ideally, we would like to optimize the network to represent the desired function everywhere
- However this requires knowledge of the function everywhere
- Instead, we draw "input-output" training instances from the function and estimate network parameters to "fit" the input-output relation at these instances
  - And hope it fits the function elsewhere as well

## Poll 1

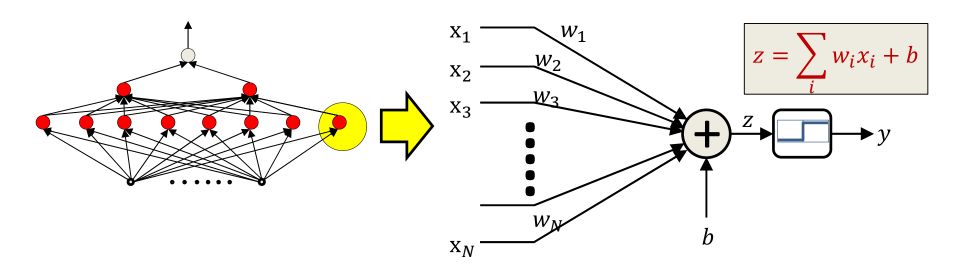
#### Poll 1

- Since neural networks are universal approximators, any network of any architecture can approximate any function to arbitrary precision (True or False):
  - True
  - False
- Which of the following are true regarding how to compose a network to approximate a given function?
  - The network architecture must have sufficient capacity to model the function
  - The network is actually a parametric function, whose parameters are its weights and biases
  - The parameters must be learned to best approximate the target function
  - The parameters can be perfectly learned from just a few training samples of the target function, even if the actual target function is unknown.

## Let's begin with a simple task

- Learning a *classifier* 
  - Simpler than regressions
- This was among the earliest problems addressed using MLPs
- Specifically, consider binary classification
  - Generalizes to multi-class

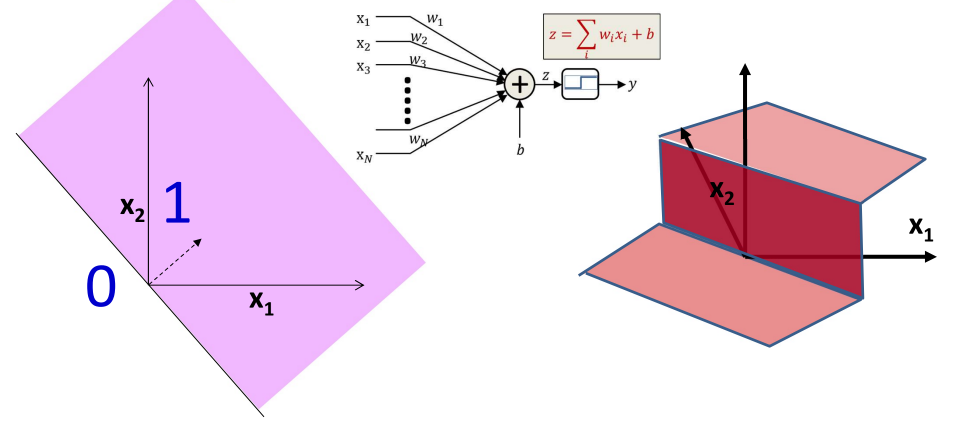
## **History: The original MLP**



- The original MLP as proposed by Minsky: a network of threshold units
  - But how do you train it?
    - Given only "training" instances of input-output pairs



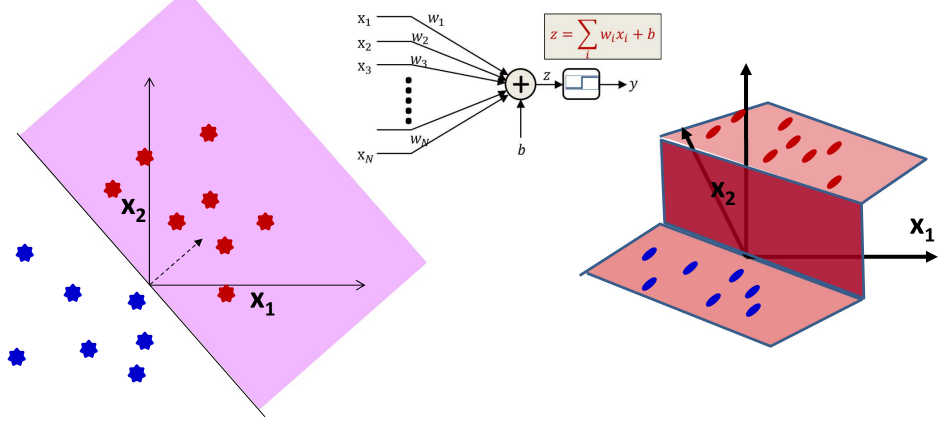
#### The simplest MLP: a single perceptron



- Learn this function
  - A step function across a hyperplane

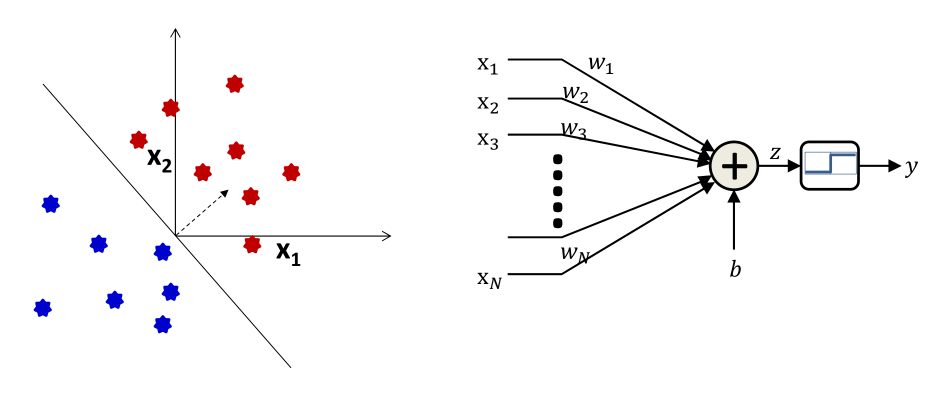


#### The simplest MLP: a single perceptron



- Learn this function
  - A step function across a hyperplane
  - Given only samples from it

#### Learning the perceptron

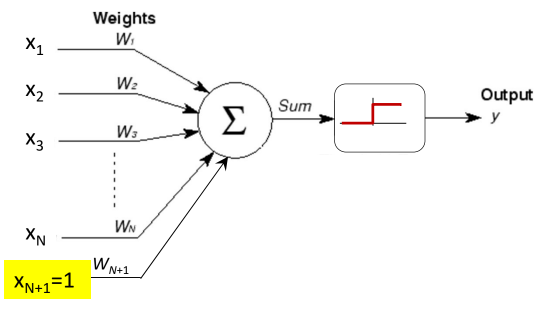


• Given a number of input output pairs, learn the weights and bias

$$- y = \begin{cases} 1 & if & \sum_{i=1}^{N} w_i X_i + b \ge 0 \\ 0 & otherwise \end{cases}$$
 Boundary:  $\sum_{i=1}^{N} w_i X_i + b = 0$ 

- Learn  $W = [w_1..w_N]^T$  and b, given several (X, y) pairs

#### Restating the perceptron

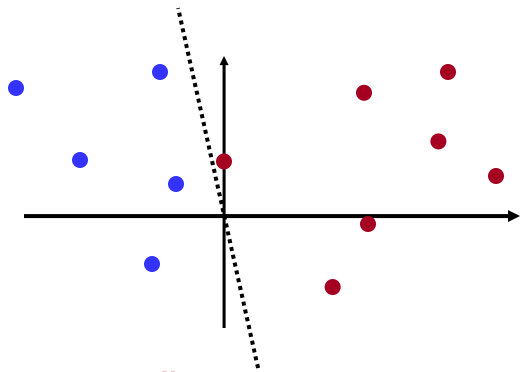


Restating the perceptron equation by adding another dimension to X

$$y = \begin{cases} 1 & if \sum_{i=1}^{N+1} w_i X_i \geq 0 \\ 0 & otherwise \end{cases}$$
 where  $X_{N+1} = 1$ 

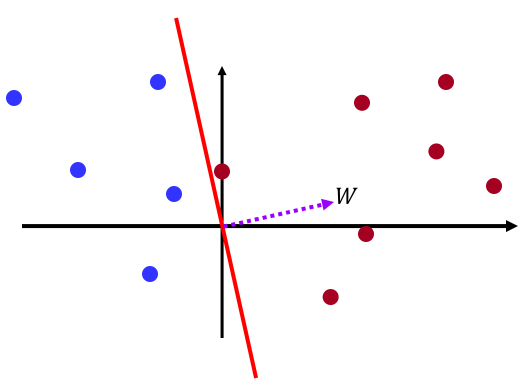
Note that the boundary  $\sum_{i=1}^{N+1} w_i X_i = 0$  is now a hyperplane through origin

## **The Perceptron Problem**



• Find the hyperplane  $\sum_{i=1}^{N+1} w_i X_i = 0$  that perfectly separates the two groups of points

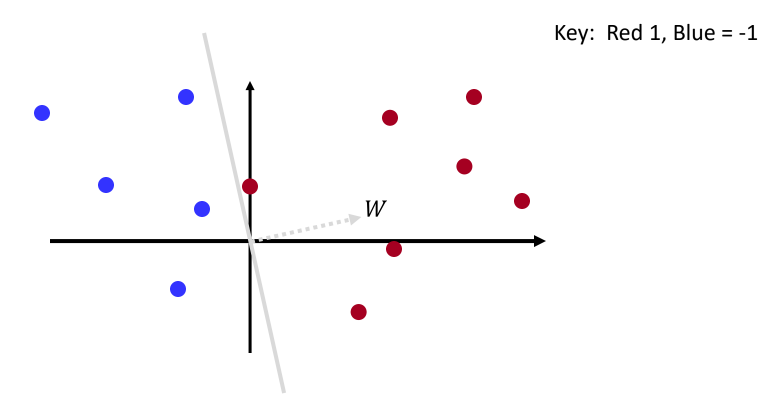
#### **The Perceptron Problem**



Key: Red 1, Blue = -1

- Find the hyperplane  $\sum_{i=1}^{N+1} w_i X_i = 0$  that perfectly separates the two groups of points
  - Let vector  $W = [w_1, w_2, ..., w_{N+1}]^T$  and vector  $X = [x_1, x_2, ..., x_N, 1]^T$
  - $-\sum_{i=1}^{N+1} w_i X_i = W^T X$  is an inner product
  - $W^T X = 0$  is the hyperplane comprising all X orthogonal to vector W
    - Learning the perceptron = finding the weight vector W for the separating hyperplane
    - W points in the direction of the positive class

#### **The Perceptron Problem**



- Learning the perceptron: Find the weights vector W such that the plane described by  $W^TX=0$  perfectly separates the classes
  - $-W^TX$  is positive for all red dots and negative for all blue ones

#### The online perceptron solution

- The popular solution, originally proposed by Rosenblatt is an *online* algorithm
  - The famous "perceptron" algorithm
- Initializes W and incrementally updates it each time we encounter an instance that is incorrectly classified
  - Guaranteed to find the correct solution for linearly separable data
  - On following slides, but will not cover in class

#### Perceptron Algorithm: Summary

- Cycle through the training instances
- Only update W on misclassified instances
- If instance misclassified:
  - If instance is positive class (positive misclassified as negative)

$$W = W + X_i$$

If instance is negative class (negative misclassified as positive)

$$W = W - X_i$$

### **Perceptron Learning Algorithm**

• Given N training instances  $(X_1, y_1), (X_2, y_2), \dots, (X_N, y_N)$ 

$$- y_i = +1 \text{ or } -1$$

- Initialize W
- Cycle through the training instances:

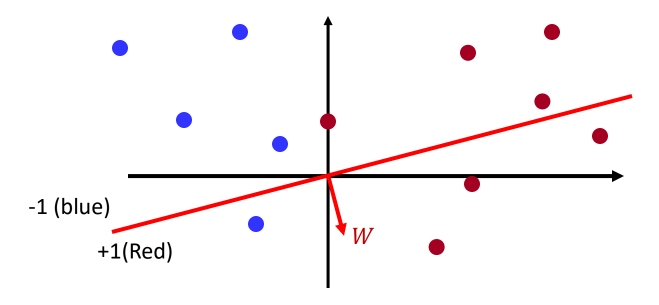
Using a +1/-1 representation for classes to simplify notation

do

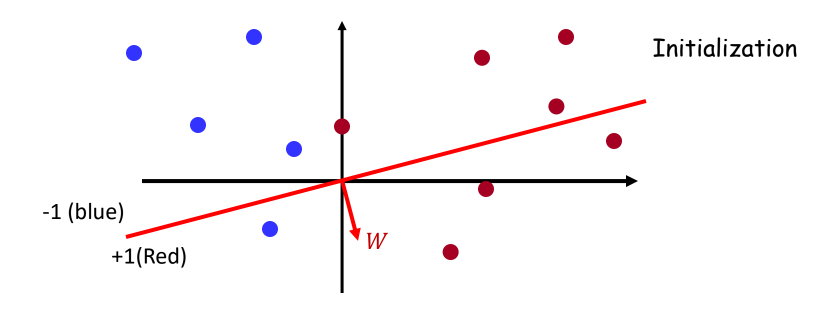
- For 
$$i=1..N_{train}$$
 
$$O(X_i) = sign(\mathbf{W}^T X_i)$$
 • If  $O(X_i) \neq y_i$  
$$W = W + y_i X_i$$

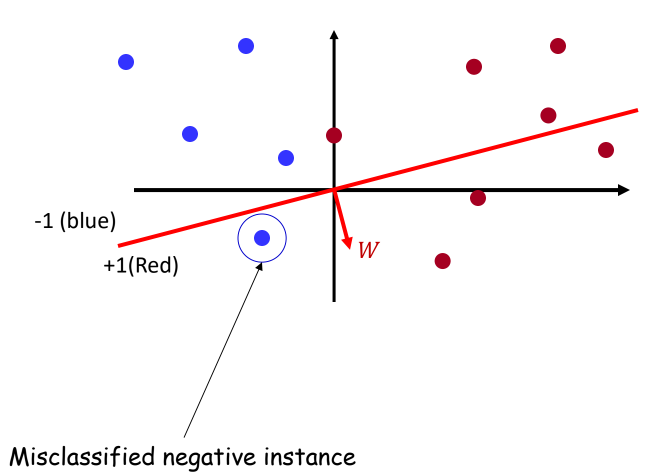
until no more classification errors

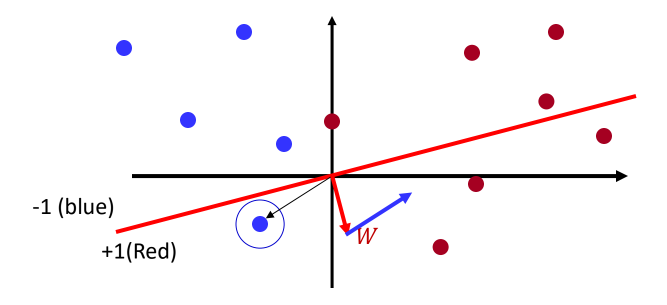
# A Simple Method: The Perceptron Algorithm



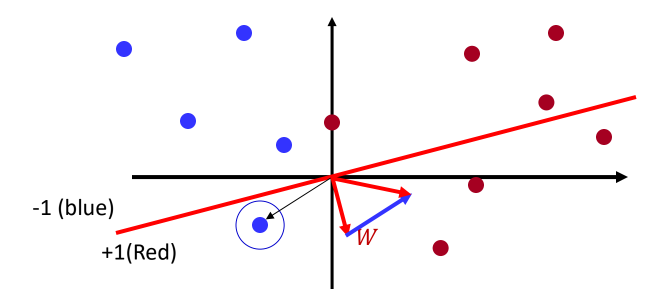
- Initialize: Randomly initialize the hyperplane
  - I.e. randomly initialize the normal vector W
- Classification rule  $sign(W^TX)$ 
  - Vectors on the same side of the hyperplane as W will be assigned +1 class,
     and those on the other side will be assigned -1
- The random initial plane will make mistakes



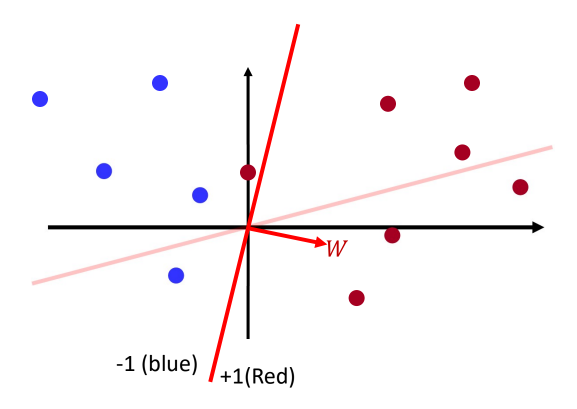




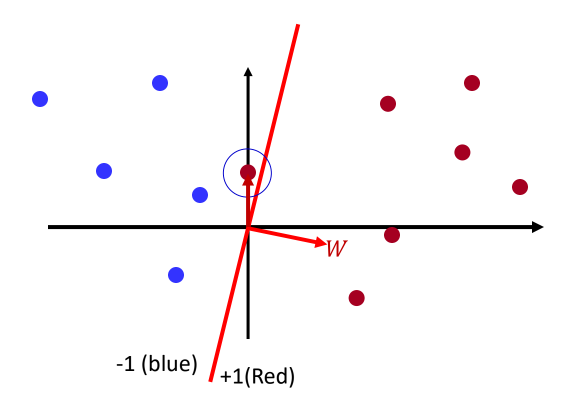
Misclassified negative instance, subtract it from W



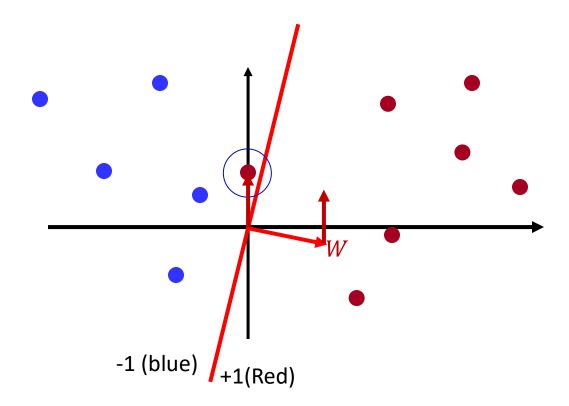
The new weight



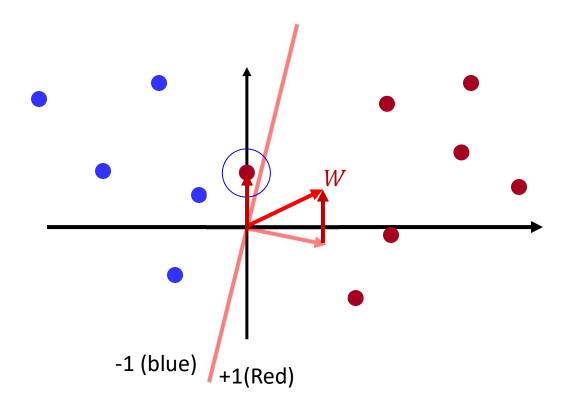
The new weight (and boundary)



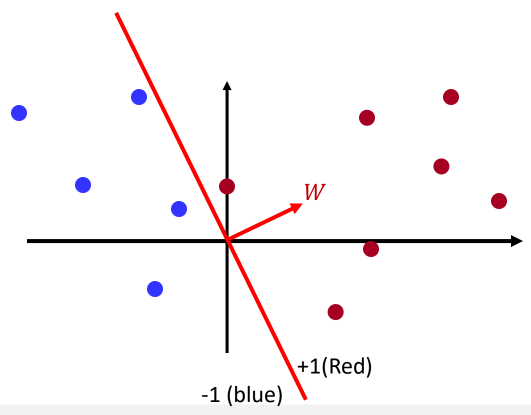
Misclassified positive instance



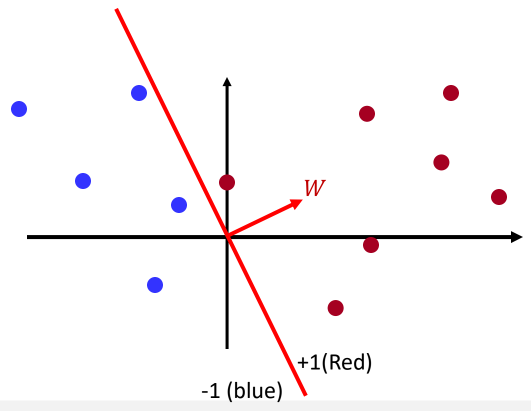
Misclassified positive instance, add it to W



The new weight vector



The new decision boundary
Perfect classification, no more updates, we are done

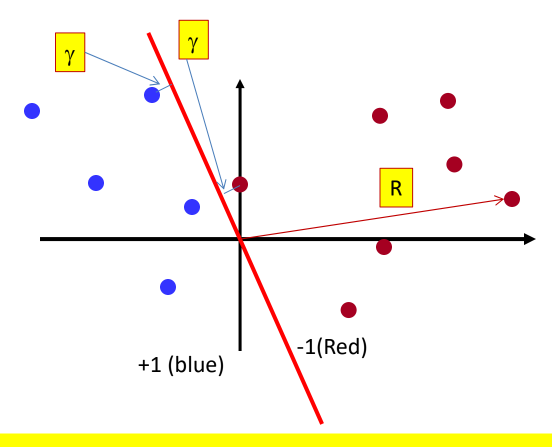


The new decision boundary
Perfect classification, no more updates, we are done

If the classes are linearly separable, guaranteed to converge in a finite number of steps

#### **Convergence of Perceptron Algorithm**

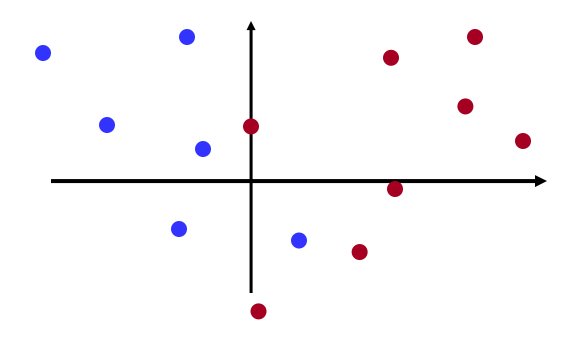
- Guaranteed to converge if classes are linearly separable
  - After no more than  $\left(\frac{R}{\gamma}\right)^2$  misclassifications
    - Specifically when W is initialized to 0
  - -R is length of longest training point
  - $-\gamma$  is the *best case* closest distance of a training point from the classifier
    - Same as the margin in an SVM
  - Intuitively takes many increments of size  $\gamma$  to undo an error resulting from a step of size R



 $\gamma$  is the best-case margin R is the length of the longest vector

# The Perceptron Solution: when classes are not linearly separable

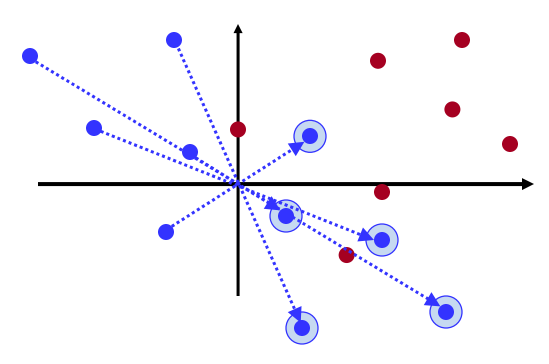
Key: Red 1, Blue = -1



- When classes are not linearly separable, not possible to find a separating hyperplane
  - No "support" plane for reflected data
  - Some points will always lie on the other side
- Model does not support perfect classification of this data

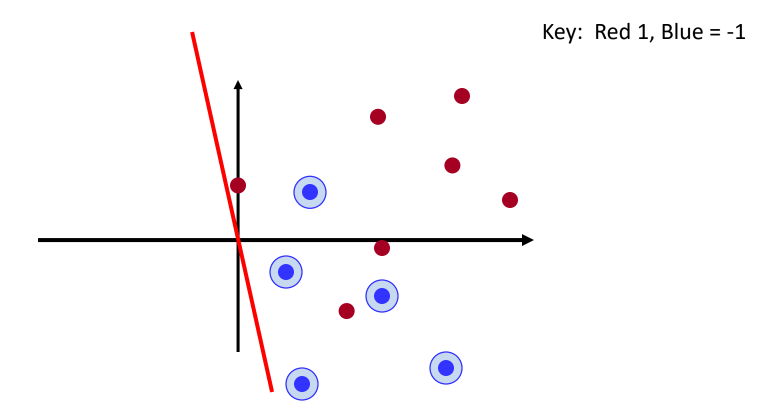
#### A simpler solution

Key: Red 1, Blue = -1



- Reflect all the negative instances across the origin
  - Negate every component of vector X
- If we use class  $y \in \{+1, -1\}$  notation for the labels (instead of  $y \in \{0, 1\}$ ), we can simply write the "reflected" values as X' = yX
  - Will retain the features X for the positive class, but reflect/negate them for the negative class

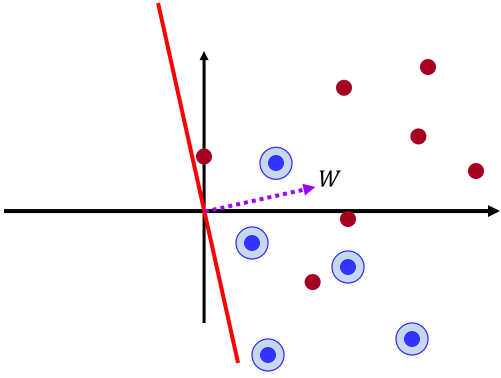
#### **The Perceptron Solution**



- Learning the perceptron: Find a plane such that all the modified (X') features lie on one side of the plane
  - Such a plane can always be found if the classes are linearly separable

## The Perceptron Solution: Linearly separable case

Key: Red 1, Blue = -1



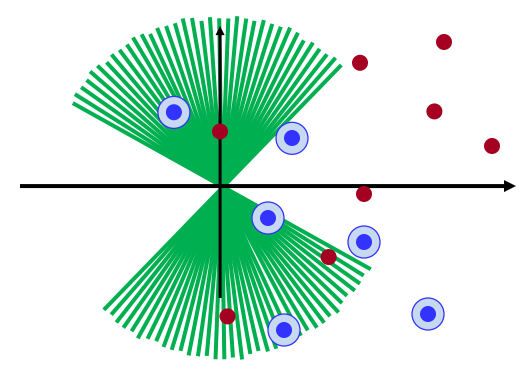
• When classes are linearly separable: a trivial solution

$$W = \frac{1}{N} \sum_{i} X_i' \qquad = \frac{1}{N} \sum_{i} y_i X_i$$

Other solutions are also possible, e.g. max-margin solution

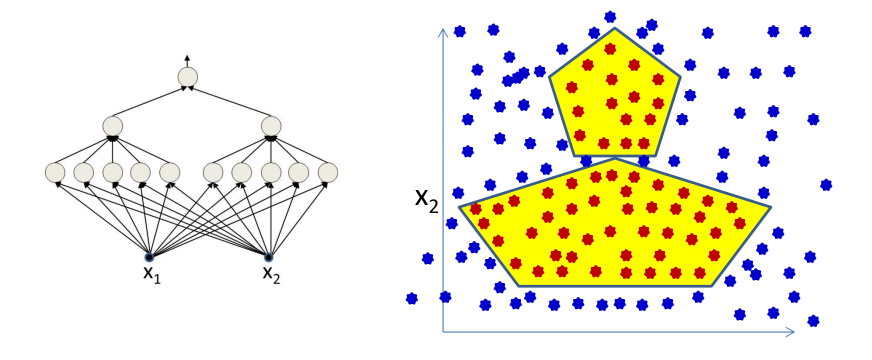
# The Perceptron Solution: when classes are not linearly separable

Key: Red 1, Blue = -1



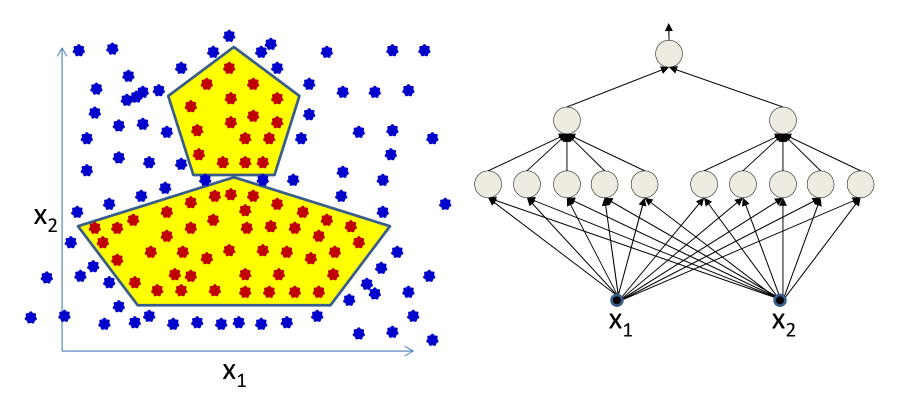
- When classes are not linearly separable, not possible to find a separating hyperplane
  - No "support" plane for reflected data
  - Some points will always lie on the other side
- Model does not support perfect classification of this data

#### **History: A more complex problem**

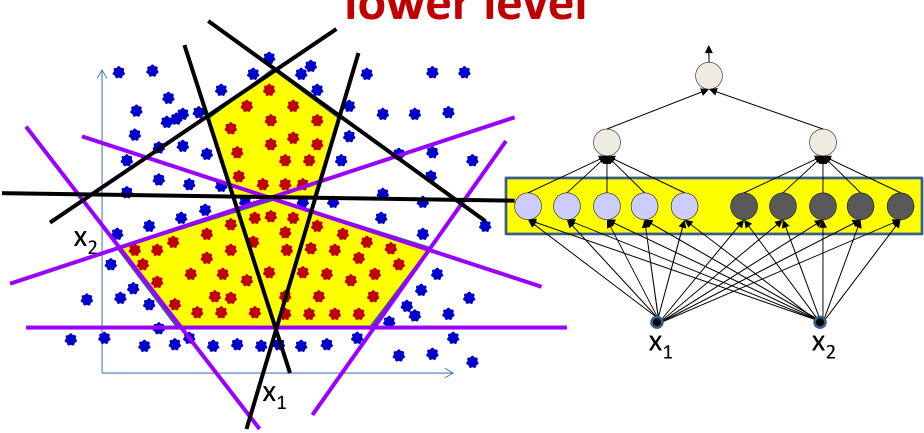


- Learn an MLP for this function
  - 1 in the yellow regions, 0 outside
- Using just the samples
- We know this can be perfectly represented using an MLP

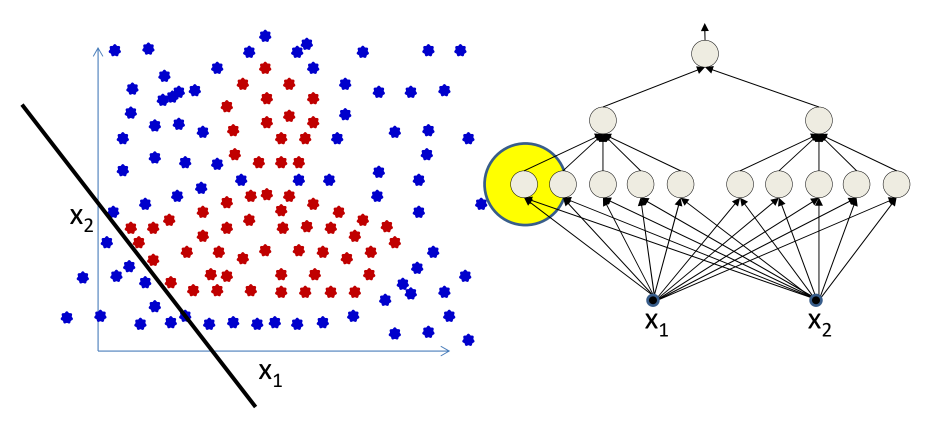
#### More complex decision boundaries



- Even using the perfect architecture...
- ... can we use perceptron learning rules to learn this classification function?



- The lower-level neurons are linear classifiers
  - They require linearly separated labels to be learned
  - The actually provided labels are not linearly separated
  - Challenge: Must also learn the labels for the lowest units! 62

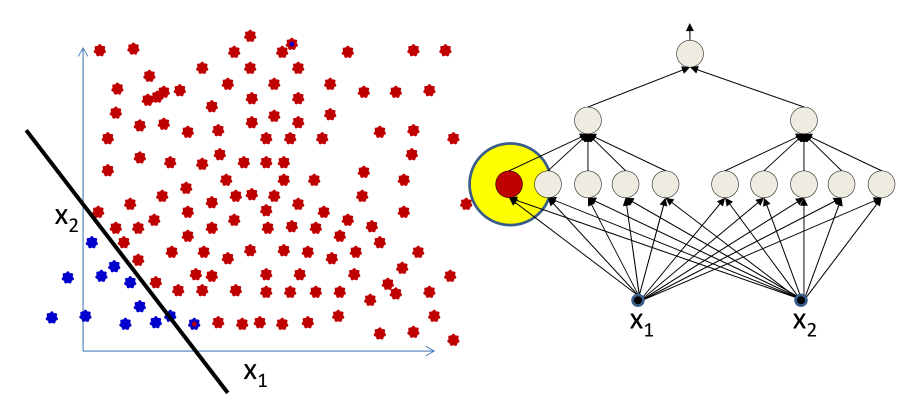


- Consider a single linear classifier that must be learned from the training data
  - Can it be learned from this data?

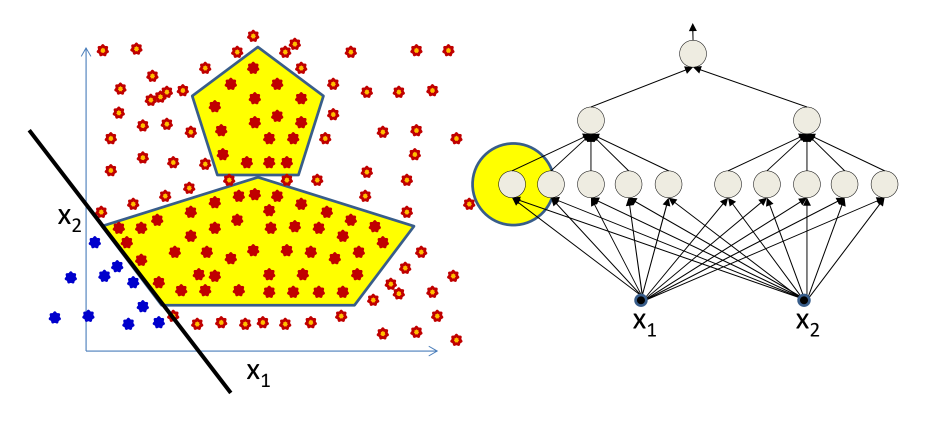
## Poll 2

#### Poll 2

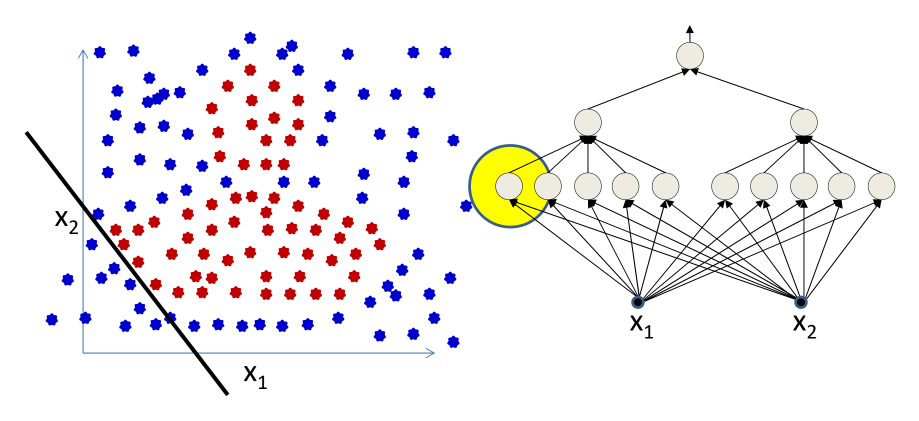
- For the double-pentagon problem, given the data shown on slide 60 and given that all but the one neuron highlighted in yellow are already correctly learned, can we use the perceptron learning algorithm to learn the one remaining neuron?
  - Yes
  - No
- What problems do you see in using the perceptron rule to learn the remaining perceptron?
  - Perceptron learning will require linearly separable classes to learn the model that classifies the data perfectly, but the data are not linearly separable
  - Perceptron learning will require relabelling the data to make them linearly separable with the correct decision boundary



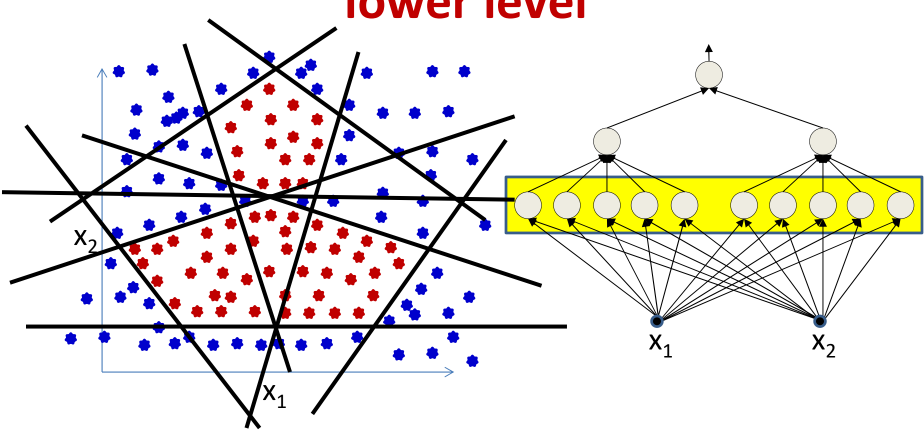
- Consider a single linear classifier that must be learned from the training data
  - Can it be learned from this data?
  - The individual classifier actually requires the kind of labelling shown here
    - Which is not given!!



- The lower-level neurons are linear classifiers
  - They require linearly separated labels to be learned
  - The actually provided labels are not linearly separated
  - Challenge: Must also learn the labels for the lowest units! 67



- For a single line:
  - Try out every possible way of relabeling the blue dots such that we can learn a line that keeps all the red dots on one side!



- This must be done for each of the lines (perceptrons)
- Such that, when all of them are combined by the higherlevel perceptrons we get the desired pattern
  - Basically an exponential search over inputs

Individual neurons represent one of the lines that compose the figure (linear classifiers)

 $X_2$   $X_1$ 

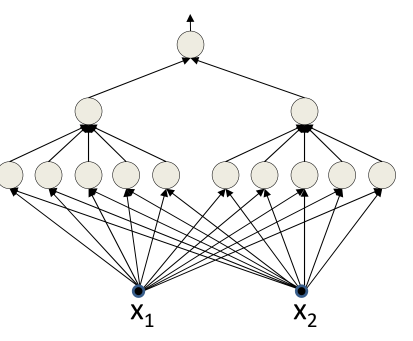
This must be done for every neuron

Getting any of them wrong will result in incorrect output!

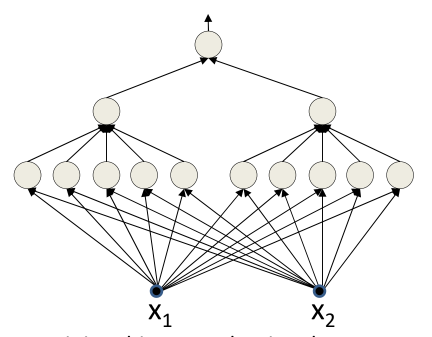
Must know the desired output of every neuron for *every* training instance, in order to learn this neuron

The outputs should be such that the neuron individually has a linearly separable task

The linear separators must combine to form the desired boundary



#### Learning a *multilayer* perceptron



Training data only specifies input and output of network

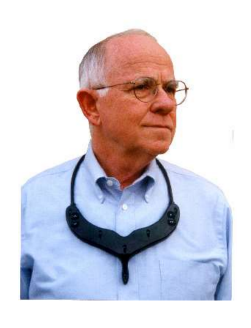
Intermediate outputs (outputs of individual neurons) are not specified

- Training this network using the perceptron rule is a combinatorial optimization problem
- We don't know the outputs of the individual intermediate neurons in the network for any training input
- Must also determine the correct output for each neuron for every training instance
- At least exponential (in inputs) time complexity!!!!!

# **Greedy algorithms: Adaline and Madaline**

- Perceptron learning rules cannot directly be used to learn an MLP
  - Exponential complexity of assigning intermediate labels
    - Even worse when classes are not actually separable
- Can we use a greedy algorithm instead?
  - Adaline / Madaline
  - On slides, will skip in class (check the quiz)

## A little bit of History: Widrow



#### **Bernie Widrow**

- Scientist, Professor, Entrepreneur
- Inventor of most useful things in signal processing and machine learning!

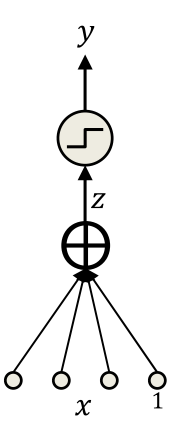
- First known attempt at an analytical solution to training the perceptron and the MLP
- Now famous as the LMS algorithm
  - Used everywhere
  - Also known as the "delta rule"

### **History: ADALINE**

$$z = \sum_{t} w_i x_i$$
 Using 1-extended vector notation to account for bias

$$y = \begin{cases} 0, & z < 0 \\ 1, & z \ge 0 \end{cases}$$

- Adaptive *linear* element (Hopf and Widrow, 1960)
- Actually just a regular perceptron
  - Weighted sum on inputs and bias passed through a thresholding function
- ADALINE differs in the learning rule



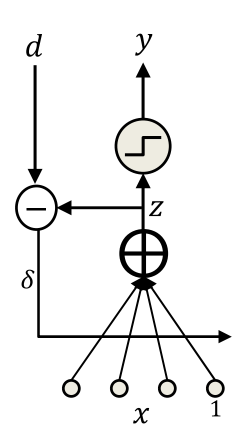
### **History: Learning in ADALINE**

$$z = \sum_{t} w_i x_i$$

$$out = \begin{cases} 0, & z < 0 \\ 1, & z \ge 0 \end{cases}$$

- During learning, minimize the squared error assuming z to be real output
- The desired output is still binary!

$$Err(x) = \frac{1}{2}(d-z)^2$$
 Error for a single input 
$$\frac{dErr(x)}{dw_i} = -(d-z)x_i$$



### **History: Learning in ADALINE**

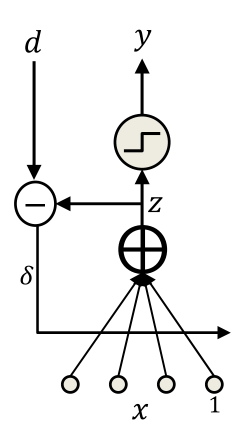
$$z = \sum_{t} w_i x_i$$

$$Err(x) = \frac{1}{2}(d-z)^2$$
 Error for a single input

$$\frac{dErr(x)}{dw_i} = -(d-z)x_i$$

If we just have a single training input,
 the gradient descent update rule is

$$w_i = w_i + \eta(d - z)x_i$$

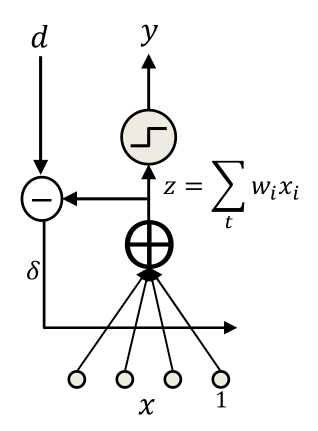


## The ADALINE learning rule

- Online learning rule
- After each input x, that has target (binary) output d, compute and update:

$$\delta = d - z$$
$$w_i = w_i + \eta \delta x_i$$

- This is the famous delta rule
  - Also called the LMS update rule



### The Delta Rule

- In fact both the Perceptron and ADALINE use variants of the delta rule!
  - Perceptron: Output used in delta rule is y

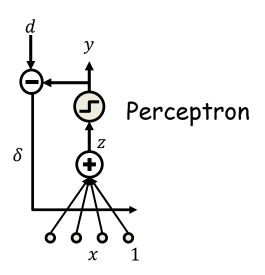
$$\delta = d - y$$

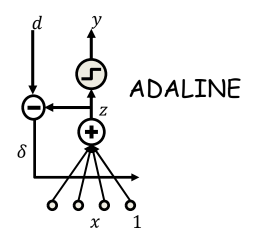
ADALINE: Output used to estimate weights is z

$$\delta = d - z$$

For both

$$w_i = w_i + \eta \delta x_i$$





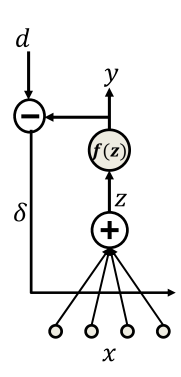
### Aside: Generalized delta rule

 For any differentiable activation function the following update rule is used

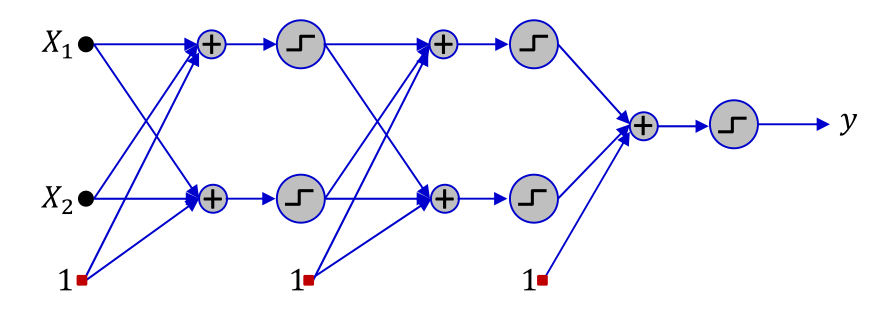
$$\delta = d - y$$

$$w_i = w_i + \eta \delta f'(z) x_i$$

- This is the famous Widrow-Hoff update rule
  - Lookahead: Note that this is *exactly* backpropagation in multilayer nets if we let f(z) represent the entire network between z and y
- It is possibly the most-used update rule in machine learning and signal processing
  - Variants of it appear in almost every problem

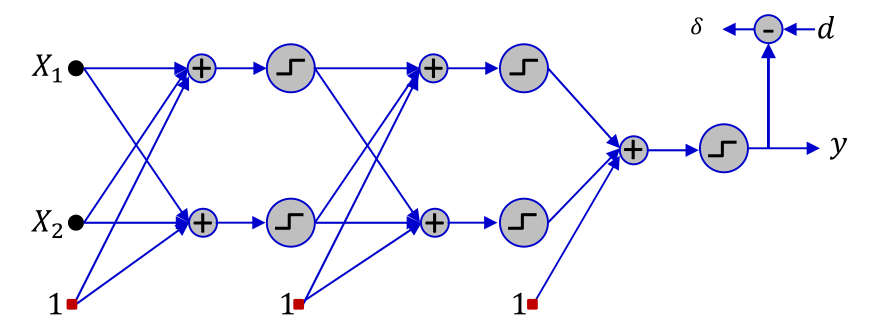


### Multilayer perceptron: MADALINE

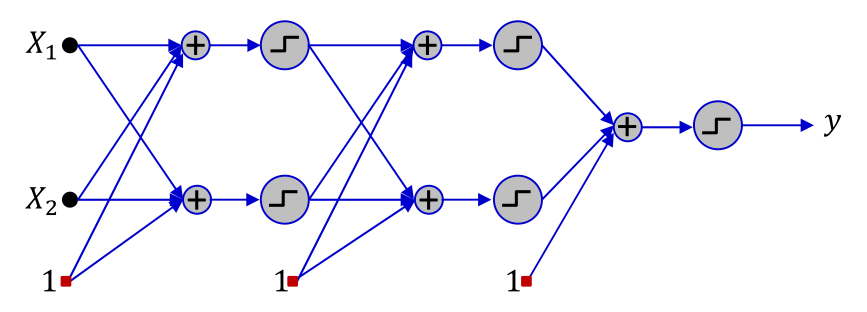


### Multiple Adaline

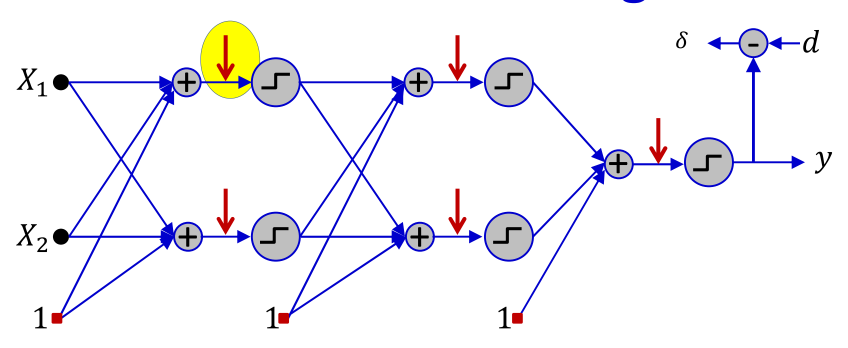
- A multilayer perceptron with threshold activations
- The MADALINE



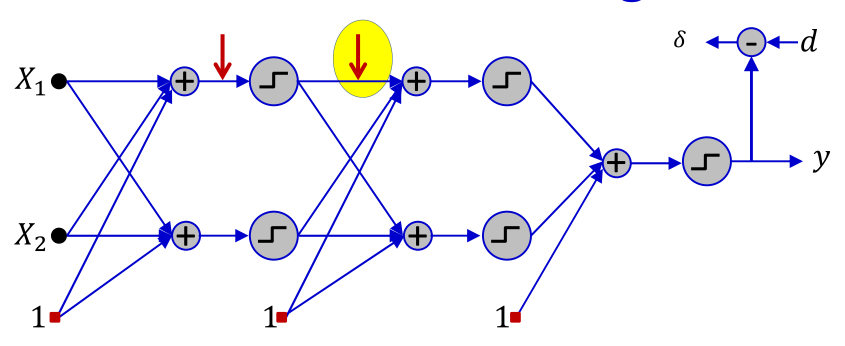
- Update only on error
  - $-\delta \neq 0$
  - On inputs for which output and target values differ



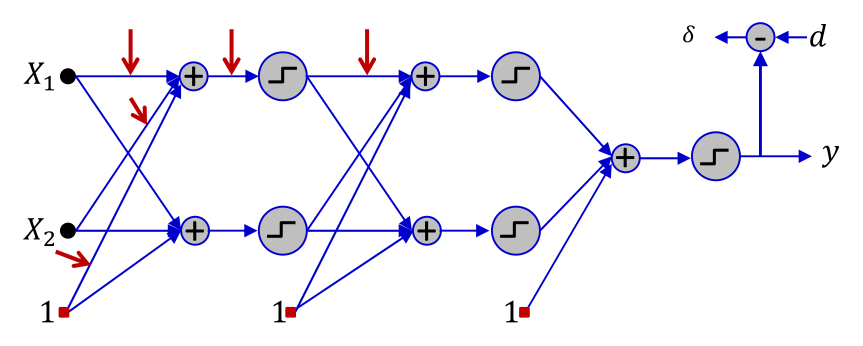
- While stopping criterion not met do:
  - Classify an input



- While stopping criterion not met do:
  - Classify an input
  - If error, find the z that is closest to 0



- While stopping criterion not met do:
  - Classify an input
  - If error, find the z that is closest to 0
  - Flip the output of corresponding unit and compute new output



- While stopping criterion not met do:
  - Classify an input
  - If error, find the z that is closest to 0
  - Flip the output of corresponding unit and compute new output
  - If error reduces:
    - Set the desired output of the unit to the flipped value
    - Apply ADALINE rule to update weights of the unit

### **MADALINE**

- Greedy algorithm, effective for small networks
- Not very useful for large nets
  - Too expensive
  - Too greedy

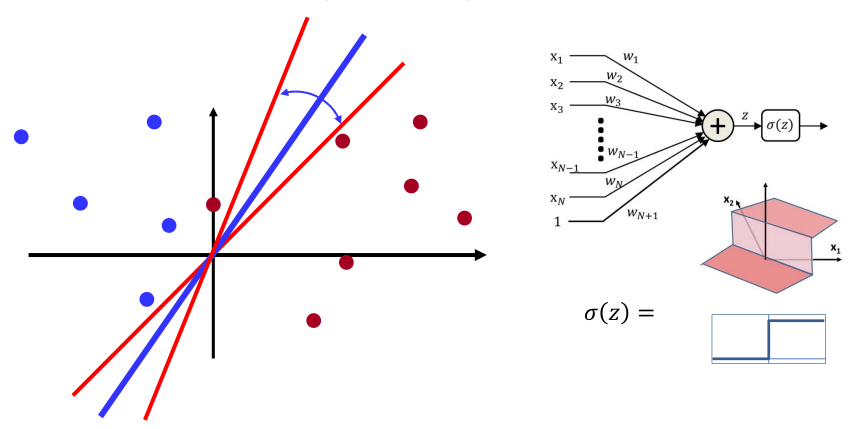
### Story so far

- "Learning" a network = learning the weights and biases to compute a target function
  - Will require a network with sufficient "capacity"
- In practice, we learn networks by "fitting" them to match the input-output relation of "training" instances drawn from the target function
- A linear decision boundary can be learned by a single perceptron (with a threshold-function activation) in linear time if classes are linearly separable
- Non-linear decision boundaries require networks of perceptrons
- Training an MLP with threshold-function activation perceptrons will require knowledge of the input-output relation for every training instance, for every perceptron in the network
  - These must be determined as part of training
  - For threshold activations, this is an NP-complexity combinatorial optimization problem

### History...

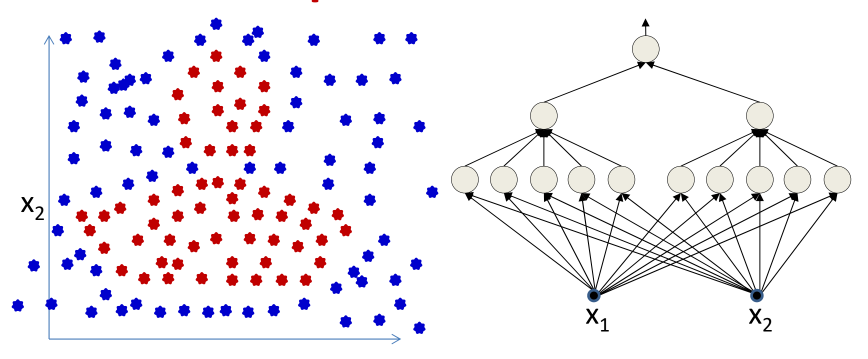
 The realization that training an entire MLP was a combinatorial optimization problem stalled development of neural networks for well over a decade!

### Why this problem?



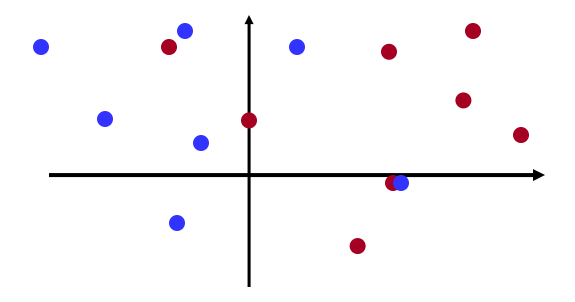
- The perceptron is a flat function with zero derivative everywhere, except at 0 where it is non-differentiable
  - You can vary the weights a *lot* without changing the error
  - There is no indication of which direction to change the weights to reduce error

# This only compounds on larger problems



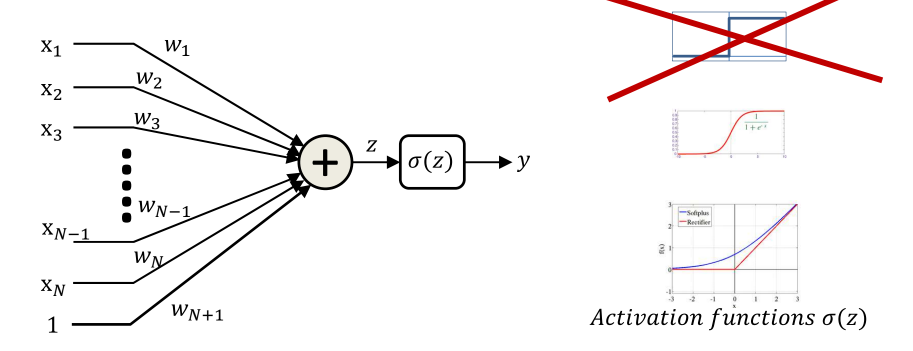
- Individual neurons' weights can change significantly without changing overall error
- The simple MLP is a flat, non-differentiable function
  - Actually a function with 0 derivative nearly everywhere, and no derivatives at the boundaries

## A second problem: What we *actually* model



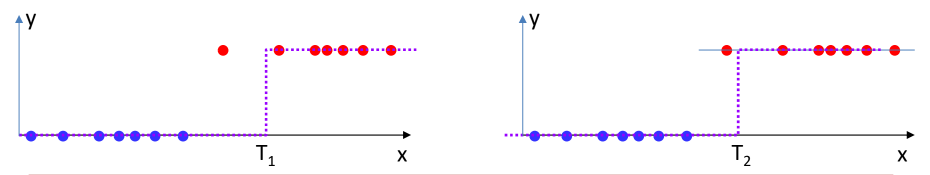
- Real-life data are rarely clean
  - Not linearly separable
  - Rosenblatt's perceptron wouldn't work in the first place

### Solution

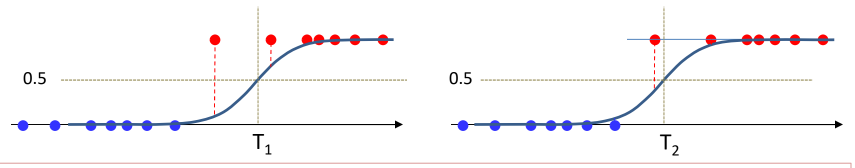


- Lets make the neuron differentiable, with non-zero derivatives over much of the input space
  - Small changes in weight can result in non-negligible changes in output
  - This enables us to estimate the parameters using gradient descent techniques..

### Differentiable activation function



- Threshold activation: shifting the threshold from T<sub>1</sub> to T<sub>2</sub> does not change classification error
  - Does not indicate if moving the threshold left was good or not



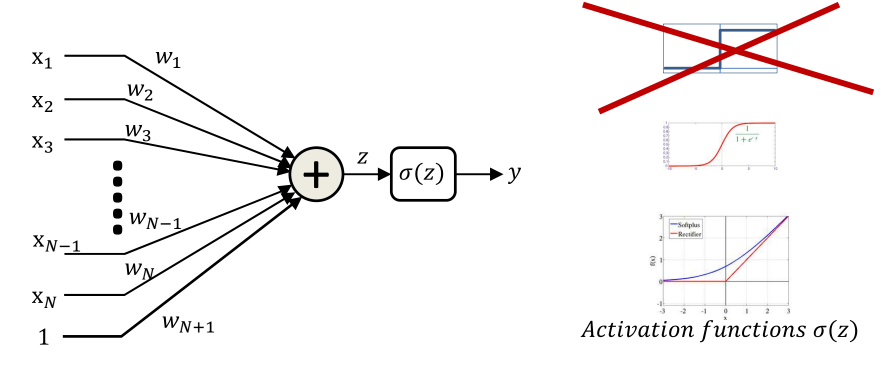
- Smooth, continuously varying activation: Classification based on whether the output is greater than 0.5 or less
  - Can now quantify how much the output differs from the desired target value (0 or 1)
  - Moving the function left or right changes this quantity, even if the classification error itself doesn't change

## Poll 3

### Poll 3

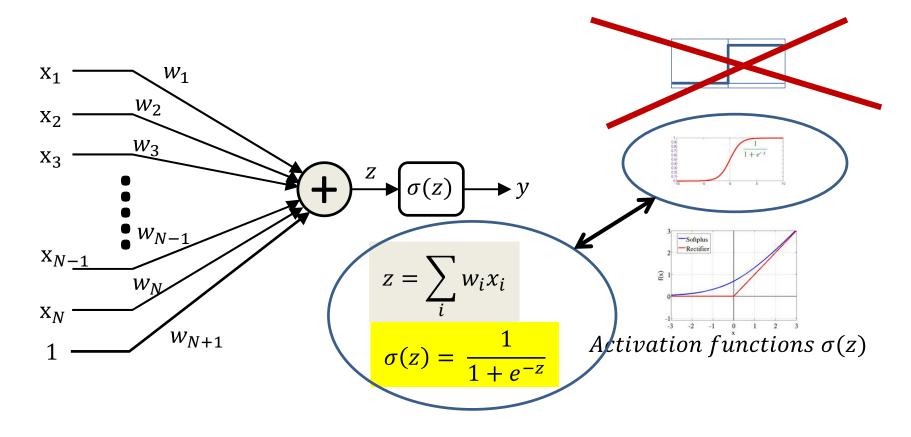
- Which of the following are true of the threshold activation
  - Increasing (or decreasing) the threshold will not change the overall classification error unless the threshold moves past a misclassified training sample
  - We cannot know if a change (increase of decrease) of the threshold moves it in the correct direction that will result in a net decrease in classification error
  - The derivative of the classification error with respect to the threshold gives us an indication of whether to increase or decrease the threshold
- Which of the following are true of the continuous activation (sigmoid)
  - Shifting the function left or right will not change the overall classification error unless the crossover point (where the function crosses 0.5) moves past a misclassified training sample
  - Shifting the function will change the total distance of the value of the function from its target value at the training instances
  - The derivative of the total distance with respect to the shift of the function gives us an indication of which direction to shift the function to improve classification error

### **Continuous Activations**



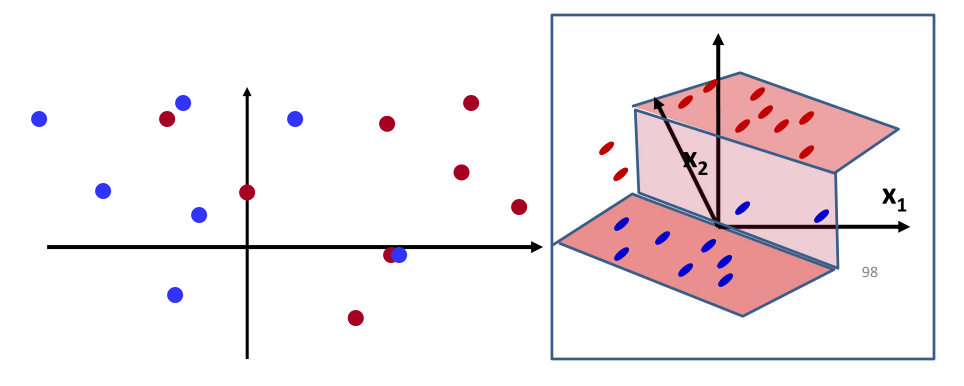
- Replace the threshold activation with continuous graded activations
  - E.g. RELU, softplus, sigmoid etc.
- The activations are differentiable almost everywhere
  - Have derivatives almost everywhere
  - And have "subderivatives" at non-differentiable corners
    - Bounds on the derivative that can substitute for derivatives in our setting
    - More on these later

### The sigmoid activation is special



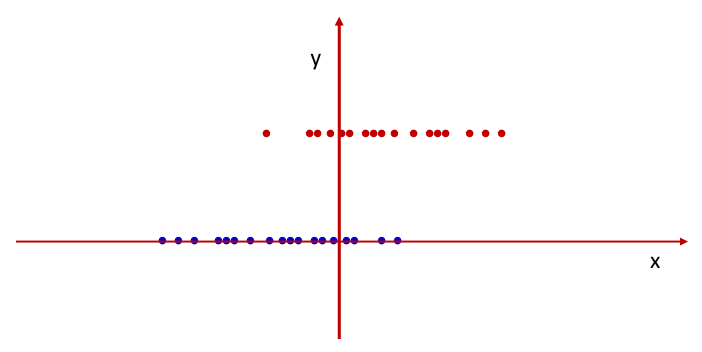
- This particular one has a nice interpretation
- It can be interpreted as P(y = 1|x)

### Non-linearly separable data

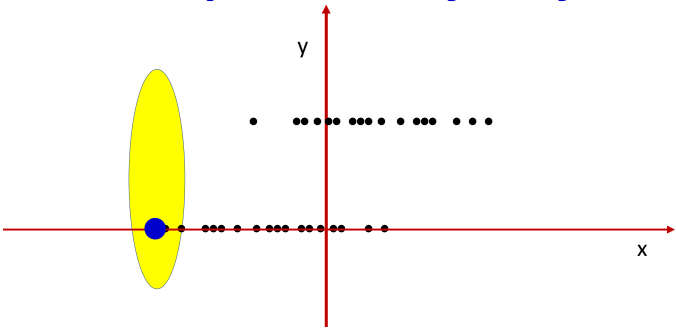


- Two-dimensional example
  - Blue dots (on the floor) on the "red" side
  - Red dots (suspended at Y=1) on the "blue" side
  - No line will cleanly separate the two colors

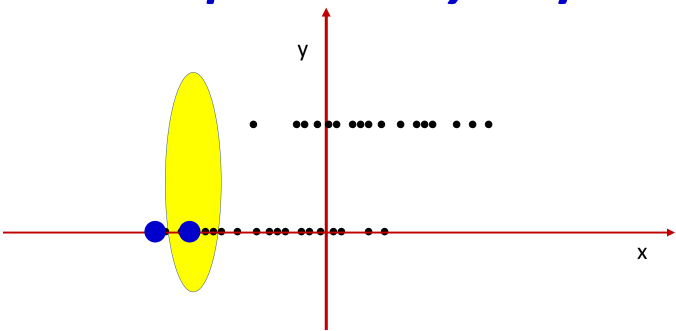
### Non-linearly separable data: 1-D example



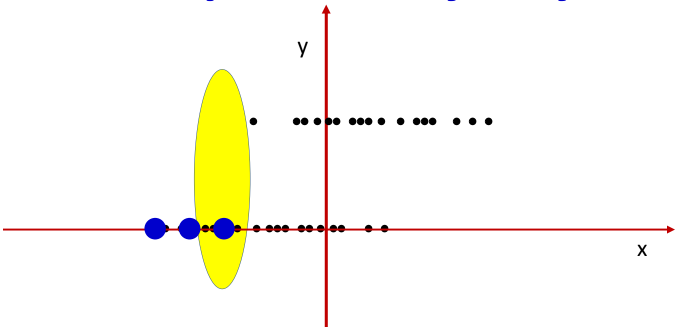
- One-dimensional example for visualization
  - All (red) dots at Y=1 represent instances of class Y=1
  - All (blue) dots at Y=0 are from class Y=0
  - The data are not linearly separable
    - In this 1-D example, a linear separator is a threshold
    - No threshold will cleanly separate red and blue dots



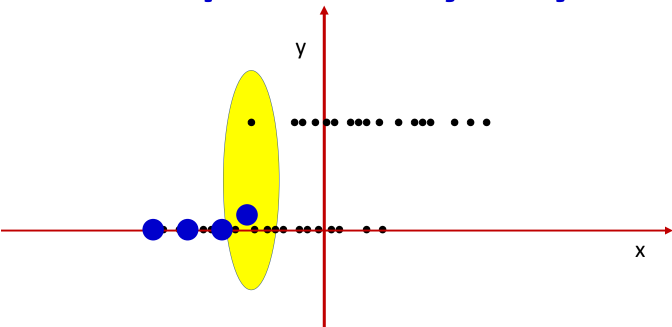
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of Y=1 at that point



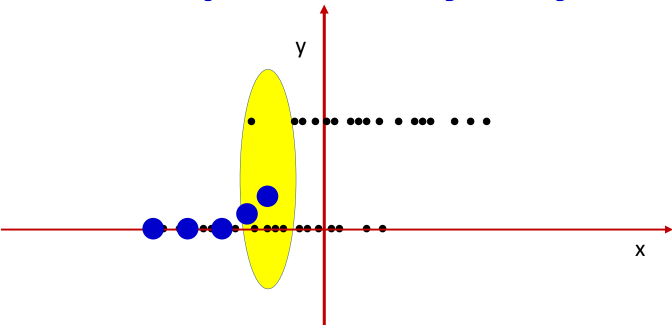
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



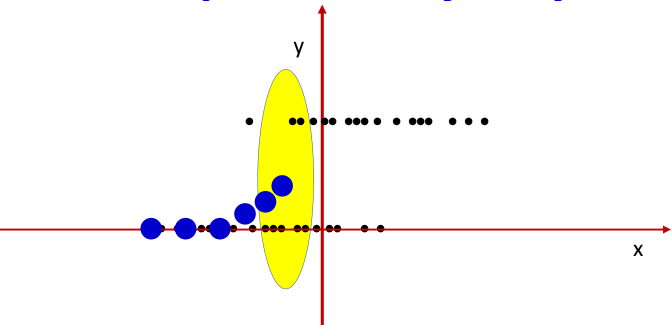
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



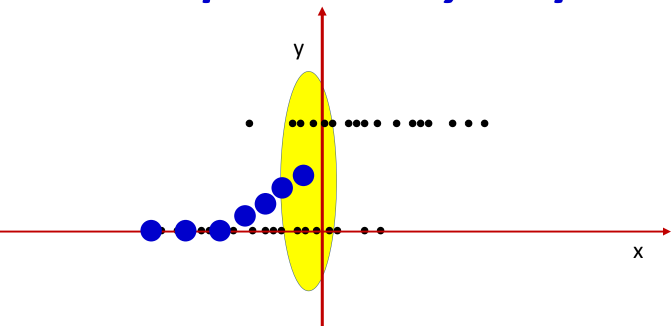
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



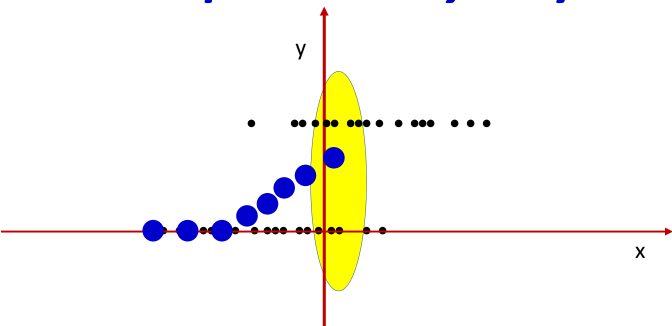
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



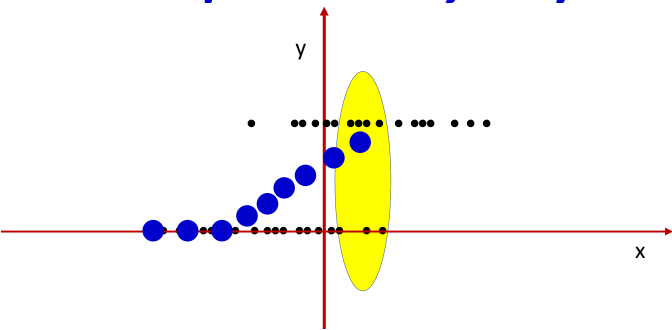
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



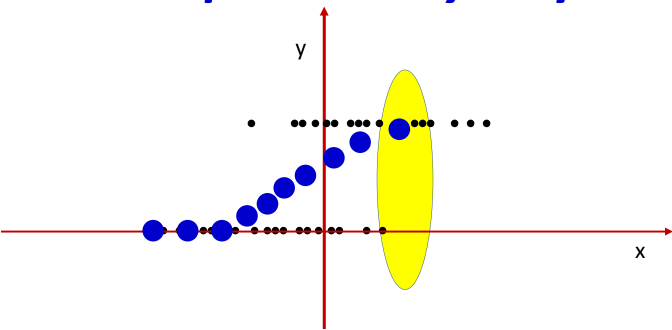
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



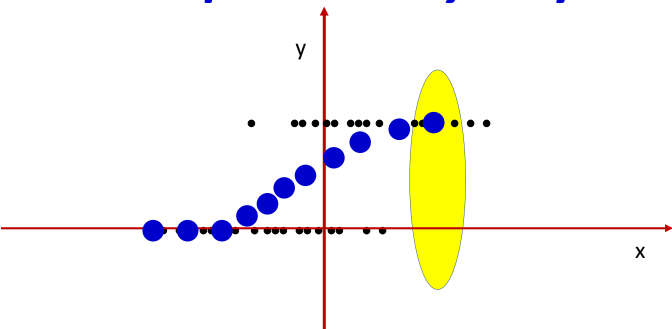
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



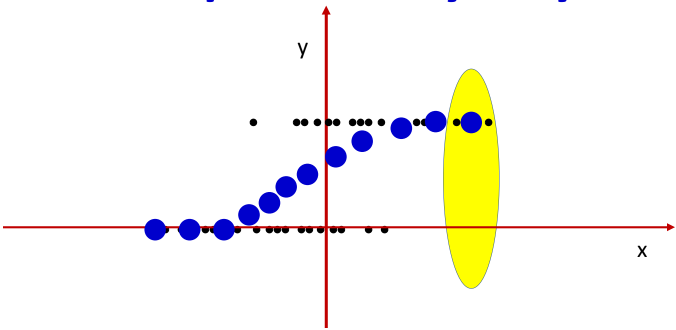
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



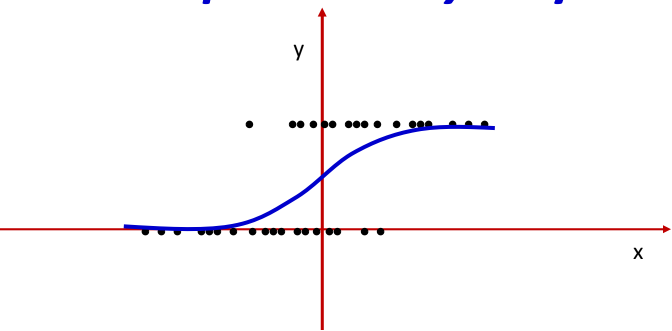
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point

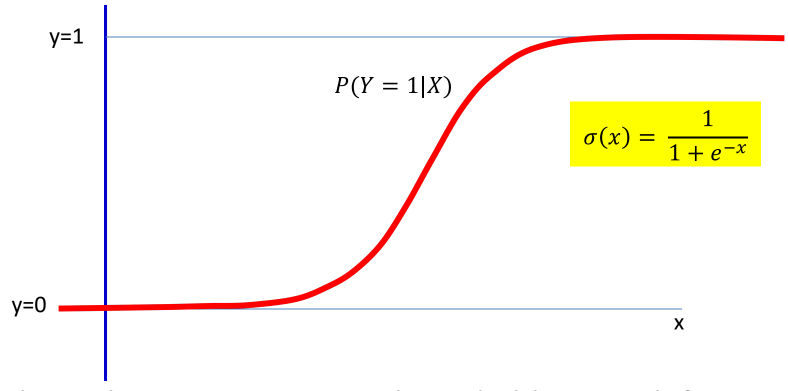


- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the probability of 1 at that point



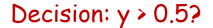
- Consider this differently: at each point look at a small window around that point
- Plot the average value within the window
  - This is an approximation of the *probability* of 1 at that point

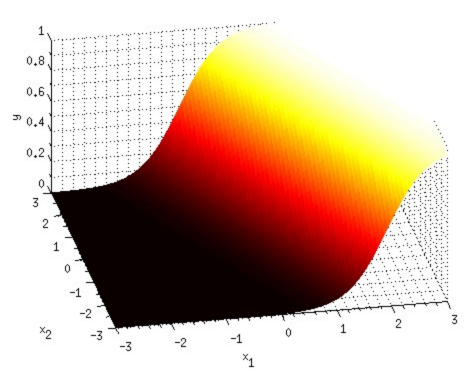
## The logistic regression model

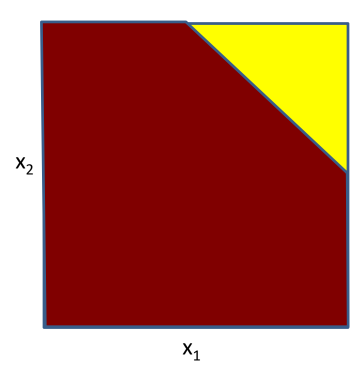


- Class 1 becomes increasingly probable going left to right
  - Very typical in many problems

## Logistic regression







When X is a 2-D variable

$$P(Y = 1|X) = \frac{1}{1 + \exp(-\sum_{i} w_{i} x_{i} - b)}$$

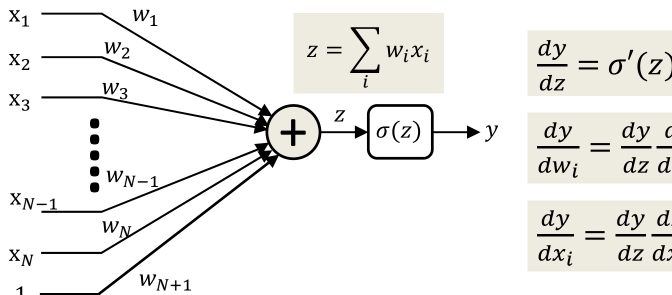
- This the perceptron with a sigmoid activation
  - It actually computes the probability that the input belongs to class 1

## Perceptrons and probabilities

 We will return to the fact that perceptrons with sigmoidal activations actually model class probabilities in a later lecture

But for now moving on..

## Perceptrons with differentiable activation functions



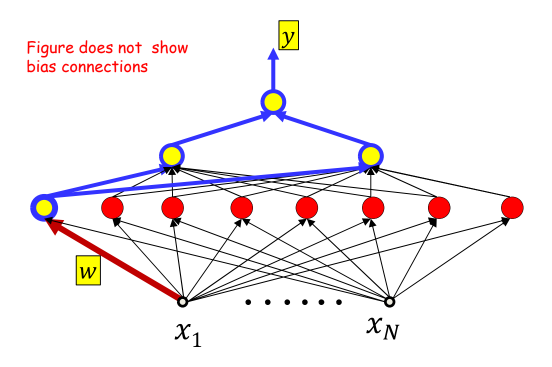
$$\frac{dy}{dz} = \sigma'(z)$$

$$\frac{dy}{dw_i} = \frac{dy}{dz} \frac{dz}{dw_i} = \sigma'(z)x_i$$

$$\frac{dy}{dx_i} = \frac{dy}{dz} \frac{dz}{dx_i} = \sigma'(z) w_i$$

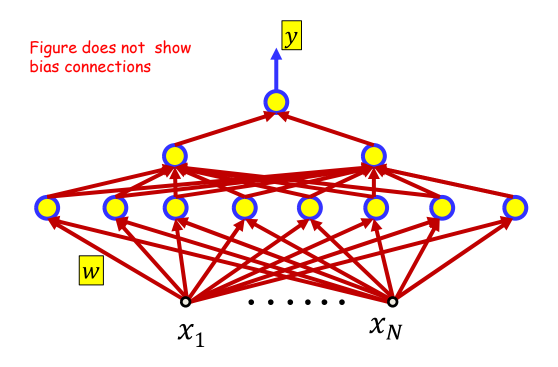
- $\sigma(z)$  is a differentiable function of z
  - $-\frac{d\sigma(z)}{dz}$  is well-defined and finite for all z
- Using the chain rule, y is a differentiable function of both inputs  $x_i$  and weights  $w_i$
- This means that we can compute the change in the output for small changes in either the input or the weights

#### Overall network is differentiable



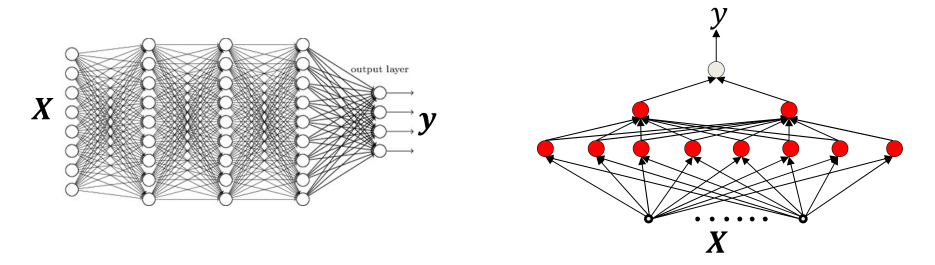
- Every individual perceptron is differentiable w.r.t its inputs and its weights (including "bias" weight)
  - Small changes in the parameters result in measurable changes in output
- Using the chain rule can compute how small perturbations of a parameter change the output of the network
  - The network output is differentiable with respect to the parameter

#### Overall function is differentiable



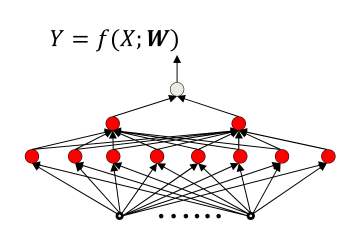
- By extension, the overall function is differentiable w.r.t every parameter in the network
  - We can compute how small changes in the parameters change the output
    - For non-threshold activations the derivative are finite and generally non-zero
- We will derive the actual derivatives using the chain rule later

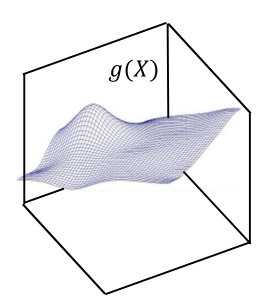
## Overall setting for "Learning" the MLP



- Given a training set of input-output pairs  $(X_1, d_1), (X_2, d_2), \dots, (X_N, d_N) \dots$ 
  - d is the desired output of the network in response to X
  - X and d may both be vectors
- ...we must find the network parameters such that the network produces the desired output for each training input
  - Or a close approximation of it
  - The architecture of the network must be specified by us

## **Recap: Learning the function**



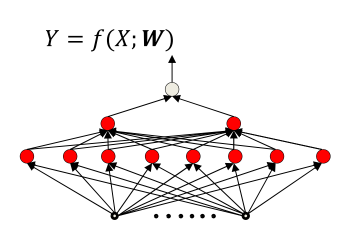


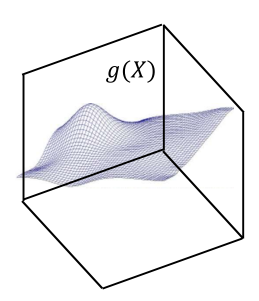
• When f(X; W) has the capacity to exactly represent g(X)

$$\widehat{\boldsymbol{W}} = \underset{W}{\operatorname{argmin}} \int_{X} div(f(X; W), g(X))dX$$

• div() is a divergence function that goes to zero when f(X; W) = g(X)

## Minimizing expected divergence

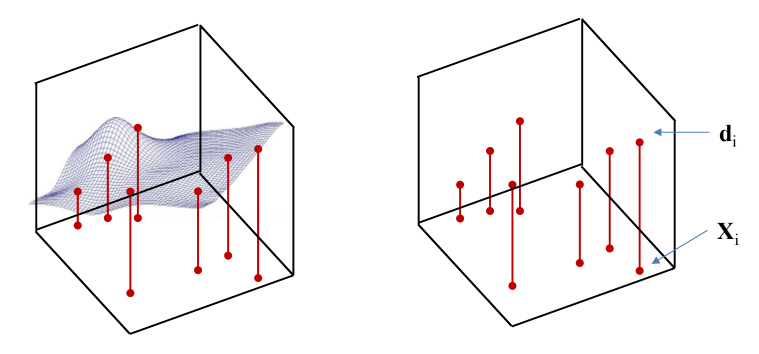




• More generally, assuming X is a random variable

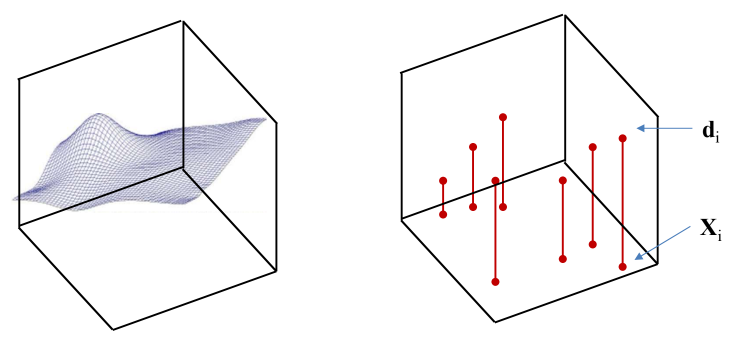
$$\widehat{W} = \underset{W}{\operatorname{argmin}} \int_{X} div(f(X; W), g(X))P(X)dX$$
$$= \underset{W}{\operatorname{argmin}} E[div(f(X; W), g(X))]$$

## Recap: Sampling the function



- We don't have g(X) so sample g(X)
  - Obtain input-output pairs for a number of samples of input  $X_i$
  - Good sampling: the samples of X will be drawn from P(X)
- Estimate function from the samples

## The *Empirical* risk



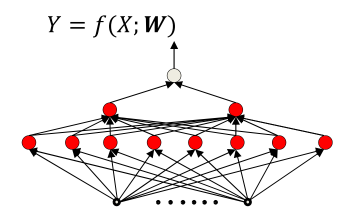
The expected divergence (or risk) is the average divergence over the entire input space

$$E[div(f(X;W),g(X))] = \int_X div(f(X;W),g(X))P(X)dX$$

The empirical estimate of the expected risk is the average divergence over the samples

$$E[div(f(X;W),g(X))] \approx \frac{1}{N} \sum_{i=1}^{N} div(f(X_i;W),d_i)$$

## **Empirical Risk Minimization**



- Given a training set of input-output pairs  $(X_1, d_1), (X_2, d_2), ..., (X_N, d_N)$ 
  - Quantification of error on the i<sup>th</sup> instance:  $div(f(X_i; W), d_i)$
  - Empirical average divergence (Empirical Risk) on all training data:

$$Loss(W) = \frac{1}{N} \sum_{i} div(f(X_i; W), d_i)$$

• Estimate the parameters to minimize the empirical estimate of expected divergence (empiricial risk)

$$\widehat{\boldsymbol{W}} = \underset{W}{\operatorname{argmin}} Loss(W)$$

I.e. minimize the *empirical risk* over the drawn samples

## **Empirical Risk Minimization**

$$Y = f(X; \boldsymbol{W})$$

Note: Its really a measure of error, but using standard terminology, we will call it a "Loss"

Note 2: The empirical risk Loss(W) is only an empirical approximation to the true risk E[div(f(X;W),g(X))] which is our actual minimization objective

Note 3: For a given training set the loss is only a function of W

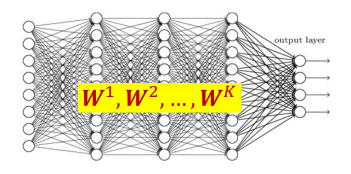
$$Loss(W) = \frac{1}{N} \sum_{i} div(f(X_i; W), d_i)$$

Estimate the parameters to minimize the empirical estimate of expected error

$$\widehat{\boldsymbol{W}} = \underset{W}{\operatorname{argmin}} Loss(W)$$

I.e. minimize the *empirical error* over the drawn samples

#### **ERM** for neural networks



#### **Actual output of network:**

$$Y_{i} = net(X_{i}; \{w_{i,j}^{k} \forall i, j, k\})$$
$$= net(X_{i}; W^{1}, W^{2}, ..., W^{K})$$

Desired output of network:  $d_i$ 

Error on i-th training input:  $Div(Y_i, d_i; W^1, W^2, ..., W^K)$ 

#### Average training error(loss):

$$Loss(\mathbf{W}^{1}, \mathbf{W}^{2}, ..., \mathbf{W}^{K}) = \frac{1}{N} \sum_{i=1}^{N} \mathbf{Div}(\mathbf{Y}_{i}, \mathbf{d}_{i}; \mathbf{W}^{1}, \mathbf{W}^{2}, ..., \mathbf{W}^{K})$$

- What is the exact form of Div()? More on this later
- Optimize network parameters to minimize the total error over all training inputs

#### **Problem Statement**

- Given a training set of input-output pairs  $(X_1, d_1), (X_2, d_2), ..., (X_N, d_N)$
- Minimize the following function

$$Loss(W) = \frac{1}{N} \sum_{i} div(f(X_i; W), d_i)$$

w.r.t W

- This is problem of function minimization
  - An instance of optimization

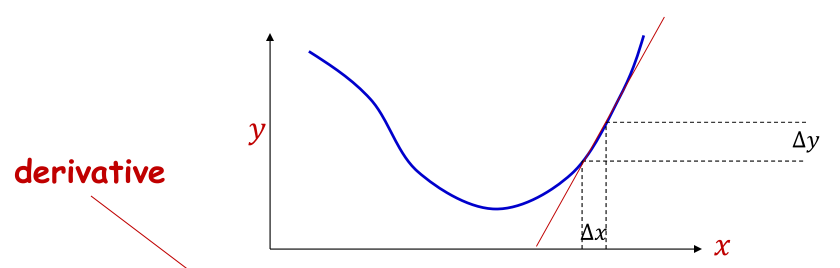
## Story so far

- We learn networks by "fitting" them to training instances drawn from a target function
- Learning networks of threshold-activation perceptrons requires solving a hard combinatorial-optimization problem
  - Because we cannot compute the influence of small changes to the parameters on the overall error
- Instead we use continuous activation functions with non-zero derivatives to enables us to estimate network parameters
  - This makes the output of the network differentiable w.r.t every parameter in the network
  - The *logistic* activation perceptron actually computes the *a posteriori* probability of the output given the input
- We define differentiable divergence between the output of the network and the desired output for the training instances
  - And a total error, which is the average divergence over all training instances
- We optimize network parameters to minimize this error
  - Empirical risk minimization
- This is an instance of function minimization.

#### A CRASH COURSE ON FUNCTION OPTIMIZATION

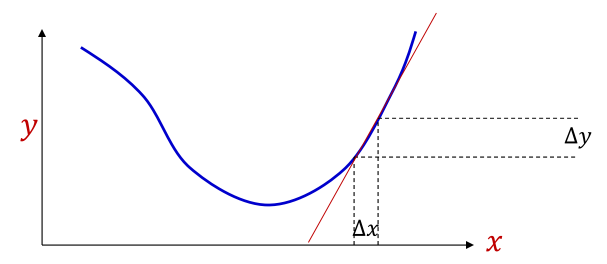
- With an initial discussion of derivatives

### A brief note on derivatives...



- A derivative of a function at any point tells us how much a minute increment to the *argument* of the function will increment the *value* of the function
  - For any y=f(x), expressed as a multiplier  $\alpha$  to a tiny increment  $\Delta x$  to obtain the increments  $\Delta y$  to the output  $\Delta y = \alpha \Delta x$
  - Based on the fact that at a fine enough resolution, any smooth, continuous function is locally linear at any point 130

## Scalar function of scalar argument



When x and y are scalar

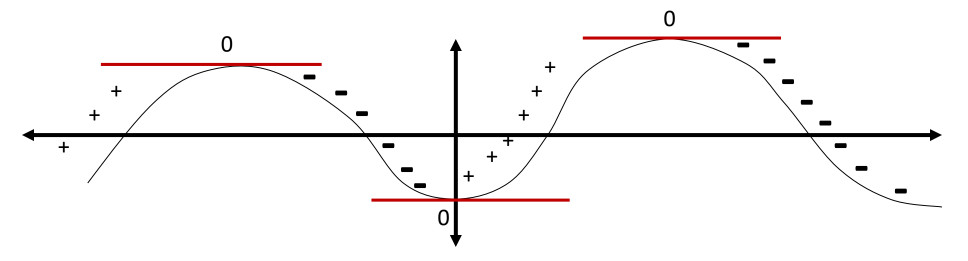
$$y = f(x)$$

Derivative:

$$\Delta y = \alpha \Delta x$$

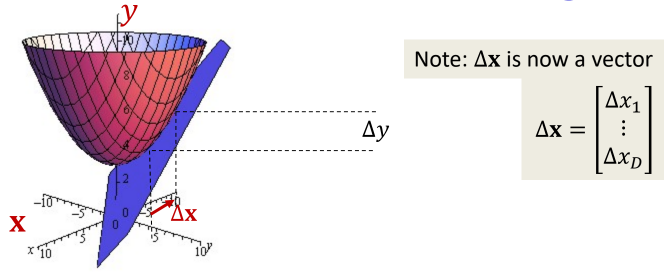
- Often represented as  $\frac{dy}{dx}$
- Or alternately (and more reasonably) as f'(x)

## Scalar function of scalar argument



- Derivative f'(x) is the *rate of change* of the function at x
  - How fast it increases with increasing x
  - The magnitude of f'(x) gives you the steepness of the curve at x
    - Larger  $|f'(x)| \rightarrow$  the function is increasing or decreasing more rapidly
- It will be positive where a small increase in x results in an *increase* of f(x)
  - Regions of positive slope
- It will be negative where a small increase in x results in a decrease of f(x)
  - Regions of negative slope
- It will be 0 where the function is locally flat (neither increasing nor decreasing)

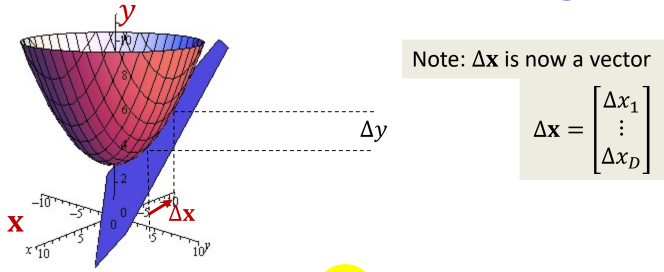
## Multivariate scalar function: Scalar function of *vector* argument



$$\Delta y = \alpha \Delta \mathbf{x}$$

- Giving us that  $\alpha$  is a row vector:  $\alpha = \begin{bmatrix} \alpha_1 & \cdots & \alpha_D \end{bmatrix}$   $\Delta y = \alpha_1 \Delta x_1 + \alpha_2 \Delta x_2 + \cdots + \alpha_D \Delta x_D$
- The partial derivative  $\alpha_i$  gives us how y increments when only  $x_i$  is incremented
- Often represented as  $\frac{\partial y}{\partial x_i}$  $\Delta y = \frac{\partial y}{\partial x_1} \Delta x_1 + \frac{\partial y}{\partial x_2} \Delta x_2 + \dots + \frac{\partial y}{\partial x_D} \Delta x_D$

# Multivariate scalar function: Scalar function of *vector* argument



$$\Delta y = \nabla_{\mathbf{x}} y \Delta \mathbf{x}$$

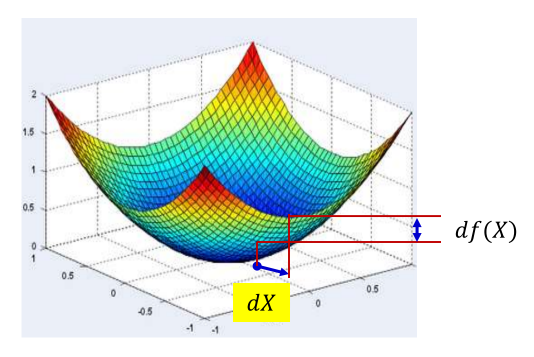
Where

$$\nabla_{\mathbf{x}} y = \begin{bmatrix} \frac{\partial y}{\partial x_1} & \cdots & \frac{\partial y}{\partial x_D} \end{bmatrix}$$

We will be using this symbol for vector and matrix derivatives

 You may be more familiar with the term "gradient" which is actually defined as the transpose of the derivative

#### **Gradient** of a scalar function of a vector



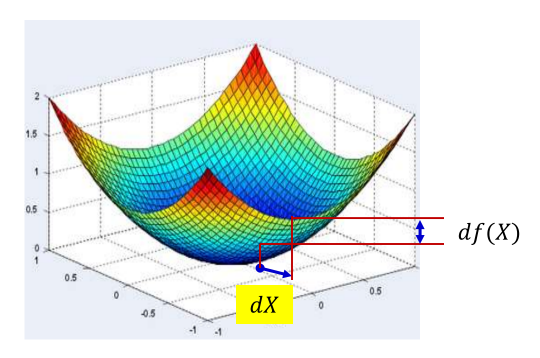
• The derivative  $\nabla_X f(X)$  of a scalar function f(X) of a multi-variate input X is a multiplicative factor that gives us the change in f(X) for tiny variations in X

$$df(X) = \nabla_X f(X) dX$$

$$- \nabla_X f(X) = \begin{bmatrix} \frac{\partial f(X)}{\partial x_1} & \frac{\partial f(X)}{\partial x_2} & \cdots & \frac{\partial f(X)}{\partial x_n} \end{bmatrix}$$

- The *gradient* is the transpose of the derivative  $\nabla_X f(X)^T$ 
  - A column vector of the same dimensionality as X

#### **Gradient** of a scalar function of a vector



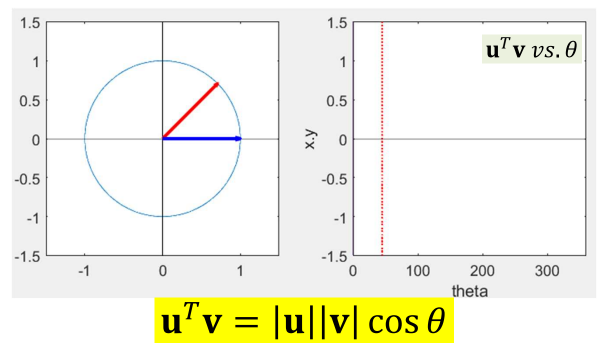
• The derivative  $\nabla_X f(X)$  of a scalar function f(X) of a multi-variate input X is a multiplicative factor that gives us the change in f(X) for tiny variations in X

$$df(X) = \nabla_X f(X) dX$$

$$- \nabla_X f(X) = \begin{bmatrix} \frac{\partial f(X)}{\partial x_1} & \frac{\partial f(X)}{\partial x_2} & \cdots & \frac{\partial f(X)}{\partial x_n} \end{bmatrix}$$

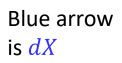
This is a vector inner product. To understand its behavior lets consider a well-known property of inner products

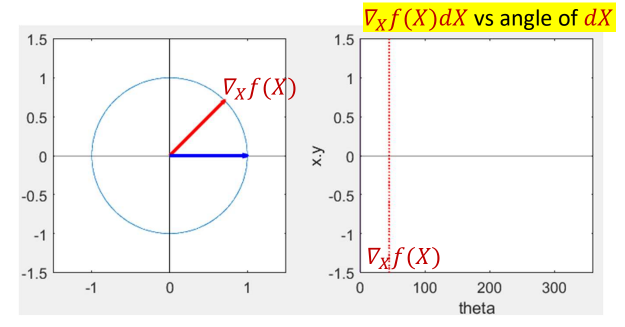
## A well-known vector property



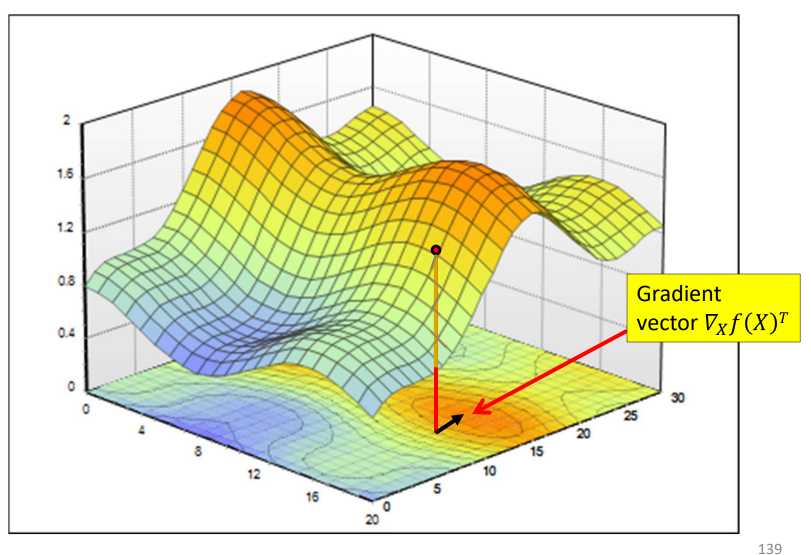
- The inner product between two vectors of fixed lengths is maximum when the two vectors are aligned
  - i.e. when  $\theta = 0$

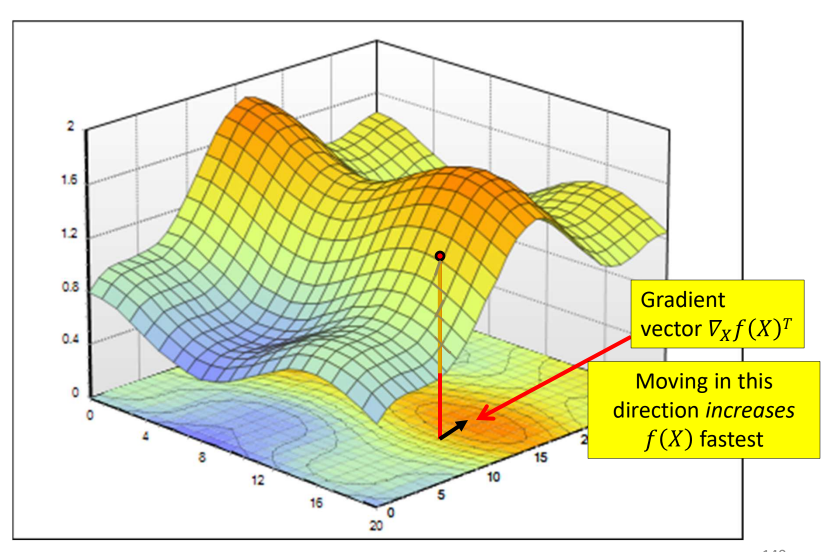
## **Properties of Gradient**

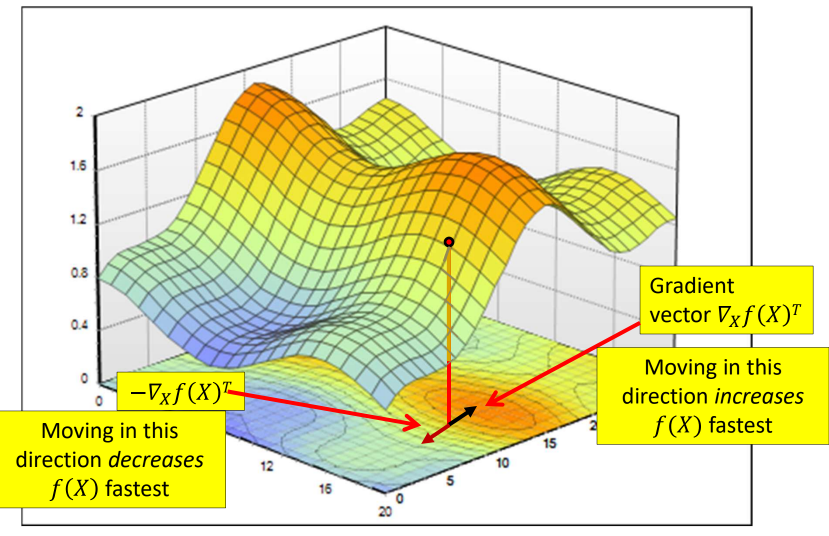


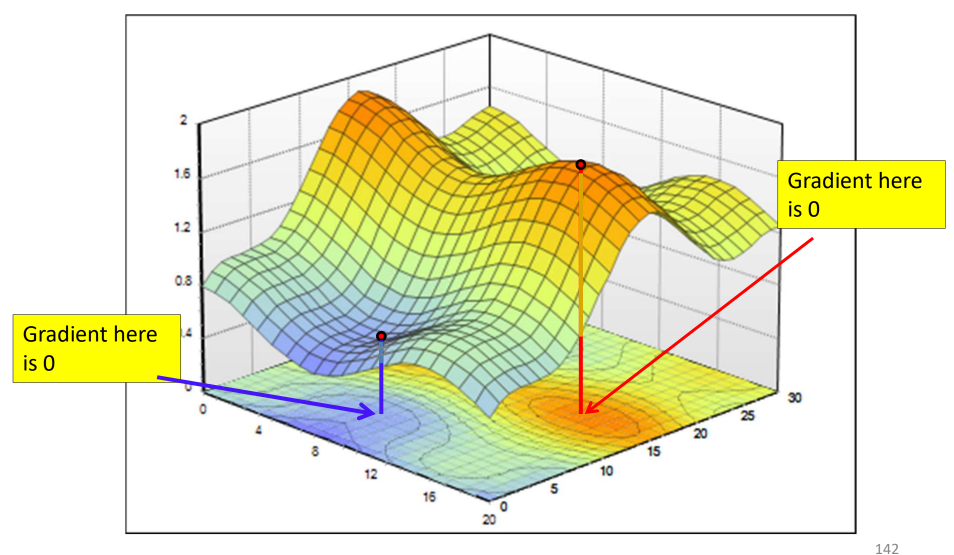


- $df(X) = \nabla_X f(X) dX$
- For an increment dX of any given length df(X) is max if dX is aligned with  $\nabla_X f(X)^T$ 
  - The function f(X) increases most rapidly if the input increment dX is exactly in the direction of  $\nabla_X f(X)^T$
- The gradient is the direction of fastest increase in f(X)

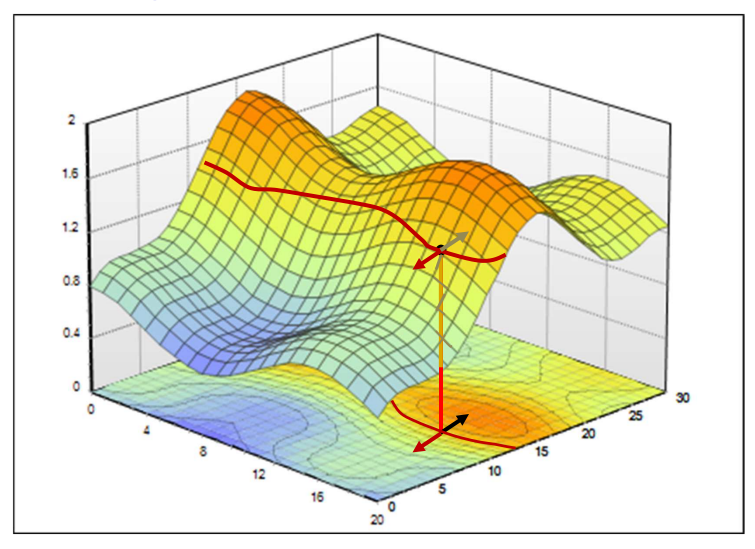








## **Properties of Gradient: 2**



• The gradient vector  $\nabla_X f(X)^T$  is perpendicular to the level curve

#### The Hessian

• The Hessian of a function  $f(x_1, x_2, ..., x_n)$  is given by the second derivative

$$\nabla_{x}^{2} f(x_{1},...,x_{n}) := \begin{bmatrix} \frac{\partial^{2} f}{\partial x_{1}^{2}} & \frac{\partial^{2} f}{\partial x_{1} \partial x_{2}} & \cdots & \frac{\partial^{2} f}{\partial x_{1} \partial x_{n}} \\ \frac{\partial^{2} f}{\partial x_{2} \partial x_{1}} & \frac{\partial^{2} f}{\partial x_{2}^{2}} & \cdots & \frac{\partial^{2} f}{\partial x_{2} \partial x_{n}} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial^{2} f}{\partial x_{n} \partial x_{1}} & \frac{\partial^{2} f}{\partial x_{n} \partial x_{2}} & \cdots & \frac{\partial^{2} f}{\partial x_{n}^{2}} \end{bmatrix}$$

## Next up

Continuing on function optimization

Gradient descent to train neural networks

A.K.A. Back propagation

## Poll 4

#### Poll 4

- Select all that are true about derivatives of a scalar function f(X) of multivariate inputs
  - At any location X, there may be many directions in which we can step, such that f(X) increases
  - The direction of the gradient is the direction in which the function increases fastest
  - The gradient is the derivative of f(X) w.r.t. X