## **Gradient Descent**

Many problems in Machine Learning are framed as optimization problems

- ullet Find the choice of parameters  $\Theta$
- That minimizes a Loss function

The best (optimal)  $\Theta$  is the one that minimizes the Average (across training examples) Loss

$$\Theta^* = \operatorname*{argmin}_{\Theta} \mathcal{L}_{\Theta}$$

Many Classical ML problems are designed such that  $\Theta^*$  has a closed-form solution • Maximum likelihood estimates for Linear Regression Closed form solutions, however, may only be feasible for Loss function of restricted form.

When a closed form solution is not possible, we may find  $\Theta^*$  via a search process known as Gradient Descent. In the Deep Learning part of the course, virtually all Loss functions will require this form of solution.

## Loss functions, review

- $\hat{\mathbf{y}}^{(i)} = h(\mathbf{x^{(i)}}; \Theta)$ , the prediction for example  $\mathbf{x^{(i)}}$  with target label  $\mathbf{y^{(i)}}$
- Per-example loss

$$\mathcal{L}_{\Theta}^{(\mathbf{i})} = L(h(\mathbf{x^{(i)}}; \Theta), \mathbf{y^{(i)}}) = L(\hat{\mathbf{y}^{(i)}}, \mathbf{y})$$

 The Loss for the entire training set is simply the average (across examples) of the Loss for the example

$$\mathcal{L}_{\Theta} = rac{1}{m} \sum_{i=1}^{m} \mathcal{L}_{\Theta}^{(\mathbf{i})}$$

Two common forms of L are Mean Squared Error (for Regression) and Cross Entropy Loss (for classification).

# **Optimization**

How do we find the  $\Theta^*$  that minimizes  $\mathcal L$  ?

$$\Theta^* = \operatorname*{argmin}_{\Theta} \mathcal{L}_{\Theta}$$

One way is via a search-like procedure known as Gradient Descent:

We start with an initial guess for  $\Theta$  and then:

• Evaluate  $\mathcal{L}_{\Theta}$  across training examples

$$\langle \mathbf{X}, \mathbf{y} \rangle = [\mathbf{x^{(i)}}, \mathbf{y^{(i)}} | 1 \leq i \leq m]$$

- Make a small change to  $\Theta$  that results in a reduced  $\mathcal{L}_{\Theta}$
- Repeat until  $\mathcal{L}_{\Theta}$  stops decreasing

Fortunately, for many functions  $\mathcal{L}_\Theta$  we can use calculus to guide the small change in  $\Theta$  in the direction of reduced  $\mathcal{L}_\Theta$ 

$$rac{\partial}{\partial \Theta} \mathcal{L}_{\Theta}$$

is the partial derivative of  $\mathcal{L}_{\Theta}$  with respect to  $\Theta$ .

• For a unit increase in  $\Theta$ :  $\mathcal{L}_{\Theta}$  increases by  $\frac{\partial}{\partial \Theta}\mathcal{L}_{\Theta}$ 

Thus, to decreases  $\mathcal{L}_{\Theta}$  we only need to add an increment in  $\Theta$  proportional to the negative of the partial derivative.

Since  $\Theta$  is a vector, the partial derivative is *also* a vector and is called the *gradient*. The iterative process we described is called gradient descent as it follows the negative of the gradient towards a minimum for  $\Theta$ .



```
In [5]: | def f(x):
             return x**2
         def deriv(f,x_0):
             h = 0.000000001
                                              #step-size
             return (f(x \ 0 + h) - f(x \ 0))/h
         def tangent(f, \times 0, \times=None):
             y 0 = f(x 0)
             slope = deriv(f, x 0)
             if x is not None:
                 r = 2
                 xmin, xmax = np.min(x), np.max(x)
                 xlo, xhi = max(x 0 - r, xmin), min(x 0 + r, xmax)
             else:
                 r = 2
                 xlo, xhi = x 0 - r, x 0 + r
             xline = np.linspace(xlo, xhi, 10)
             yline = y 0 + slope*(xline - x 0)
             return xline, yline
         def plot tangent(f, x_s, x, ax, show_tangent=True):
             # Plot function
             = ax.plot(x, f(x))
             # Plot tangent point x s
             y s = f(x s)
             ax.scatter(x s, y s, color='r', s=90)
             # Plot tangent line
             if show tangent:
                 xtang, ytang = tangent(f, x s, x)
                 ax.plot(xtang, ytang, 'q--')
```

```
In [6]: def plot_step(f, x_s, x, show_tangent=True, visible=True):
    fig, ax = plt.subplots(1, 1, figsize=(12,6))

y_s = f(x_s)

# Plot the function, the point, and optionally: the tangent line
    _= plot_tangent(f, x_s, x, ax, show_tangent=show_tangent)

= ax.set_xlabel("$\Theta$", fontsize=16)
    _= ax.set_ylabel("$L$", fontsize=16, rotation=0)

if not visible:
    plt.close(fig)

return fig, ax
```

```
In [7]: def plot gradient descent(max steps=4, alphas=[ 0.1, 0.4, 0.7, 1.0 ]):
             fig, axs = plt.subplots(len(alphas), max steps, figsize=(20,min(12, 6 * len
         (alphas))))
             axs = axs.reshape( (len(alphas), max steps) ) # Take care of special case w
         here len(alpha) == 1
             for a idx, alpha in enumerate(alphas):
                 x s = x 0
                 for step in range(0, max steps):
                     ax = axs[a idx, step]
                     = ax.set xlabel("$\Theta$", fontsize=16)
                     = ax.set ylabel("$L$", fontsize=16, rotation=0)
                     = ax.set title('$\\alpha$={a:3.2f}'.format(a=alpha))
                     y s = f(x s)
                     # Obtain tangent line at x0
                     = plot tangent(f, x s, x, ax)
                     # Update x s
                     slope = deriv(f, x s)
                     x s = x s + alpha \overline{*} (- slope)
             _= fig.tight_layout()
             plt.close(fig)
             return fig, axs
```

```
In [8]: alpha = 0.4
 x = np.linspace(-5, +5, 30)
```

## **Gradient Descent: Overview**

Let's illustrate the process with an example

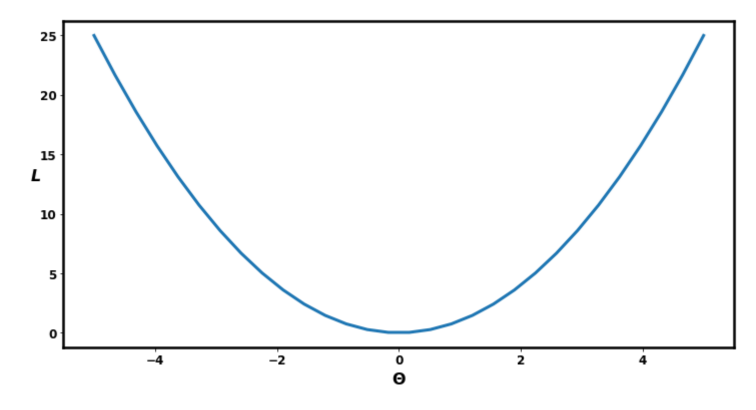
Let's plot a simple loss function as an illustration.

In this simple example:  $\Theta$  is a vector of length 1.

```
In [9]: fig, ax = plt.subplots(1,1, figsize=(12,6))
    _= ax.plot(x, f(x), linewidth=3)
    _= ax.set_xlabel("$\Theta$", fontsize=16)
    _= ax.set_ylabel("$L$", fontsize=16, rotation=0)
    plt.close(fig)
```

In [10]: fig

Out[10]:



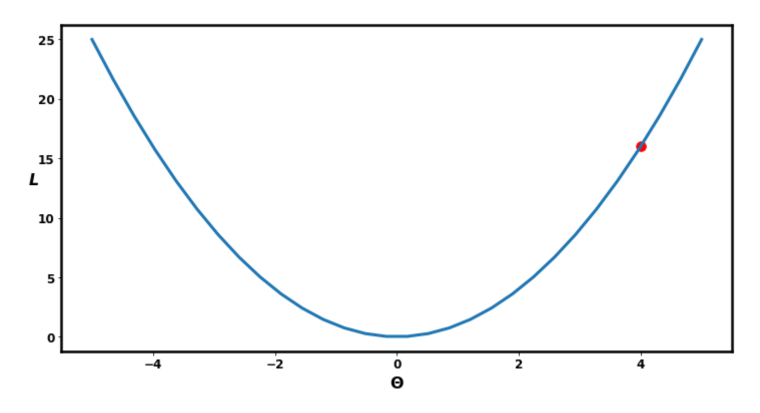
Let's start off with a guess for  $\Theta$ 

In [11]: 
$$x_0 = 4$$
  
 $x_s = x_0$ 

```
In [12]: fig, ax = plot_step(f, x_s, x, show_tangent=False, visible=False)
```

In [13]: fig

Out[13]:



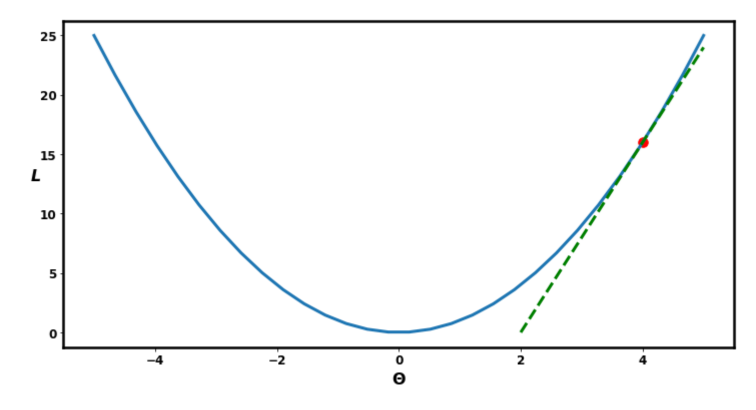
Clearly not at a minimum.

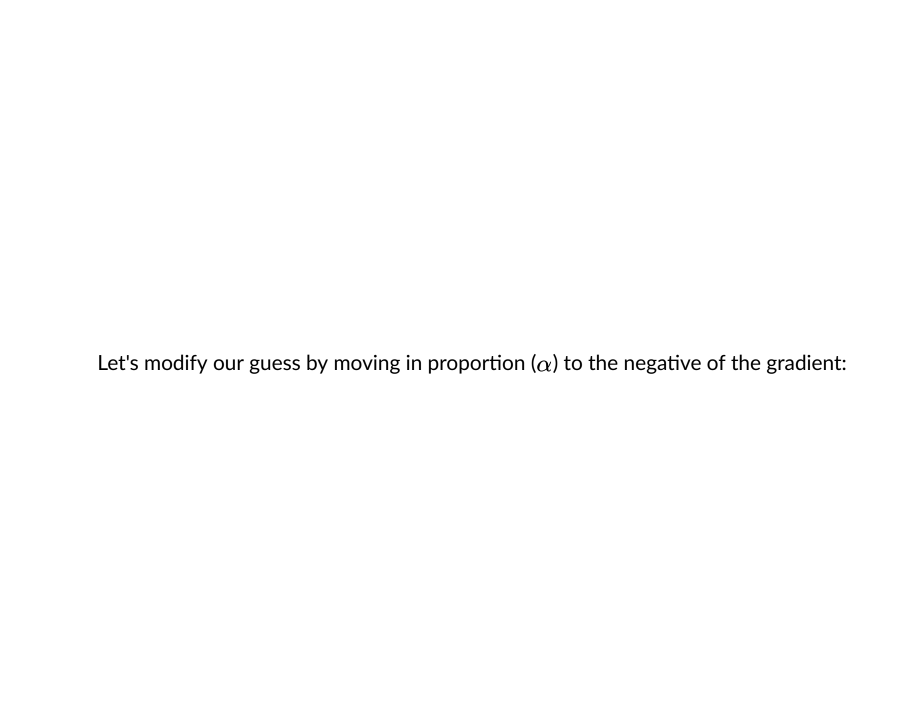
Compute the gradient of  $\mathcal{L}_\Theta$  at initial guess  $x_s$ 

```
In [14]: x_s = x_0
fig, ax = plot_step(f, x_s, x, visible=False)
```

In [15]: fig

Out[15]:



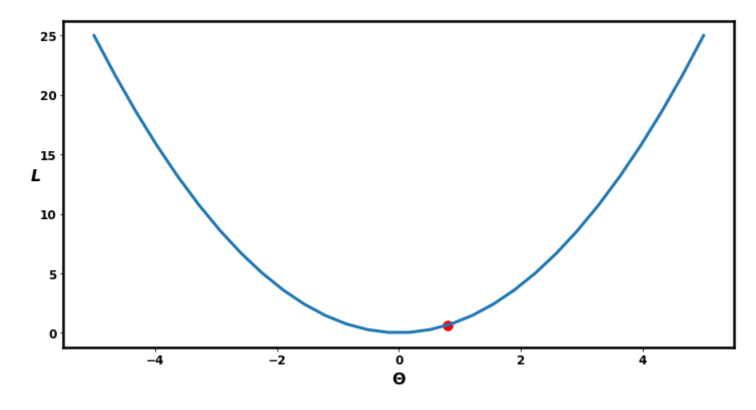


```
In [16]: # Update x_s
slope = deriv(f, x_s)
x_s = x_s + alpha * (- slope)
```

```
In [17]: fig, ax = plot_step(f, x_s, x, show_tangent=False, visible=False)
```

In [18]: fig

Out[18]:



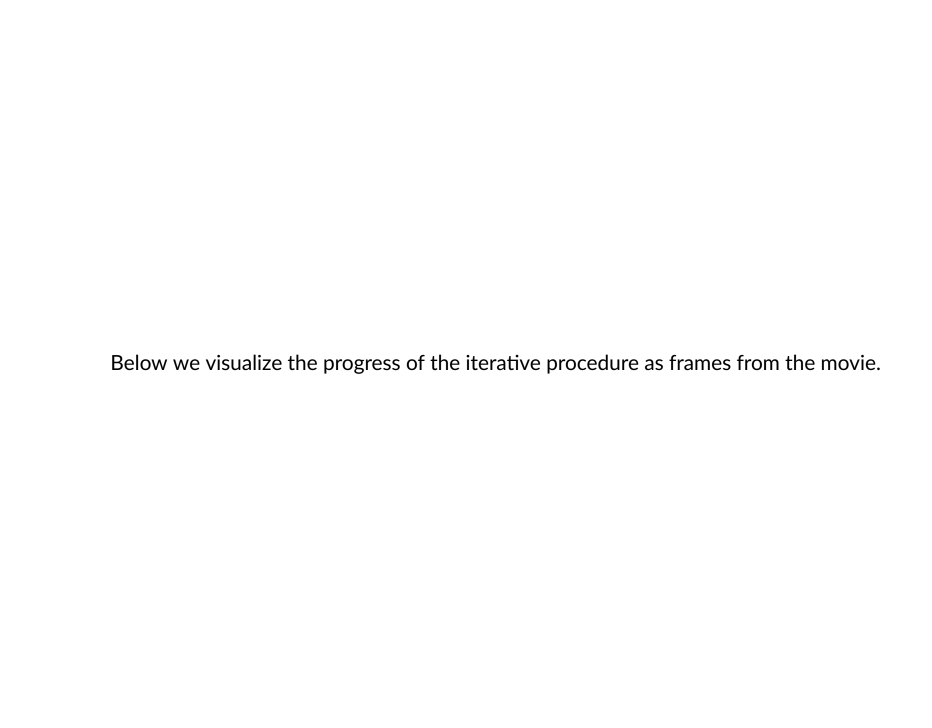
By following the gradient as we did: we wind up at a new  $\Theta$  where  $\mathcal{L}_{\Theta}$  is reduced compared ot that at the original guess.

Taking the gradient of the  $\mathcal L$  at the new point, we continue the iterative process.

```
In [19]: if CREATE_MOVIE:
    _= gdh.create_gif2(x, f, x_0, out="images/gd.gif", alpha=alpha)
```

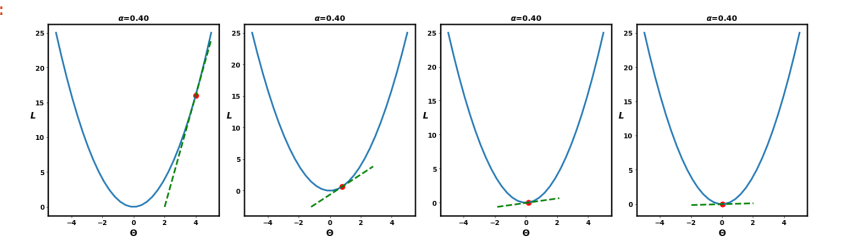
```
In [33]: _= gdh.display_gif("images/gd.gif")
```

```
In [21]: fig, axs = plot_gradient_descent(alphas= [ alpha ])
```



In [22]: fig

#### Out[22]:



## **Gradients: vector derivatives**

We illustrated the use of Gradient Descent to find the minimum of a function of a single variable.

The same procedure works when the function is of higher dimension.

Let's illustrate with the MSE Loss often used in Linear Regression, when  $\mathbf{x^{(i)}}$  (and hence  $\Theta$ ) is of dimension n.

$$\mathbf{y} = \Theta^T \cdot \mathbf{x}$$

With (n+1) features (including the constant)

- $\Theta$  is a vector of length (n+1)
- $rac{\partial}{\partial \Theta} \mathcal{L}_{\Theta} = 
  abla_{\Theta} \mathcal{L}_{\Theta}$ , is a vector of length (n+1)

$$abla_{\Theta}\mathcal{L}_{\Theta} = egin{pmatrix} rac{\partial}{\partial \Theta_0}\mathcal{L}_{\Theta} \ rac{\partial}{\partial \Theta_1}\mathcal{L}_{\Theta} \ dots \ rac{\partial}{\partial \Theta_n}\mathcal{L}_{\Theta} \end{pmatrix}$$

Using MSE Loss as the Loss function

$$\mathcal{L}_{\Theta} = ext{MSE}(\mathbf{y}, \hat{\mathbf{y}}, \Theta) = rac{1}{m} \sum_{i=1}^{m} (\mathbf{y^{(i)}} - \hat{\mathbf{y}^{(i)}})^2$$

$$abla_{\Theta} \mathcal{L}_{\Theta} = egin{pmatrix} rac{\partial}{\partial \Theta_0} \mathrm{MSE}(\mathbf{y}, \hat{\mathbf{y}}, \Theta) \ rac{\partial}{\partial \Theta_1} \mathrm{MSE}(\mathbf{y}, \hat{\mathbf{y}}, \Theta) \ dots \ rac{\partial}{\partial \Theta_n} \mathrm{MSE}(\mathbf{y}, \hat{\mathbf{y}}, \Theta) \end{pmatrix}$$

#### Whereas in our code

- We computed derivatives numerically
- We will compute them below analytically, using calculus

Analytic (closed form) derivatives are much faster to compute.

• During the Deep Learning part of the course, we will see how to *automatically* obtain analytic derivatives

$$\frac{\partial}{\partial \Theta_{j}} \text{MSE}(\mathbf{y}, \hat{\mathbf{y}}, \Theta) = \frac{1}{m} \sum_{i=1}^{m} \frac{\partial}{\partial \Theta_{j}} (\mathbf{y^{(i)}} - \hat{\mathbf{y}^{(i)}})^{2} \qquad \text{definition}$$

$$= \frac{1}{m} \sum_{i=1}^{m} 2 * (\mathbf{y^{(i)}} - \hat{\mathbf{y}^{(i)}}) \frac{\partial}{\partial \Theta_{j}} (-\hat{\mathbf{y}^{(i)}}) \qquad \text{chain rule}$$

$$= -\frac{1}{m} \sum_{i=1}^{m} 2 * (\mathbf{y^{(i)}} - \hat{\mathbf{y}^{(i)}}) \frac{\partial}{\partial \Theta_{j}} (\Theta * \mathbf{x^{(i)}}) \qquad \hat{\mathbf{y}^{(i)}} = \Theta^{T} \cdot \mathbf{x^{(i)}}$$

$$= -\frac{1}{m} \sum_{i=1}^{m} 2 * (\mathbf{y^{(i)}} - \hat{\mathbf{y}^{(i)}}) \mathbf{x_{j}^{(i)}}$$

$$= -\frac{2}{m} \sum_{i=1}^{m} (\mathbf{y^{(i)}} - \hat{\mathbf{y}^{(i)}}) \mathbf{x_{j}^{(i)}}$$

Thus the gradient for Linear Regression can be written in matrix form as

$$abla_{m{ heta}} \operatorname{MSE}(X, m{ heta}) = = rac{2}{m} \mathbf{X}^T (\Theta^T \mathbf{X} - \mathbf{y}) \quad \operatorname{since} \hat{\mathbf{y}} = \Theta^T \mathbf{x}$$

This will be particularly useful when working with NumPy as the gradient calculation is a vector operation that is implemented so as to be fast.

## **Gradient Descent versus MLE**

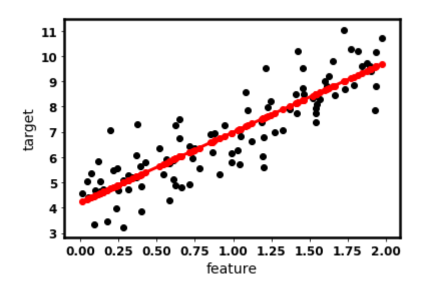
For Linear Regression, there is a closed form solution for finding the optimal  $\Theta$ .

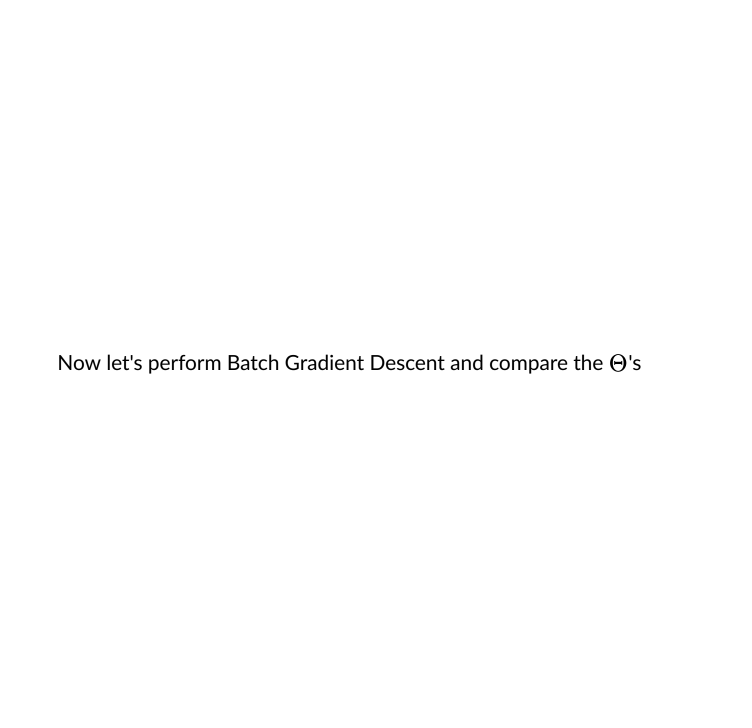
We will demonstrate that the Gradient Descent search comes arbitrarily close.

Let's illustrate Batch Gradient Descent on an example.

First, we use sklearn's LinearRegression as a baseline against which we will compare the  $\Theta$  obtained from Gradient Descent.

```
In [23]: X_lr, y_lr = gdh.gen_lr_data()
    clf_lr = gdh.fit_lr(X_lr,y_lr)
    fig, ax = gdh.plot_lr(X_lr, y_lr, clf_lr)
    theta_lr = (clf_lr.intercept_, clf_lr.coef_)
```





The  $\Theta$ 's are equal up to 15 decimal points.



alpha =  $0.1 \text{ n_iterations} = 1000 \text{ m} = 100 \text{ theta} = \text{np.random.randn}(2,1) \text{ for iteration in range}(\text{n_iterations}): gradients = <math>2/\text{m} * \text{X_b.T.dot}(\text{X_b.dot}(\text{theta}) - \text{y}) \text{ theta} = \text{theta} - \text{alpha} * \text{gradients}$ 

- We use the closed form, analytic expression for the gradient
- We update

$$\Theta = \Theta - \alpha * gradient$$

Notice that the "step size" ( $\alpha*$  gradient)

- Is "big" when the gradient is large
- Is "small" when the gradient is small (close to optimal)

Since the  $\Theta$ 's computed by Gradient Descent and Linear Regression are the same, it's no surprise that the predictions are too. • As demonstrated in the following code

```
In [25]: X_new = np.array([[0], [2]])
    gd_y_pred = gdh.predict(X_new, theta_lr)
    clf_y_pred = clf_lr.predict(X_new)

gd_y_pred == clf_y_pred
```

# **Gradient Descent in depth**

There are many subtleties to Gradient Descent.

As Gradient Descent will be a *key tool* in the Deep Learning part of the course, we briefly explore a few issues below.

## How big should lpha be ?

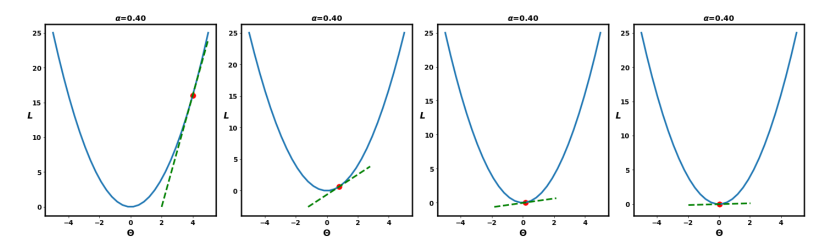
The "step size" we take along the direction of the gradient is  $\alpha$ .

Does the choice of  $\alpha$  matter?

Here are 4 steps with lpha=0.40

In [26]: fig, axs = plot\_gradient\_descent(alphas= [ alpha ])
fig

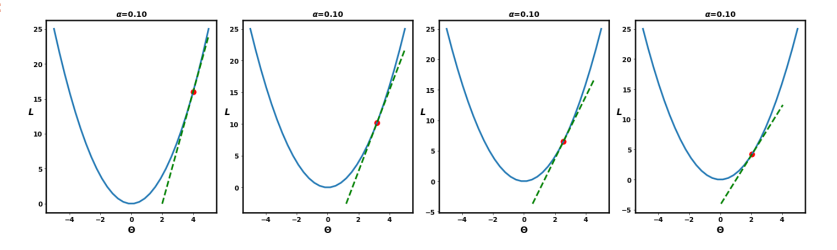
Out[26]:



And with a much smaller lpha=0.1

In [27]: fig, axs = plot\_gradient\_descent(alphas= [ 0.1 ])
fig

### Out[27]:

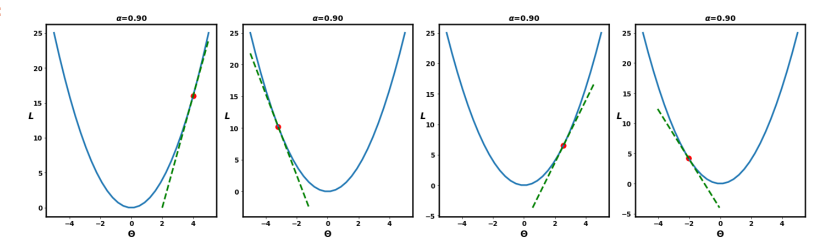


Convergence toward the optimal is much slower.

What if we used a larger lpha=0.9 ?

In [28]: fig, axs = plot\_gradient\_descent(alphas= [ 0.9 ])
fig

### Out[28]:



You can see that we over-shoot the optimal repeatedly. This may be problematic • For more complex loss functions: we may "skip" over a local optimum An adaptive learning rate schedule may be the solution:

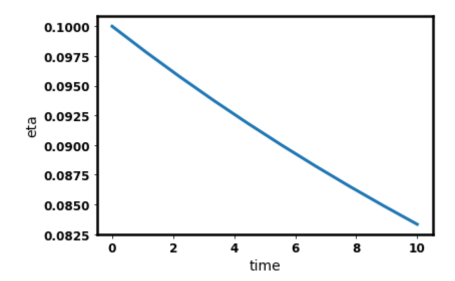
- Take big steps at first
- Take smaller steps toward end

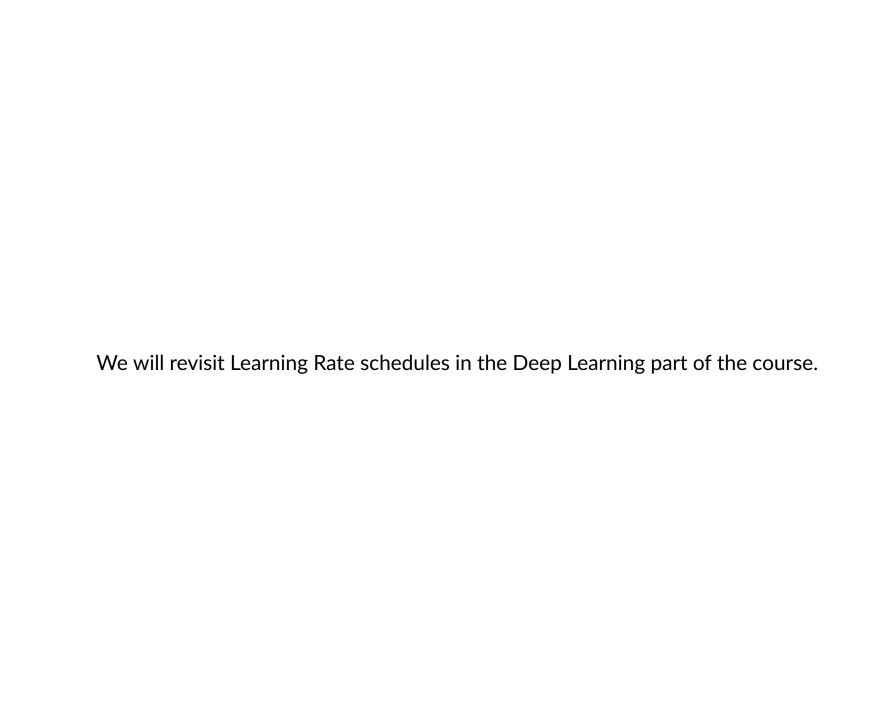
```
In [29]: t0, t1 = 5, 50 # learning schedule hyperparameters

def learning_schedule(t):
    return t0 / (t + t1)

t = np.linspace(0, 10, 10)

fig = plt.figure()
    ax = fig.add_subplot(1,1,1)
    _ =ax.plot(t, learning_schedule(t))
    _ = ax.set_xlabel("time")
    _ = ax.set_ylabel("eta")
```





### Mini batch Gradient Descent

The Average Loss function in Classical Machine Learning has the form

$$\mathcal{L}_{\Theta} = rac{1}{m} \sum_{i=1}^{m} \mathcal{L}_{\Theta}^{(\mathbf{i})}$$

That is, it is composed of m sub-expressions where m is the number of training examples.

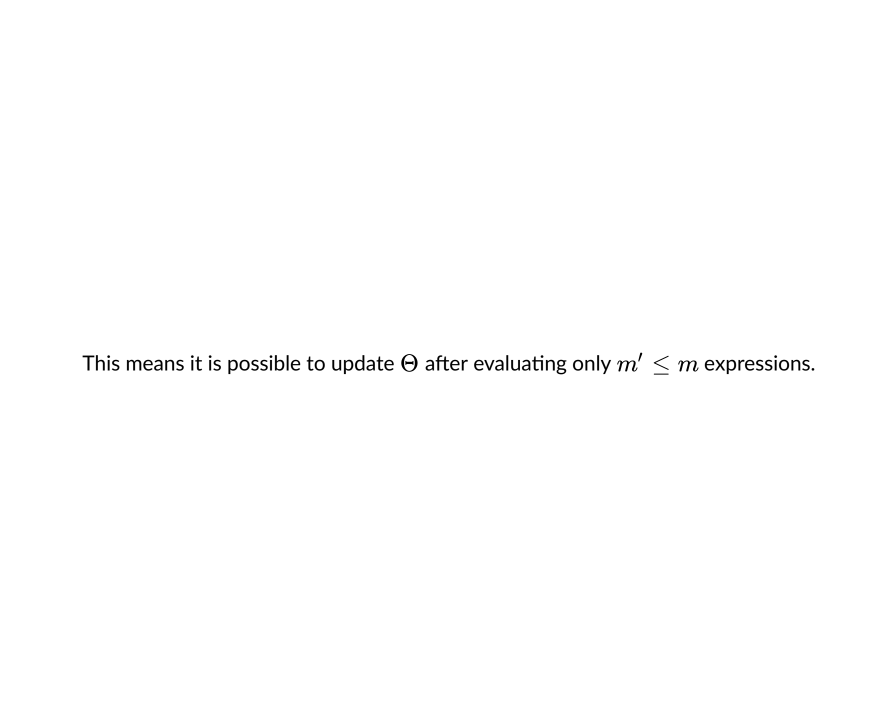
• Each subexpression requires a computation and a derivative

Thus, for large sets of training examples, Gradient Descent can be expensive.

It may be possible to approximate  $\mathcal{L}_{\Theta}$  using fewer than m expressions.

- ullet Choose a random subset ( of size  $m' \leq m$  ) of examples:  $I = \{i_1, \dots, i_{m'}\}$
- Approximate  $\mathcal{L}_{\Theta}$  on I

$$\mathcal{L}_{\Theta} pprox rac{1}{|I|} \sum_{i \in I} \mathcal{L}_{\Theta}^{(\mathbf{i})}$$



Whereas Gradient Descent computes an exact  $\mathcal{L}_{\Theta}$  to perform a single update of  $\Theta$ :

Mini batch Gradient Descent

- ullet Takes b=m/m' smaller steps, each updating  $\Theta$
- ullet Each small step using an approximation of  $\mathcal{L}_\Theta$  based on  $m' \leq m$  examples

#### It does this by

- Choosing batch size m'
- Partitioning the set of example indices  $\{i|1\leq i\leq m\}$ 
  - into b batches of size m'
  - lacksquare Batch  $i':b_{(i')}$  is one partition consisting of m' example indices
  - ullet Each small step uses a single batch to approximate  $\mathcal{L}_\Theta$  and update  $\Theta$

The collection of b small steps (comprising all examples) is called an *epoch* 

So one epoch of Mini batch Gradient Descent performs  $\boldsymbol{b}$  updates.

When batch size m'=m, we have our original algorithm known as Batch Gradient Descent.

How does one choose  $m' \leq m$ ?

- ullet Want m' large enough so approximations aren't too noisy
  - Don't want losses of the mini-batches of each epoch to be too different
- Often determined by *external* considerations
  - GPU memory (preview of Deep Learning)

# Initializing $\Theta$

As we will see in the Deep Learning part of the course

• Initial  $\Theta$  is not a trivial choice

Consider a Loss function like the Hinge Loss

- ullet Our initial choice of  $\Theta$  could leave us in a *flat* area of the Loss function
- No derivative, but maybe not optimal
- No way to escape!

## When to stop

Deciding when to stop the iterative process is another choice to be made

- Stop when decrease in  $\mathcal{L}_{\Theta}$  is "too small"

# Improvements to Gradient Descent

Simon Ruder survey (https://arxiv.org/abs/1609.04747)

<u>Gradient Descent Cheatsheet (https://towardsdatascience.com/10-gradient-descent-optimisation-algorithms-86989510b5e9)</u>

The update step

$$\Theta = \Theta - lpha * rac{\partial \mathcal{L}_{\Theta}}{\partial \Theta}$$

where  $\alpha$  is the learning rate.

The improvements to Gradient Descent modify

- $\alpha$ , the learning rate
- $\frac{\partial \mathcal{L}_{\Theta}}{\partial \Theta}$  the gradient

In order to be able to flexibly change the definition of both the gradient and the learning rate at each time step t, e will re-write the update step at time t as

$$\Theta_{(t)} = \Theta_{(t-1)} - lpha' * V_{(t)}$$

 $V_{(t)}$  will be our modified gradient and lpha' our modified learning rate.

## Momentum: modify the gradient

In vanilla Gradient Descent, the gradients at time (t-1) and time t are completely independent.

This has the potential for gradients to rapidly change direction (recall, they are a vector).

To smooth out jumps we could compute a modified gradient  $V_{(t)}$  as:

$$V_{(t)} = eta_V * V_{(t-1)} + (1-eta_V) * rac{\partial \mathcal{L}_{\Theta}}{\partial \Theta}$$

(Initialize  $V_0=0$ )

That is, the modified gradient is a weighted combination of the previous gradient and the new gradient.

Typically  $eta_V pprox 0.9$  so the old gradient dominates.

 $V_{\left(t
ight)}$  is the exponentially weighted moving average of the gradient.

Hence, there is "momentum" in the gradients in that they can't jump suddenly.

## RMSprop: Modify the learning rate

Let

$$S_{(t)} = eta_S * S_{(t-1)} + (1-eta_S) * \left(rac{\partial \mathcal{L}_{\Theta}}{\partial \Theta}
ight)^2$$

That is,  $S_{(t)}$  is the exponentially weighted *variance* of the gradient.

(Initialize  $S_0 = 0$ )

Rather than using a learning rate of  $\alpha$ , the RMSprop algorithm uses

$$lpha' = rac{1}{\sqrt{S_{(t)} + \epsilon}} * lpha$$

The intuition is that if the gradient with respect to  $\Theta_j$  is noisy (i.e., large variance) we want to damp updates in that component.

This also has the advantage that

- A rarely updated element  $\Theta_i$ , having a low variance,
- Will have a relatively larger update when it is encountered than a more frequently encountered feature.

Typically  $eta_S pprox 0.9$  so the old variance dominates.

Why the extra  $\epsilon$ ? We've seen this before (e.g.,  $\log(x+\epsilon)$ ): it's to avoid mathematical issues of certain functions (inverse, log) when the argument is 0.

## AdaM: Modify both the gradient and the learning rate

The AdaM (Adaptive Moment) algorithm modifies both

- The gradient
- The learning rate

via exponentially weighted moving averages of the gradient as well as its variance.

$$egin{aligned} V_{(t)} &= eta_V * V_{(t-1)} + (1-eta_V) rac{\partial \mathcal{L}_{\Theta}}{\partial \Theta} \ S_{(t)} &= eta_S * S_{(t-1)} + (1-eta_S) * ig(rac{\partial \mathcal{L}}{\partial \Theta}ig)^2 \ lpha' &= rac{1}{\sqrt{S_{(t)} + \epsilon}} * lpha \end{aligned}$$

#### Bias correction

You will have observed that we initialized to 0 the moving averages for gradients (  $V_0=0$ ) and the variance of the gradients ( $S_0=0$ ).

So the values are "biased" towards 0 with the bias having greatest effect for small t (i.e., when the number of "actual" values is small).

We can correct for the bias by dividing by  $(1 - \beta^t)$ :

$$egin{array}{lll} \hat{V} &=& rac{V_{(t)}}{1-eta_V^t} \ \hat{S} &=& rac{S_{(t)}}{1-eta_S^t} \end{array}$$

$$\hat{S} = rac{S_{(t)}}{1-eta_S^t}$$

```
In [30]: print("Done")
```

Done