Machine Learning

Topic: Gradient Descent

Supervised Machine Learning in one slide

- 1. Pick data X, labels Y, model $M(\theta)$ and loss function $L(X, Y; \theta)$
- 2. Initialize model parameters θ , somehow

3. Measure model performance with the loss function $L(X, Y; \theta)$

HOW?

4. Modify parameters θ somehow, hoping to improve $L(X, Y; \theta)$

5. Repeat 3 and 4 until you stop improving or run out of time

A common approach to picking the next parameters

HOW?

1. Measure how the the loss changes when we change the parameters θ slightly

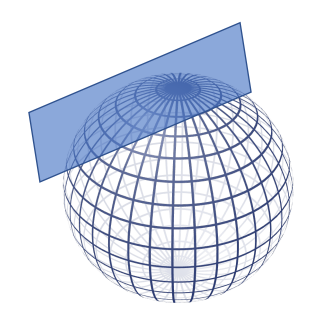
2. Pick the next set of parameters to be close to the current set, but in the direction that most changes the loss function for the better

3. Repeat

Slope vs gradient

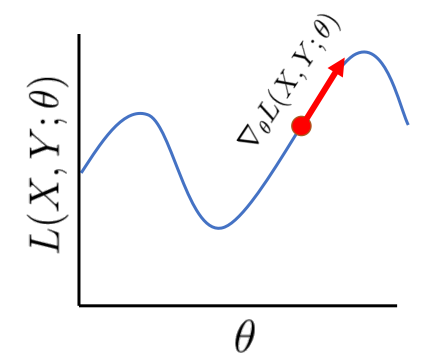
• Slope of $f(\theta)$ is a scalar describing a line perpendicular to the tangent of the function at that point .

• Gradient $\nabla f(\theta)$ is a vector describing a hyperplane perpendicular to the tangent at θ



What does the gradient tell us?

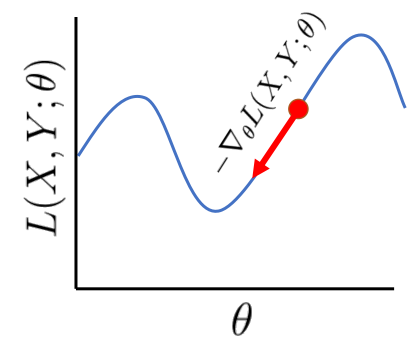
- If the loss function and hypothesis function encoded by the model are differentiable* (i.e., the gradient exists)
- We can evaluate the gradient for some fixed value of our model parameters θ and get the *direction* in which the loss *increases* fastest



^{*}or subdifferentiable

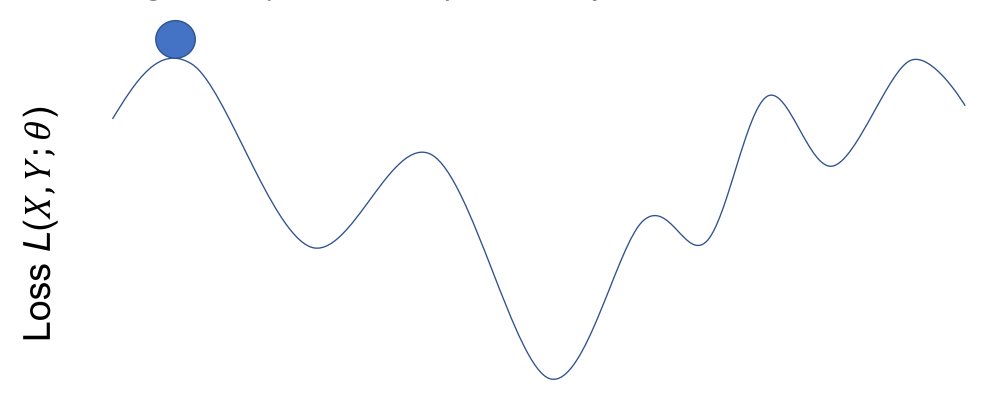
What does the gradient tell us?

• We want to decrease our loss, so let's go the other way instead



Gradient Descent: Promises & Caveats

- Much faster than guessing new parameters randomly
- Finds the global optimum only if the objective function is convex



 θ : the value of some parameter

Gradient Descent Pseudocode

Initialize $\theta^{(0)}$

Repeat until stopping condition met:

$$\theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(X, Y; \theta^{(t)})$$

Return $\theta^{(t_{max})}$

 $\theta^{(t)}$ are the parameters of the model at time step t

X, *Y* are the input data vectors and the output values.

 $\nabla L(X,Y;\theta^{(t)})$ is the gradient of the loss function with respect to model parameters $\theta^{(t)}$ η controls the step size

 $\theta^{(t_{max})}$ is the set of parameters that did best on the loss function.

Design choices

Initialize $\theta^{(0)}$

Repeat until stopping condition met:

$$\theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(X, Y, \theta^{(t)})$$

Return $\theta^{(t_{max})}$

- Initialization of θ
- Convergence criterion (i.e. when to stop)
- How much data to use (batch size)
- Step size for updating model parameters
- Choosing a loss function

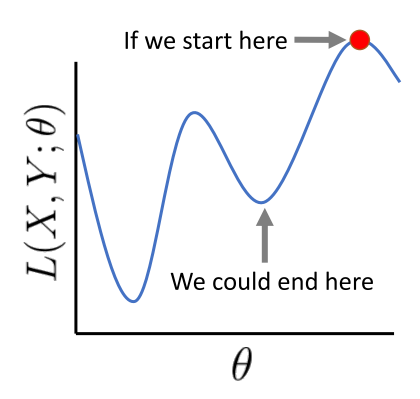
Parameter Initialization

Common initializations:

- $\bullet \ \theta^{(0)} = 0$
- $\theta^{(0)}$ = random values

What happens if our initialization is bad?

- Convergence to a *local* minimum
- No way to determine if you've converged to the global minimum



Convergence criterion: when to stop

• Stop when the gradient is close (within ε) to 0 (i.e., we reached a minimum)

Stop after some fixed number of iterations

 Stop when the loss on a validation set stops decreasing (This helps prevent overfitting)

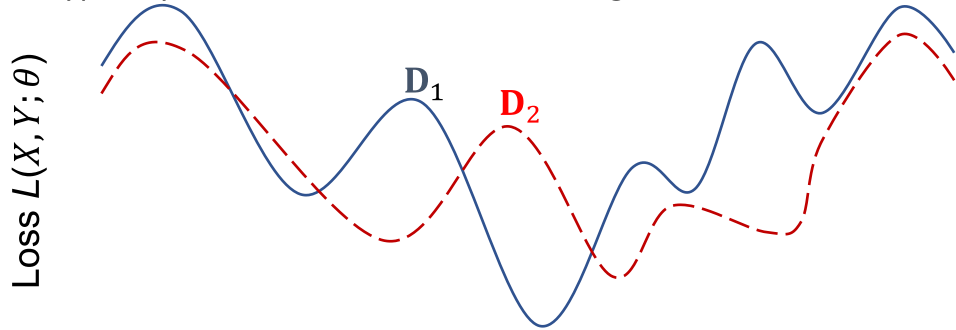
Batch Size: How much data?

- Call D the set X,Y pairs we measure loss on
- In **batch gradient descent**, the loss is a function of both the parameters θ and the set of all training data **D**. (What if if $|\mathbf{D}| > \text{memory}$?)
- In **stochastic gradient descent**, loss is a function of the parameters and a different single random training sample at each iteration.
- In mini-batch gradient descent, random subsets of the data (e.g. 100 examples) are used at each step in the iteration.

Different data, different loss

- Call D the set X,Y pairs we measure loss on.
- If D changes, then the landscape of the loss function changes

You typically won't know how it has changed.



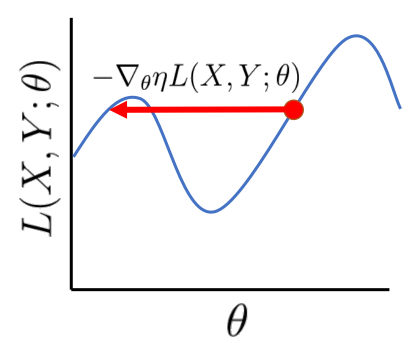
 θ : the value of some parameter

How much data to use in each step?

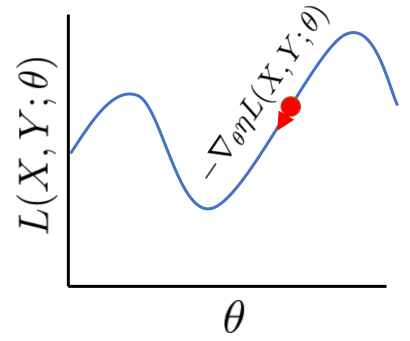
- All of it (batch gradient descent)
 - The *most accurate* representation of your training loss
 - It can be slow
 - Not possible if data does not fit in RAM
- Just one data point (stochastic gradient descent)
 - A noisy, inaccurate representation of your training loss
 - very fast
 - Random shuffling is important
- More than one data point, but less than all (mini-batch gradient descent)
 - Most common approach today
 - Balances *speed* and *accuracy*
 - Random shuffling is important
 - Usually want batch size to be as large as possible for your machine

Step Size: how far should we go?

- ullet The gradient we calculated was based on a fixed value of heta
- As we move away from this point, the gradient changes



If the step size is too large, we may overshoot the minimum



If the step size is too small, we need to take more steps (more computation)

Add Momentum

Initialize $\theta^{(0)}$, $V^{(0)}$

Repeat until stopping condition met:

$$V^{(t+1)} = mV^{(t)} - \eta \nabla L(X, Y, \theta^{(t)})$$

$$\theta^{(t+1)} = \theta^{(t)} + V^{(t+1)}$$

Return $\theta^{(t_{max})}$

There are many variants on gradient descent

 Lots of kinds of momentum/step size selection algorithms (e.g. ADAM)

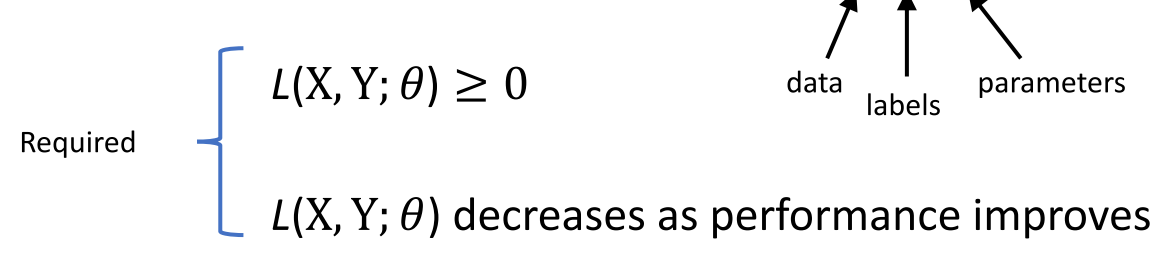
Lots of 2nd order algorithms (e.g. BGFS)

This is an entire field of study.

Check out classes taught in IEMS on this.

Loss functions

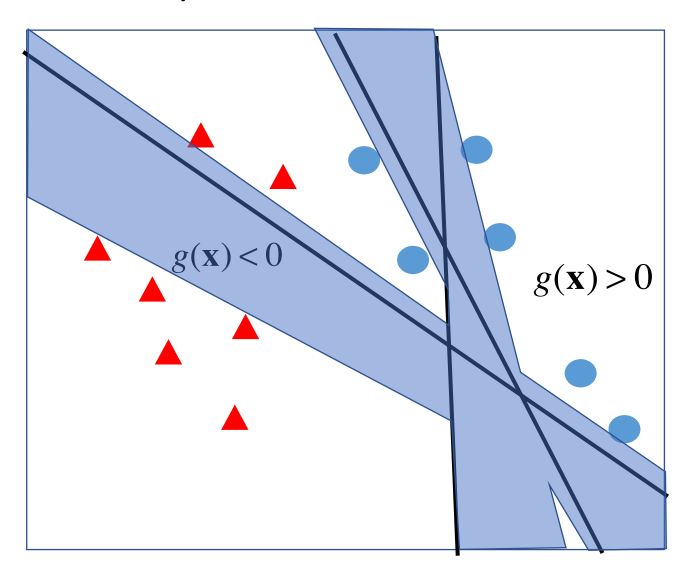
A good objective (loss) function $L(X, Y; \theta)$



Required for gradient descent $L(X,Y;\theta)$ is differentiable*, with respect to θ helpful For gradient descent $\mathbf{0} < |\nabla L| \ll \infty$

^{*}or subdifferentiable

Example: 0 1 loss



$$g(\mathbf{x}) = w_0 + w_1 x_1 + w_2 x_2 = 0$$

$$SSE = \sum_{i}^{n} (y_i - h(\mathbf{x}_i))^2$$

SSE is same everywhere in the blue Gradient 0 in the blue region!

The 0 1 Loss function

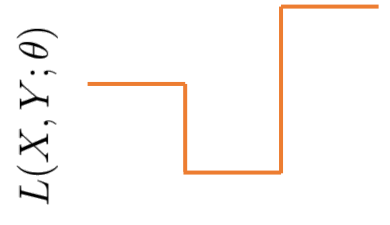
ullet A generic term for machine learning model parameters is heta

• Loss = 1 if $y \neq h_{\theta}(x)$, else it's 0

A count of mislabeled items

Results in a step function

Not useful for for gradient descent



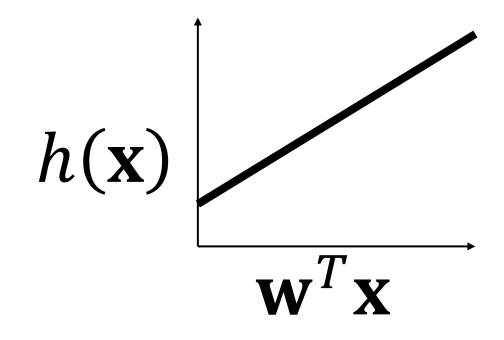
 θ

Perceptron Problem: The step function

$$h(\mathbf{x})$$
 $T_{\mathbf{X}}$

$$h(x) = \begin{cases} 1 & if \ 0 < \mathbf{w}^T \mathbf{x} \\ -1 & else \end{cases}$$

Solution: Remove the step function



$$h(\mathbf{x}) = \mathbf{w}^T \mathbf{x}$$

Squared loss: we now have a gradient

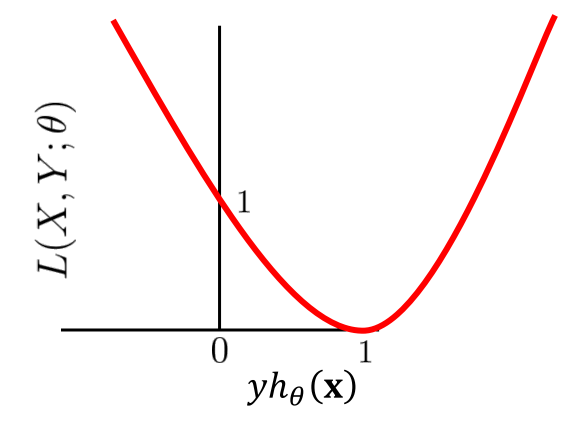
• Our hypothesis function is now $h_{\theta}(\mathbf{x})$ where θ are the model parameters.

• We write our loss function as...

$$L_s(X, Y; \theta) = \frac{1}{2N} \sum_{i=1}^{N} (y_i - h_{\theta}(x_i))^2$$

• If we use a linear model, then..

$$h_{\theta}(\mathbf{x}) = \theta^T \mathbf{x}$$



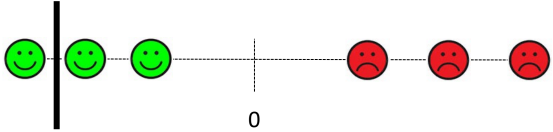
A simple example: where do you draw the line?

Happy faces have label y = +1 and sad faces have label y = -1.

We have a linear model with 2 parameters: $\hat{y} = \mathbf{w}^T \mathbf{x} = w_0 x_0 + w_1 x_1$

Our loss function will be sum-of-squared-errors:

$$L(X, Y, \mathbf{w}) = \frac{1}{2N} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$



How does the loss change as we move the line defined by w_0 ? Can we use that to decide where to move it? What does w_1 do?

Measuring loss for a linear unit

• Model's hypothesis $h_{\theta}(\mathbf{x})$ function outputs a label estimate \hat{y} , given its parameters θ . Let's call them the weights, \mathbf{w}

$$\hat{\mathbf{y}} = h_{\theta}(\mathbf{x}) = \mathbf{w}^T \mathbf{x}$$

• Sum of squared errors loss function:

 \mathbf{y}_i is the true label for example i

$$L(X,Y,\mathbf{w}) = \frac{1}{2N} \sum_{i=1}^{N} (\mathbf{y}_i - \widehat{\mathbf{y}}_i)^2$$
This ½ makes the derivative simpler

N is a normalizing factor so different batch sizes have comparable loss. Safe to remove.

i is the index to the ith example \mathbf{x}_i and its label \mathbf{y}_i

If we consider a single example, then...

$$L(X, Y, \mathbf{w}) = \frac{1}{2N} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$

Setting the number of data points N = 1 results in...

$$L(X, Y, \mathbf{w}) = \frac{1}{2} (y - \hat{y})^2$$

For each dimension i, take the partial derivative

$$\frac{\partial L}{\partial w_i} = \frac{\partial L}{\partial \hat{y}} \frac{\partial \hat{y}}{\partial w_i}$$
 gives the change of our loss function L with respect to weight w_i

Our loss function is :
$$L = \frac{1}{2}(y - \hat{y})^2$$
$$= \frac{y^2}{2} + \frac{\hat{y}^2}{2} - y\hat{y}$$

therefore...
$$\frac{\partial L}{\partial \hat{v}} = \hat{y} - y$$

For each dimension i, take the partial derivative

$$\frac{\partial L}{\partial w_i} = \frac{\partial L}{\partial \hat{y}} \frac{\partial \hat{y}}{\partial w_i}$$
 gives the change of our loss function L with respect to weight w_i

From the previous slide.... $\frac{\partial L}{\partial \hat{y}} = \hat{y} - y$

Our estimator is a linear unit : $\hat{y} = \mathbf{w}^T \mathbf{x}$

therefore... $\frac{\partial L}{\partial \hat{y}} = \mathbf{w}^T \mathbf{x} - y$

Let's calculate $\frac{\partial \hat{y}}{\partial w_i}$

Our estimator is : $\hat{y} = \mathbf{w}^T \mathbf{x} = w_0 x_0 + \dots w_i x_i + \dots w_d x_d$

Now... w_i is the only parameter we're varying right now.

So $\forall w_j$ where $j \neq i$ are constant in this partial derivative.

Therefore,
$$\frac{\partial \hat{y}}{\partial w_i} = x_i$$

Portion of the gradient for a linear model for weight i

$$\frac{\partial L}{\partial w_i} = \frac{\partial L}{\partial \hat{y}} \frac{\partial \hat{y}}{\partial w_i} = (\mathbf{w}^T \mathbf{x} - y) x_i$$
$$= -(y - \mathbf{w}^T \mathbf{x}) x_i$$

Let's now bring back having N data points (instead of just 1)

$$\frac{\partial L}{\partial w_i} \propto -\sum_{i=1}^{N} (y - \mathbf{w}^T \mathbf{x}) x_i$$

Means "proportional to". If you don't want to deal with a scaling factor like 1/N, you use this

For the whole set of model parameters...

From the previous slide: $\frac{\partial L}{\partial w_i} \propto -\sum_{i=1}^{N} (y - \mathbf{w}^T \mathbf{x}) x_i$

$$\nabla L(X,Y;\theta^{(t)}) = \left[\frac{\partial L}{\partial w_0}, \dots \frac{\partial L}{\partial w_i}, \frac{\partial L}{\partial w_i}, \frac{\partial L}{\partial w_d}\right]$$

The gradient can now be used here

Initialize $\theta^{(0)}$

Repeat until stopping condition met:

$$\theta^{(t+1)} = \theta_t - \eta \nabla L(X, Y; \theta^{(t)})$$

Return $\theta^{(t_{max})}$

 $\theta^{(t)}$ are the parameters of the model at time step t

 $\nabla L(X,Y;\theta^{(t)})$ is the gradient of the loss function with respect to model parameters $\theta^{(t)}$

 η controls the step size

 $\theta^{(t_{max})}$ is the set of parameters that did best on the loss function.

Squared loss: we now have a gradient

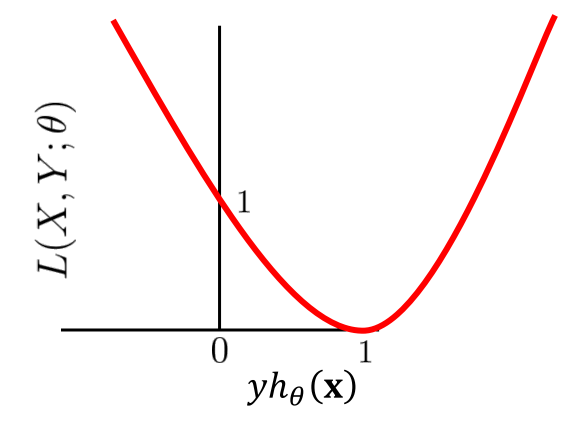
• Our hypothesis function is now $h_{\theta}(\mathbf{x})$ where θ are the model parameters.

• We write our loss function as...

$$L_s(X, Y; \theta) = \frac{1}{2N} \sum_{i=1}^{N} (y_i - h_{\theta}(x_i))^2$$

• If we use a linear model, then..

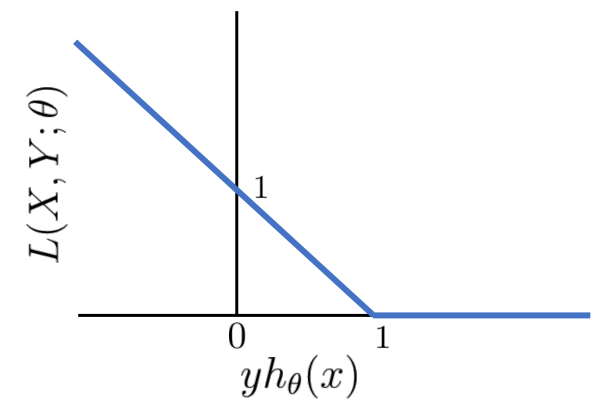
$$h_{\theta}(\mathbf{x}) = \theta^T \mathbf{x}$$



Hinge Loss

$$L_H(X, Y; \theta) = \frac{1}{N} \sum_{i=1}^{N} \max(0, 1 - y_i h_{\theta}(x_i))$$

 Loss only happens if the data is on the wrong side of the line.



Hinge Loss

The loss function:

$$L_H(X, Y; \theta) = \frac{1}{N} \sum_{i=1}^{N} \max(0, 1 - y_i h_{\theta}(x_i))$$

It gradient is...

Tradient is...
$$\nabla_{\theta}L_H(X,Y;\theta)=\frac{1}{N}\sum_{i=1}^N[[1-y_ih_{\theta}(x_i)>0]](-y_i\nabla_{\theta}h_{\theta}(x_i))$$

For example, if we use a linear model $h_{\theta}(x_i) = \theta^T x_i$,

$$\nabla_{\theta} L_H(X, Y; \theta) = \frac{1}{N} \sum_{i=1}^{N} [[1 - y_i \theta^T x_i > 0]](-y_i x_i)$$

Regularization

Revisiting Overfitting

- Overfitting occurs when your model begins to "memorize" the training data
 - Can detect overfitting from an increasing gap between training and validation loss.
 - Performance on the training set improves, but performance on the validation set does not.



Revisiting Overfitting: Regularization

- Big idea (**Occam's Razor**) Given two models with equal performance, prefer the *simpler* model.
 - E.g., models with fewer parameters or smaller coefficients
- Regularization can be applied to any loss function

$$L_R(X, Y; \theta) = L(X, Y; \theta) + \lambda R(\theta)$$

 ${\bf \cdot}$ The amount of regularization is controlled by the hyperparameter λ

L1- and L2-regularization

• Recall the l_p -norm:

$$\ell_p(\theta) = \sqrt[p]{\sum_{i=1}^d |\theta_i|^p}$$

• l_1 -regularization penalizes high values of the l_1 -norm of the model parameters:

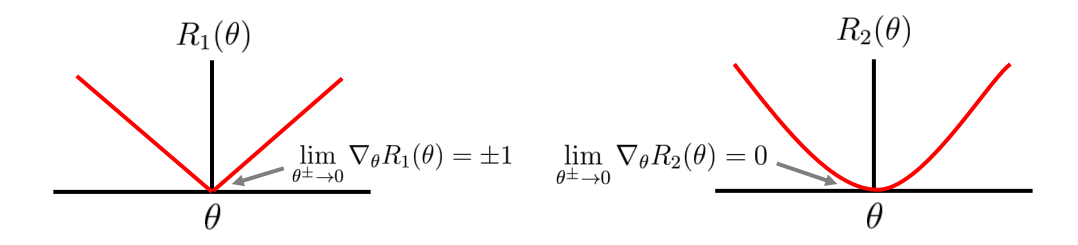
$$R_1(\theta) = \sum_{i=1}^d |\theta_i|$$

• l_2 -regularization penalizes high values of the l_2 -norm:

$$R_2(\theta) = \frac{1}{2} \sum_{i=1}^{d} |\theta_i|^2$$

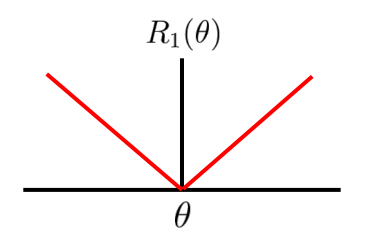
L1-regularization and sparsity

- L1-regularization encourages the model parameters to be *sparse*
 - This is a form of feature selection
 - Only features with non-zero coefficients contribute to the model's prediction
- This is because the gradient of L1-regularization moves model parameters towards 0 at a constant rate



L1-regularization and sparsity

- The gradient of the L1-regularizer is bounded (between -1 and +1, inclusive) but not unique at $\theta=0$.
- Arbitrarily set the gradient at this point to 0.
- The resulting function is the *sign* function



$$\nabla_{\theta} R_1(\theta) = \operatorname{sign}(\theta) = \begin{cases} +1, \ \theta > 0 \\ 0, \ \theta = 0 \\ -1, \ \theta < 0 \end{cases}$$

Regularization and offset (aka bias)

• Many ML models include a bias term, b.

• Example: A linear model:
$$h_{ heta}(x) = heta^T x + b$$

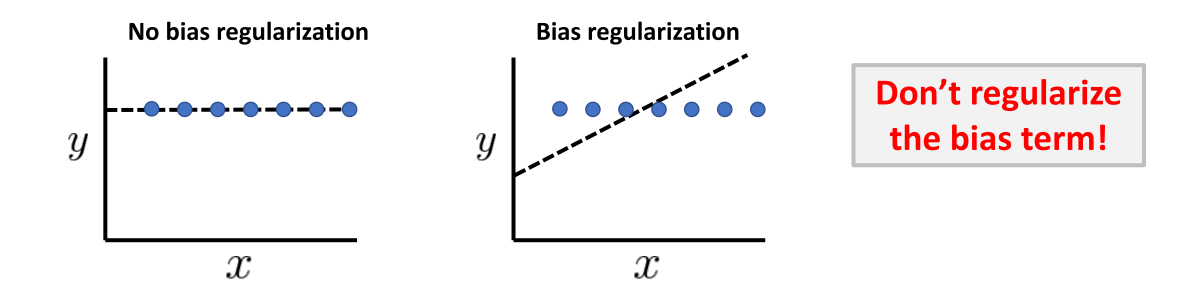
• Or equivalently, by augmenting θ and x, like we did with perceptrons...

$$\theta' = [\theta_1, \theta_2, \dots, \theta_d, b], \ x' = [x_1, x_2, \dots, x_d, 1]$$

What happens if we regularize the bias term?

Regularization and offset (aka bias)

- Recall that "regularizing" a model parameter means encouraging that model parameter to tend towards 0.
- How would a linear model represent horizontal line?
- How does shrinking the bias affect its ability to do so?



A familiar example

 A lot of algorithms can be constructed by simply combining a loss function and a regularizer

• For example, hinge loss and L2-regularization with a linear hypothesis

$$L_H(S, Y; \theta) + \lambda R_2(\theta) = \frac{1}{N} \sum_{i=1}^{N} \max(0, 1 - y_i \theta^T x_i) + \frac{\lambda}{2} \|\theta\|_2^2$$

Does this look like something we've studied?

Soft margin SVMs!

Learning a maximum margin separator via gradient descent

- Finding an exact solution for SVMs can be difficult
 - Convex quadratic programming problem
 - $\sim O(n^3)$
- Finding a good approximate solution for SVMs using gradient descent is much easier and computationally faster

Multi-class Classification

Using your linear model, once learned

• Regression:

$$\hat{y} = h_{\theta}(x)$$

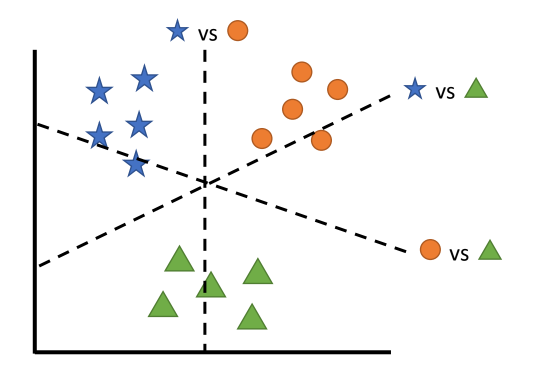
Binary classification:

$$\hat{y} = \begin{cases} 1, & \text{if } h_{\theta}(x) > 0 \\ -1, & \text{otherwise} \end{cases}$$

- What about classification with more than 2 classes (e.g., C classes)? We will discuss 2 approaches:
 - One-vs-One classification
 - One-vs-All classification

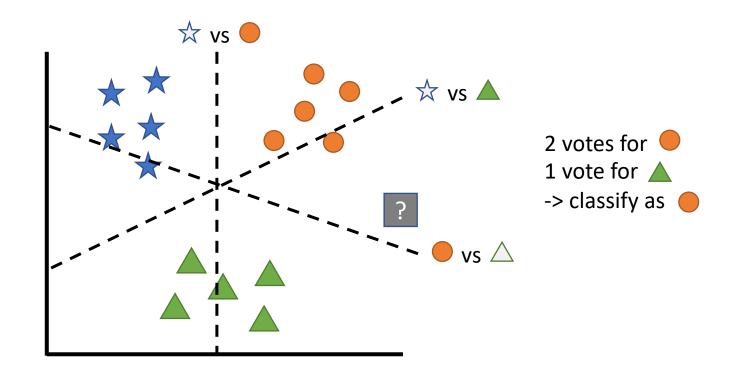
One-vs-One Classification

- Train a binary classifier to disambiguate between each *pair* of classes
- Final prediction is the majority vote among all binary classifiers



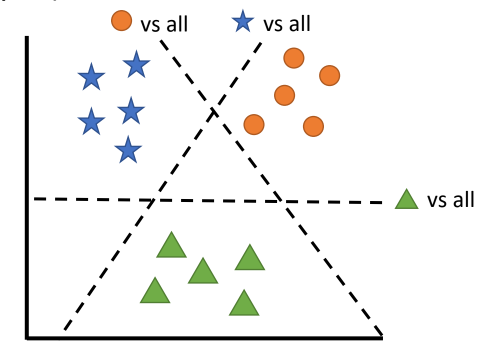
One-vs-One Classification

- Train a binary classifier to disambiguate between each *pair* of classes
- Final prediction is the majority vote among all binary classifiers



One-vs-All Classification

- Train a binary classifier on whether an example does or does not belong to a class
- Predict based on the highest confidence score (i.e., the regression output)



$$\hat{y} = \underset{c \in C}{\operatorname{argmax}} \ h_{\theta_c}(x)$$