# EE 046211 - Technion - Deep Learning

Tal Daniel (https://taldatech.guthub.io)

# **Tutorial 03 - Optimization & Gradient Descent Algorithms**



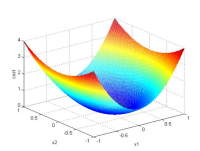
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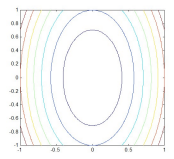
```
In [7]: # imports for the tutorial
   import numpy as np
   import pandas as pd
   import matplotlib.pyplot as plt
   from mpl_toolkits.mplot3d import Axes3D
   from sklearn.model_selection import train_test_split
   %matplotlib notebook
   # %matplotlib inline

# pytorch imports
   import torch
   import torch.nn as nn
   import torchvision.transforms as transforms
   import torchvision.datasets as dsets
```

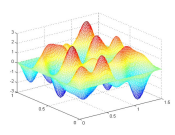
# Unimodal vs. Multimodal Optimization

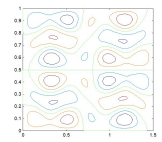
• Unimodal - only one optimum, that is, the local optimum is also global.





• Multimodal - more than one optimum.





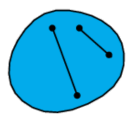
Most search schemes are based on the assumption of unimodal surface. The optimum determined in such cases is called local optimum design.

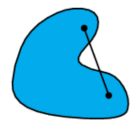
The **global optimum** is the best of all *local optimum* designs.



• Definition:

$$orall x_1, x_2 \in X, orall t \in [0,1]: \ f(tx_1 + (1-t)x_2) \leq tf(x_1) + (1-t)f(x_2)$$



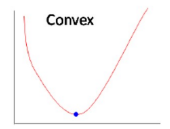


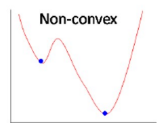
convex

concave

Image Source (http://mathworld.wolfram.com/Convex.html)

- Convex functions are unimodal.
  - However, unimodal functions are not always convex, but they are usually still easy to optimize.







# **Optimality Conditions**

- If  $f:\mathbb{R}^d o\mathbb{R}$  has *local* optimum at  $x_0$  then  $\nabla f(x_0)=0$ .

    $\nabla f(x_0)$  is also called **the gradient** at  $x_0$ .

   The Hessian Matrix :  $H(f)(x)_{i,j}=\frac{\partial^2}{\partial x_i\partial x_j}f(x)\in\mathbb{R}^{d\times d}$

$$H = egin{bmatrix} rac{\partial^2 f}{\partial x_1^2} & rac{\partial^2 f}{\partial x_1 \partial x_2} & \cdots & rac{\partial^2 f}{\partial x_1 \partial x_d} \ rac{\partial^2 f}{\partial x_2 x_1} & rac{\partial^2 f}{\partial x_2^2} & \cdots & rac{\partial^2 f}{\partial x_2 \partial x_d} \ dots & dots & dots & dots \ rac{\partial^2 f}{\partial x_d x_1} & rac{\partial^2 f}{\partial x_d \partial x_2} & \cdots & rac{\partial^2 f}{\partial x_d^2} \ \end{pmatrix}$$

- If the **Hessian** matrix is:
  - Positive Definite (all eigenvalues positive) at  $x_0 o \textit{local minimum}$ .
  - Negative Definite (all eigenvalues negative) at  $x_0 o \textit{local maximum}.$
  - Both **positive and negative** eigenvalues at  $x_0 o saddle$  point.



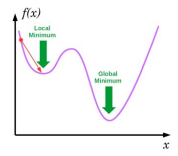




· Note: the Hessian matrix is symmetric if the second partial derivatives are continuous, but this is not always true (Schwarz's theorem (https://en.wikipedia.org/wiki/Symmetry of second derivatives)).

# (Batch) Gradient Descent

- · Generic optimization algorithm capable of finding optimal solutions to a wide range of problems.
- The general idea is to tweak parameters **iteratively** to minimize a cost function.
- It measures the local gradient of the error function with regards to the parameter vector (\(\theta\) or \(w\)), and it goes down in the direction of the descending gradient.
- · Once the gradient is zero the minimum is reached (=convergence).
- · Pseudocode:
  - Require: Learning rate  $\alpha_k$
  - ullet Require: Initial parameter vector w
  - While stopping criterion not met do
    - Compute gradient:  $g \leftarrow \nabla f(x, w)$
    - $\circ$  Apply update:  $w \leftarrow w lpha_k g$
    - $\bullet$   $k \leftarrow k+1$
  - end while



• **Convergene**: When the cost function is *convex* and its slope does not change abruptly, (Batch) GD with a (small enough) *fixed* learning rate will eventually converge to the optimal solution (but the time is depndent on the rate).

# 3

## Stochastic Gradient Descent (Mini-Batch Gradient Descent)

- The main problem with (Batch) GD is that it uses the **whole** training set to compute the gradients. But what if that training set is huge or each sample has a very large number of features? Computing the gradient can take a very long time.
- Stochastic Gradient Descent on the other hand, samples just one instance randomly at every step and computes the gradients based on that
  single instance (remember the Perceptron algorithm?). This makes the algorithm much faster but due to its randomness, it is much less stable.
  Instead of steady decreasing untill reaching the minimum, the cost function will bounce up and down, decreasing only on average. With time,
  it will get very close to the minimum, but once it is there it will continue to bounce around!
  - When we have outliers, SGD may take us to undesirable areas.
- The final parameters are good but not optimal.
- When the cost function is very irregular, this bouncing can actually help the algorithm escape local minima, so SGD has better chance to find the global minimum.
- How to find optimal parameters using SGD?
  - Reduce the learning rate gradually: this is called learning rate schedule
    - But don't reduce too quickly or you will get stuck at a local minimum or even frozen!
- Mini-Batch Gradient Descent same idea as SGD, but instead of one instance each step, m samples.
  - Get a little bit closer to the minimum than SGD but a little harder to escape local minima.

#### · Pseudocode:

- lacktriangle Require: Learning rate  $lpha_k$
- ullet Require: Initial parameter w
- While stopping criterion not met do
  - $\circ~$  Sample a minibatch of m examples from the training set (m=1 for SGD)
  - $\circ \ \mbox{ Set } \{x_1, \dots, x_m, \}$  with corresponding targets  $\{y_1, \dots, y_m\}$
  - Compute gradient:  $g \leftarrow \frac{1}{m} \sum_{i=1}^m f'(x_i, w, y_i)$  Apply update:  $w \leftarrow w \alpha_k g$

  - $\circ$   $k \leftarrow k+1$
- end while

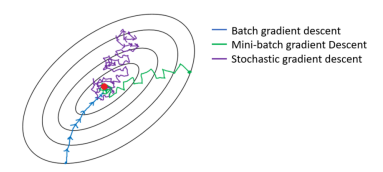


Image Source (https://towardsdatascience.com/gradient-descent-algorithm-and-its-variants-10f652806a3)

#### **GD Comparison Summary**

Method	Accuracy	Update Speed	Memory Usage	Online Learning
Batch Gradient Descent	Good	Slow	High	No
Stochastic Gradient Descent	Good (with softening)	Fast	Low	Yes
Mini-Batch Gradient Descent	Good	Medium	Medium	Yes (depends on the MB size)

- "Online" samples arrive while the algorithm runs (that is, when the algorithm starts running, not all samples exist)
- Note: all of the Gradient Descent algorithms require scaling if the feaures are not within the same range!

### **Update Speed**

- Assume: number of samples N=100 and we wish to train for 10 epochs (iterating over all of the samples 10 times).
- Denote batch\_size with m.
- batch\_size and num\_epochs are hyper-parameters: parameters that are chosen by the user prior to the training stage, they are not learned.

Method	# Gradients Updates (=iterations)
Batch Gradient Descent, $m=100$	1 * 10
Stochastic  Gradient Descent,  m=1	100*10 = 1000
Mini-Batch Gradient Descent, $m=10$	10*10=100

• The update speed affects how we tune the learning rate, which is also a hyper-parameter (usually, larger batches enable using higher learning rates).

### Challenges

- · Choosing a learning rate.
  - Defining learning schedule.
- Working with features of different scales (e.g. heights (cm), weights (kg) and age (scalar)).
- Avoiding local minima (or suboptimal minima).

# The Learning Rate

- Learning Rate hyperparameter it is the size of step to be taken in each iteration.
  - ullet Too  $\mathit{small} o \mathsf{the}$  algorithm will have to go through many iterations to converge, which will take a long time.
  - ullet Too  $\mathit{high} 
    ightarrow \mathsf{might}$  make the algorithm diverge as it may miss the minimum.

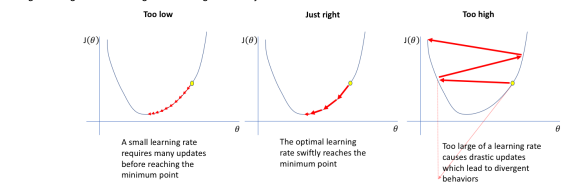


Image Source (https://www.jeremyjordan.me/nn-learning-rate/)



### **Example - (Multivariate) Linear Least Squares**

- · Problem Formulation
  - $ullet y \in \mathbb{R}^N$  vector of values
  - $oldsymbol{X} \in \mathbb{R}^{N imes L}$  data matrix with N examples and L features
  - $oldsymbol{w} \in \mathbb{R}^L$  the *parameters* to be learned, a **weight for each feature**
- Goal: find w that best fits the measurement y, that is, find a weighted linear combination of the feature vector to best fit the measurement y
- · Mathematiacally, the problem is:

$$\min_w f(w;x,y) = \min_w \sum_{i=1}^N \left|\left|x_i w - y_i
ight|
ight|^2$$

· In vector form:

$$\min_{w}f(w;x,y)=\min_{w}\left|\left|Xw-Y
ight|
ight|^{2}$$



### (Multivariate) LLS - Analytical Solution

· Mathematically:

$$\min_{w} f(w; x, y) = \min_{w} ||Xw - Y||^2 = \min_{w} (Xw - Y)^T (Xw - Y) = \min_{w} (w^T X^T Xw - 2w^T X^T Y + Y^T Y)$$

• The derivative:

$$abla_w f(w;x,y) = (X^TX + X^TX)w - 2X^TY = 0 
ightarrow w = (X^TX)^{-1}X^TY \ X^TX \in \mathbb{R}^{L imes L}$$

• Notice how the gradient is dependent on the features, which is why scaling them is important when applying gradient descent (however, scaling is not necessary if we use the closed-form solution, the least squares solution).

```
In [2]: # let's load the cancer dataset
dataset = pd.read_csv('./datasets/cancer_dataset.csv')
# print the number of rows in the data set
number_of_rows = len(dataset)
# reminder, the data looks like this
dataset.sample(10)
```

Out[2]:

	id	diagnosis	radius_mean	texture_mean	perimeter_mean	area_mean	smoothness_mean	compactness_mean	concavit
416	905978	В	9.405	21.70	59.60	271.2	0.10440	0.06159	О
315	894089	В	12.490	16.85	79.19	481.6	0.08511	0.03834	C
373	901288	М	20.640	17.35	134.80	1335.0	0.09446	0.10760	C
115	864685	В	11.930	21.53	76.53	438.6	0.09768	0.07849	C
379	9013838	М	11.080	18.83	73.30	361.6	0.12160	0.21540	C
296	891936	В	10.910	12.35	69.14	363.7	0.08518	0.04721	C
172	87164	М	15.460	11.89	102.50	736.9	0.12570	0.15550	С
117	864729	М	14.870	16.67	98.64	682.5	0.11620	0.16490	С
321	894618	М	20.160	19.66	131.10	1274.0	0.08020	0.08564	C
542	921644	В	14.740	25.42	94.70	668.6	0.08275	0.07214	О

10 rows × 33 columns

```
In [8]: def batch_generator(x, y, batch_size, shuffle=True):
            This function generates batches for a given dataset x.
            N, L = x.shape
            num_batches = N // batch_size
            batch_x = []
            batch_y = []
            if shuffle:
                # shuffle
                rand gen = np.random.RandomState(0)
                shuffled_indices = rand_gen.permutation(np.arange(N))
                x = x[shuffled_indices, :]
                y = y[shuffled_indices, :]
            for i in range(N):
                batch_x.append(x[i, :])
                batch_y.append(y[i, :])
                if len(batch_x) == batch_size:
                    yield np.array(batch_x).reshape(batch_size, L), np.array(batch_y).reshape(batch_size, 1)
                    batch_x = []
                    batch_y = []
            if batch_x:
                yield np.array(batch_x).reshape(-1, L), np.array(batch_y).reshape(-1, 1)
```

- Pseudocode for Linear Regression:
  - lacktriangle Require: Learning rate  $lpha_k$
  - ullet Require: Initial parameter w
  - While stopping criterion not met do
    - $\circ~$  Sample a minibatch of m examples from the training set ( $m=1~{
      m for}~{
      m SGD}$ )
    - $\circ \ \ \text{Set} \ \tilde{X} = [x_1, \dots, x_m] \ \ \text{with corresponding targets} \ \tilde{Y} = [y_1, \dots, y_m]$   $\circ \ \ \ \text{Compute gradient:} \ g \leftarrow 2\tilde{X}^T \tilde{X} w 2\tilde{X}^T \tilde{Y}$

    - $\circ \ \ \mathsf{Apply} \ \mathsf{update} \colon w \leftarrow w \alpha_k g$
    - $\circ$   $k \leftarrow k+1$
  - end while

```
In [9]: # multivaraite mini-batch gradient descent
          X = dataset[['radius_mean', 'area_mean']].values
          Y = dataset[['perimeter_mean']].values
          # Scaling
          X = (X - X.mean(axis=0, keepdims=True)) / X.std(axis=0, keepdims=True)

Y = (Y - Y.mean(axis=0, keepdims=True)) / Y.std(axis=0, keepdims=True)
          N = X.shape[0]
          batch_size = 10
          num_batches = N // batch_size
          print("total batches:", num_batches)
```

total batches: 56

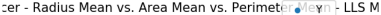
```
In [10]: num_iterations = 10
         alpha k = 0.001
         batch_gen = batch_generator(X, Y, batch_size, shuffle=True)
         # initialize w
         w = np.zeros((L, 1))
         for i in range(num_iterations):
             for batch_i, batch in enumerate(batch_gen):
                 batch_x, batch_y = batch
                 if batch_i % 50 == 0:
                     print("iter:", i, "batch:", batch_i, " w = ")
                     print(w)
                 gradient = 2 * batch_x.T @ batch_x @ w - 2 * batch_x.T @ batch_y
                 w = w - alpha_k * gradient
             batch_gen = batch_generator(X, Y, batch_size, shuffle=True)
         lls_sol = X @ w
         iter: 0 batch: 0 w =
         [[0.]
          [0.]]
         iter: 0 batch: 50 w =
         [[0.4391737]
          [0.41735388]]
         iter: 1 batch: 0 w =
         [[0.45621431]
          [0.43296123]]
         iter: 1 batch: 50 w =
         [[0.50458588]
          [0.46953377]]
         iter: 2 batch: 0 w =
         [[0.50659246]
          [0.46976394]]
         iter: 2 batch: 50 w =
         [[0.51664408]
          [0.46913792]]
         iter: 3 batch: 0 w =
         [[0.51715596]
          [0.46786206]]
         iter: 3 batch: 50 w =
         [[0.52339741]
          [0.46367398]]
         iter: 4 batch: 0 w =
         [[0.52374047]
          [0.4622501]]
         iter: 4 batch: 50 w =
         [[0.5295512 ]
          [0.45779193]]
         iter: 5 batch: 0 w =
         [[0.52985556]
          [0.456353 ]]
         iter: 5 batch: 50 w =
         [[0.53556725]
          [0.45194588]]
         iter: 6 batch: 0 w =
         [[0.53584598]
          [0.45050492]]
         iter: 6 batch: 50 w =
         [[0.54149192]
          [0.44617918]]
         iter: 7 batch: 0 w =
         [[0.54174661]
          [0.44473749]]
         iter: 7 batch: 50 w =
         [[0.54733087]
          [0.44049501]]
         iter: 8 batch: 0 w =
         [[0.54756198]
          [0.43905271]]
```

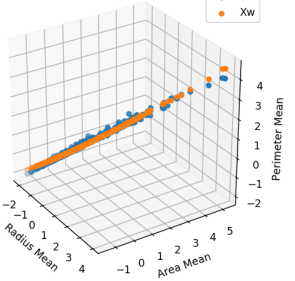
iter: 8 batch: 50 w =

[[0.55308576] [0.43489257]] iter: 9 batch: 0 w = [[0.55329364] [0.43344969]] iter: 9 batch: 50 w =

[[0.55875783] [0.42937075]]

```
In [11]: # plot
         plot 3d lls(X[:,0], X[:, 1], Y, lls sol, "Breast Cancer - Radius Mean vs. Area Mean vs. Perimeter Mean - L
         LS Mini-Batch GD")
         print("w:")
         print(w)
```





[[0.55894282] [0.42792729]]



## Learning Rate Scheduling (Annealing)

- When training deep networks, it is usually helpful to anneal (gradually change the rate) the learning rate over time.
  - Physics intuition: with a high learning rate, the system contains too much kinetic energy and the parameter vector bounces around chaotically, unable to settle down into deeper, but narrower parts of the loss function.
- . Knowing when to decay the learning rate can be tricky: decay it slowly and you'll be wasting computation bouncing around chaotically with little improvement for a long time. But decay it too aggressively and the system will cool too quickly, unable to reach the best position it can.
- There are three common types of implementing the learning rate decay: step deacy, exponential decay and 1/t decay.
  - Recently, cyclic learning schedulers, such as One-cycle learning rate scheduler (https://www.kaggle.com/residentmario/one-cycle-learning-<u>rate-schedulers</u>) or *cosine* scheduling, have been gaining popularity as well.
- Step decay: Reduce the learning rate by some factor every few epochs.
  - Typical values might be reducing the learning rate by a half every 5 epochs, or by 0.1 every 20 epochs. These numbers depend heavily on the type of problem and the model.
  - One heuristic you may see in practice is to watch the validation error while training with a fixed learning rate, and reduce the learning rate by a constant (e.g. 0.5) whenever the validation error stops improving.
- Exponential decay: has the mathematical form:

$$\alpha = \alpha_0 \exp(-kt),$$

where  $\alpha_0$ , k are hyperparameters and t is the iteration number (but you can also use units of epochs).

- $\alpha_0$  is the initial learning rate.
- k is also referred to as the gamma  $(\gamma)$  hyperparameter.

• 1/t decay: has the mathematical form:

$$\alpha = \frac{\alpha_0}{1 + kt},$$

where  $\alpha_0, k$  are hyperparameters and t is the iteration number.

· Cosine annealing: has the mathematical form:

$$lpha = lpha_{min} + rac{1}{2}(lpha_0 - lpha_{min}) \left(1 + \cos\!\left(rac{t}{t_{max}}\pi
ight)
ight),$$

where  $lpha_{min}$  is the minimum learning rate (deafult is 0) and  $t_{max}$  is number of iterations to perform a cycle.

- In practice, we find that the step decay is slightly preferable because the hyperparameters it involves (the fraction of decay and the step timings in units of epochs) are more interpretable than the hyperparameter k.
- · Lastly, if you can afford the computational budget, you can try a slower decay and train for a longer time.



# Learning Rate Scheduling in PyTorch

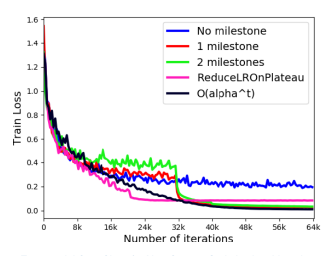
- We will use learning rate scheduling to train the neural network models later in the course.
- PyTorch offers several schedulers which can be found <a href="https://pytorch.org/docs/stable/optim.html#how-to-adjust-learning-rate">https://pytorch.org/docs/stable/optim.html#how-to-adjust-learning-rate</a>).
- A typical workflow with schedulers (learning rate scheduling should be applied after optimizer's update):

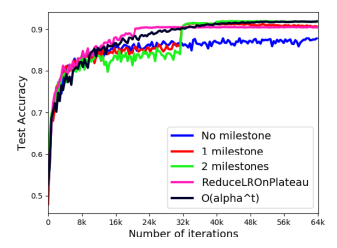
```
scheduler = ...
for epoch in range(100):
    train(...)
    validate(...)
    scheduler.step()
```

- torch.optim.lr\_scheduler.StepLR
- torch.optim.lr\_scheduler.MultiStepLR
- $\bullet \quad \texttt{torch.optim.lr\_scheduler.ExponentialLR}$
- torch.optim.lr\_scheduler.CosineAnnealingLR
- And more...

### Reducing LR on Plateau

- Reduce learning rate when a metric has stopped improving. Usually the validation accuracy.
- Models often benefit from reducing the learning rate by a factor of 2-10 once learning does not improve.
- This scheduler reads a metrics quantity and if no improvement is seen for a patience number of epochs, the learning rate is reduced.
- In PyTorch: torch.optim.lr\_scheduler.ReduceLROnPlateau





- Exponential Step Sizes for Non-Convex Optimization, Li et al. 2020 (https://www.researchgate.net/publication/339252130 Exponential Step Sizes for Non-Convex Optimization).
- · Plots of the train loss and test accuracy for training a 20-layer Residual Network to do image classification on CIFAR-10.
- · The number of milestones in the legend denotes how many times we can choose to decrease the step size during training.

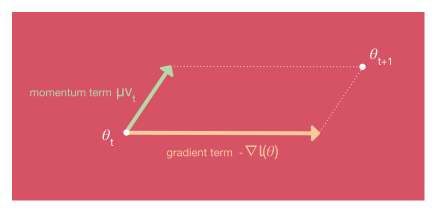


### **Momentum & Nesterov Momentum**

- Gradient descent is simple and has many virtues, but **speed** is not one of them.
- For a step-size small enough, gradient descent makes a monotonic improvement at every iteration. It should always converge (sometimes to a local minimum).
- Momentum update is another optimization approach that almost always enjoys better convergence rates in deep networks.
  - It can be seen as a "global" (equally for all parameters) adaptive learning rate.
- This update can be motivated from a physical perspective of the optimization problem. In particular, the loss can be interpreted as the height of a *hilly terrain*.
  - Initializing the parameters with random numbers is equivalent to setting a particle with zero initial velocity at some location. The optimization process can then be seen as equivalent to the process of simulating the parameter vector (i.e. a particle) as rolling on the landscape.
  - Since the force on the particle is related to the gradient of potential energy (i.e.  $F = -\nabla U$ ), the force felt by the particle is precisely the (negative) gradient of the loss function.
  - ullet Moreover, F=ma so the (negative) gradient is in this view proportional to the acceleration of the particle.
  - The physics view suggests an update in which the gradient only directly influences the velocity (and maintains information about the acceleration), which in turn has an effect on the position.
- Momentum proposes the following tweak to gradient descent, giving gradient descent a short-term memory:

$$z^{k+1} = eta z_k - lpha 
abla f(w^k) \ w^{k+1} = w^k + z^{k+1}$$

- lacksquare  $\alpha$  is the learning rate.
- When  $\beta=0$  , we recover gradient descent. But for  $\beta=0.99$  (sometimes 0.999, if things are really bad), this appears to be the boost we need. Our iterations regain that speed and boldness it lost, speeding to the optimum with a renewed energy.
- $\beta$  is a variable that is sometimes called *momentum*.
- Effectively, this variable damps the velocity and reduces the kinetic energy of the system, or otherwise the particle would never come to a stop at the bottom of a hill.
- · With Momentum update, the parameter vector will build up velocity in any direction that has consistent gradient.
- Momentum Demo (https://distill.pub/2017/momentum/)
- Note: Momentum usually works in larger batches and may break in smaller batches.

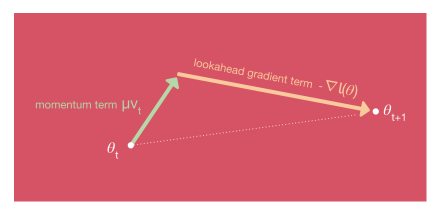


• Image Source (https://dominikschmidt.xyz/nesterov-momentum)

#### **Nesterov Momentum**

- · Nesterov Momentum is a slightly different version of the momentum update that has recently been gaining popularity.
- It enjoys stronger theoretical convergence guarantees for **convex functions** and in practice it also consistenly works slightly better than standard momentum.
- The core idea behind Nesterov momentum is that when the current parameter vector is at some position x, then looking at the momentum update above, we know that the momentum term alone (i.e. ignoring the second term with the gradient) is about to nudge the parameter vector by  $\beta * z_k$ .
- Therefore, if we are about to compute the gradient, we can treat the future approximate position  $x + \beta * z_k$  as a "lookahead" this is a point in the vicinity of where we are soon going to end up.
- Hence, it makes sense to compute the **gradient** at  $x + \beta * z_k$  instead of at the "old/stale" position x, since while the gradient term always points in the right direction, the momentum term may not. If the momentum term points in the wrong direction or overshoots, the gradient can still "go back" and correct it in the same update step.
- Nesterov Momentum:

$$z^{k+1} = eta z^k - lpha 
abla f(w^k + eta z^k) \ w^{k+1} = w^k + z^{k+1}$$



• Image Source (https://dominikschmidt.xyz/nesterov-momentum)



## **Momentum in PyTorch**

• torch.optim.SGD(model.parameters(), lr=learning\_rate, momentum=0.9, nesterov=True)

```
In [ ]: # simple optimizer and lr scheduling example
        # courtesy of: deeplearningwizard.com/deep learning/boosting models pytorch/lr scheduling/
        from torch.optim.lr_scheduler import ReduceLROnPlateau
        STEP 1: LOADING DATASET
        train_dataset = dsets.MNIST(root='./data',
                                    train=True,
                                     transform=transforms.ToTensor(),
                                    download=True)
        test_dataset = dsets.MNIST(root='./data',
                                    train=False,
                                   transform=transforms.ToTensor())
        STEP 2: MAKING DATASET ITERABLE
        batch_size = 100
        n_{iters} = 6000
        num_epochs = n_iters / (len(train_dataset) / batch_size)
        num_epochs = int(num_epochs)
        train_loader = torch.utils.data.DataLoader(dataset=train_dataset,
                                                    batch_size=batch_size,
                                                    shuffle=True)
        test_loader = torch.utils.data.DataLoader(dataset=test_dataset,
                                                   batch_size=batch_size,
                                                   shuffle=False)
        STEP 3: CREATE MODEL CLASS
        class SimpleModel(nn.Module):
                  _init__(self, input_dim, hidden_dim, output_dim):
                super(SimpleModel, self).__init__()
                # Linear function
                self.fc1 = nn.Linear(input_dim, hidden_dim)
                # Non-linearity
                self.relu = nn.ReLU()
                # Linear function (readout)
                self.fc2 = nn.Linear(hidden_dim, output_dim)
            def forward(self, x):
                # Linear function
                out = self.fc1(x)
                # Non-linearity
                out = self.relu(out)
                # Linear function (readout)
                out = self.fc2(out)
                return out
        ...
        STEP 4: INSTANTIATE MODEL CLASS AND DEVICE
        input_dim = 28 * 28
        hidden dim = 100
        output_dim = 10
        device = torch.device("cuda:0" if torch.cuda.is_available() else "cpu")
        model = SimpleModel(input_dim, hidden_dim, output_dim).to(device)
        ...
        STEP 5: INSTANTIATE LOSS CLASS
        criterion = nn.CrossEntropyLoss()
        STEP 6: INSTANTIATE OPTIMIZER CLASS
        learning rate = 0.1
        optimizer = torch.optim.SGD(model.parameters(), lr=learning_rate, momentum=0.9, nesterov=True)
        STEP 7: INSTANTIATE STEP LEARNING SCHEDULER CLASS
```

```
# lr = lr * factor
# mode='max': Look for the maximum validation accuracy to track
# patience: number of epochs - 1 where loss plateaus before decreasing LR
        # patience = 0, after 1 bad epoch, reduce LR
# factor = decaying factor
scheduler = ReduceLROnPlateau(optimizer, mode='max', factor=0.1, patience=0, verbose=True)
STEP 7: TRAIN THE MODEL
iter = 0
for epoch in range(num_epochs):
    for i, (images, labels) in enumerate(train_loader):
        # Send images and labels to device
        images = images.view(-1, 28 * 28).to(device)
        labeles = labels.to(device)
        # Forward pass to get output/logits
        outputs = model(images)
        # Calculate Loss: softmax --> cross entropy loss
        loss = criterion(outputs, labels)
        # Clear gradients w.r.t. parameters
        optimizer.zero_grad()
        # Getting gradients w.r.t. parameters
        loss.backward()
        # Updating parameters
        optimizer.step()
        iter += 1
        if iter % 500 == 0:
            # Calculate Accuracy
            correct = 0
            total = 0
            # Iterate through test dataset
            for images, labels in test_loader:
                # Send images and labels to device
                images = images.view(-1, 28 * 28).to(device)
                labeles = labels.to(device)
                # Forward pass only to get logits/output
                outputs = model(images)
                # Get predictions from the maximum value
                _, predicted = torch.max(outputs.data, 1)
                # Total number of labels
                total += labels.size(0)
                # Total correct predictions
                # Without .item(), it is a uint8 tensor which will not work when you pass this number to t
he scheduler
                correct += (predicted == labels).sum().item()
            accuracy = 100 * correct / total
            # Print Loss
            # print('Iteration: {}. Loss: {}. Accuracy: {}'.format(iter, Loss.data[0], accuracy))
    # Decay Learning Rate, pass validation accuracy for tracking at every epoch
    print('Epoch {} completed'.format(epoch))
    print('Loss: {}. Accuracy: {}'.format(loss.item(), accuracy))
    print('-' * 20)
    scheduler.step(accuracy) # accuracy is used to track down a plateau
```



## **Adaptive Learning Rate Methods**

Adapative learning methods compute individual learning rates for different parameters. Previously, we performed an update for all parameters w (or  $\theta$ ) at once as every parameter  $w_i$  used the same learning rate  $\alpha$ .

Popular algorithms include: AdaGrad, Rprop, RMSprop, Adam and more...



### **Adagrad**

- · Adagrad: one of the first adaptive learning rate algorithm, with the basic idea of adapting the learning rate to the parameters by performing smaller updates (i.e. low learning rates) for parameters associated with frequently occurring features, and larger updates (i.e. high learning rates) for parameters associated with infrequent features.
  - For this reason, it works well with sparse data.
- Adagrad uses a different learning rate for every parameter  $\boldsymbol{w}_i$  at every time step k
- We denote:
  - lacksquare  $\alpha$  the learning rate.
  - $g_k = 
    abla f(w^k)$ , the gradient at time step k, and  $g_{i,k}$  the partial derivative w.r.t. the parameter  $w_i$  at time step k.
  - $m{G}_k \in \mathbb{R}^{d imes d}$  a diagonal matrix, where each element  $G_{i,i}^k$  is the sum of squares of the gradients w.r.t  $w_i$  up to time step k,
  - $\epsilon$ , a "smoothing" term that prevents division by zero, deafult is  $10^{-8}$ , but can range from  $10^{-4}$  to  $10^{-8}$ .
- · The Adagrad update rule:

$$w_i^{k+1} = w_i^k - rac{lpha}{\sqrt{G_{i,i}^k + \epsilon}} \cdot g_{i,k}$$

- · Interestingly, without the square root operation, the algorithm performs much worse.

In vectorized form, we use the matrix-vector product 
$$\odot$$
 between  $G_k$  and  $g_k$ : 
$$w^{k+1} = w^k - \frac{\alpha}{\sqrt{G^k + \epsilon}} \odot g_k$$

- · Adagrad eliminates the need to manually tune the learning rate, which is nice.
- Most implementations use a default value of 0.01 for the learning rate.
- However, its main weakness is the accumulation of the squared gradients (a positive quantity) in the denominator which keeps growing during training and causes the learning rate to shrink and eventually become very small, at which point the algorithm doesn't acquire additional knowledge.



# Adagrad in PyTorch

• torch.optim.Adagrad(model.parameters(), lr=learning\_rate, initial\_accumulator\_value=0, eps=1e-10)



- RMSprop: an unpublished (no official paper) optimization algorithm designed for neural networks, first proposed by Geoffrey Hinton in <u>lecture 6</u> (<u>slide 29</u>) (<a href="https://www.cs.toronto.edu/~tijmen/csc321/slides/lecture\_slides\_lec6.pdf">https://www.cs.toronto.edu/~tijmen/csc321/slides/lecture\_slides\_lec6.pdf</a>) of the online course "Neural Networks for Machine Learning".
- The RMSProp update adjusts the Adagrad method in a very simple way in an attempt to reduce its aggressive, monotonically decreasing learning rate.
- · In particular, it uses a moving average of squared gradients instead.
- · We denote:
  - ullet  $\alpha$  the learning rate.
  - $ullet g_k = 
    abla f(w^k)$
  - ullet  $\mathbb{E}[g^2]$  moving average of squared gradients (stored in a cache with squared gradients from previous iterations).
  - $\beta$  moving average parameter (good default value 0.9).
- The RMSprop update rule:

$$egin{aligned} \mathbb{E}[g^2]_{k+1} &= eta \mathbb{E}[g^2]_k + (1-eta)g_k^2 \ w^{k+1} &= w^k - \dfrac{lpha}{\sqrt{\mathbb{E}[g^2]_{k+1}}} 
abla f(w^k) \end{aligned}$$

• The learning rate is adapted by dividing by the root of squared gradient, but since we only have the estimate of the gradient on the current minibatch, we need instead to use the moving average of it.



# **RMSprop in PyTorch**

- torch.optim.RMSprop(model.parameters(), lr=learning\_rate, alpha=0.99)
  - ullet alpha is eta from the equations above, the moving average parameter.



### **Adam - Adaptive Moment Estimation**

- · Adam: another optimization method that computes adaptive learning rates for each parameter.
- Adam combines the advantages of Adagrad, RMSprop and Momentum: It uses the squared gradients to scale the learning rate like RMSprop and it takes advantage of momentum by using moving average of the gradient instead of the gradient itself like SGD with momentum.
- In addition to storing an exponentially decaying average of past squared gradients like Adadelta and RMSprop, Adam also keeps an
  exponentially decaying average of past gradients similar to momentum.
  - Whereas momentum can be seen as a ball running down a slope, Adam behaves like a heavy ball with friction, and thus prefers flat minima in the error surface.
- We denote:
  - $\alpha$  the learning rate.
  - *m* moving average of gradients. Estimates the first moment (mean) of the gardients.
  - v moving average of squared gradients. Estimates the second momemnt (variance) of the gradients.
  - $\beta_1$  moving average parameter for m (default: 0.9).
  - $\beta_2$  moving average parameter for v (default: 0.999).

• The Adam update rule:

$$\mathbb{E}[g]_{k+1} = m_{k+1} = \beta_1 m_k + (1 - \beta_1) \nabla f(w^k) = \beta_1 m_k + (1 - \beta_1) g_k \\ \mathbb{E}[g^2]_{k+1} = v_{k+1} = \beta_2 v_k + (1 - \beta_2) (\nabla f(w^k))^2 = \beta_2 v_k + (1 - \beta_2) g_k^2$$

Then, we use an **unbiased** estimation:

$$\hat{m}_{k+1} = rac{m_{k+1}}{1-eta_1^{k+1}} \ \hat{v}_{k+1} = rac{v_{k+1}}{1-eta_2^{k+1}}$$

(the  $\beta$ 's are taken with the power of the current iteration)

$$w^{k+1} = w^k - rac{lpha}{\sqrt{\hat{v_k}} + \epsilon} \hat{m_k}$$

 $w^{k+1}=w^k-\frac{\alpha}{\sqrt{\hat{v_k}}+\epsilon}\hat{m_k}$  •  $\epsilon$ , a "smoothing" term that prevents diviosn by zero, deafult's is  $10^{-8}$ , but can range from  $10^{-4}$  to  $10^{-8}$ .

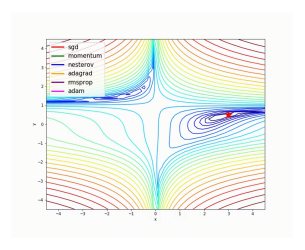


# Adam in PyTorch

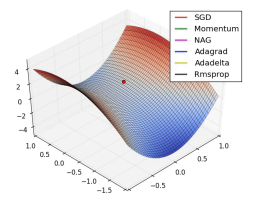
• torch.optim.Adam(model.parameters(), lr=learning\_rate, betas=(0.9, 0.999))



# 🕻 Comparison Between Methods



- · Contours of a loss surface and time evolution of different optimization algorithms.
- Notice the "overshooting" behavior of momentum-based methods, which makes the optimization look like a ball rolling down the hill.
- Image Source (https://github.com/ilguyi/optimizers.numpy)



- A visualization of a saddle point in the optimization landscape, where the curvature along different dimension has different signs (one
  dimension curves up and another down).
- Notice that SGD has a very hard time breaking symmetry and gets *stuck* on the top.
- Conversely, algorithms such as RMSprop will see very low gradients in the saddle direction. Due to the denominator term in the RMSprop
  update, this will increase the effective learning rate along this direction, helping RMSProp proceed.
- Image credit: <u>Alec Radford (https://twitter.com/alecrad)</u>



#### **Recommended Videos**



- These videos do not replace the lectures and tutorials.
- · Please use these to get a better understanding of the material, and not as an alternative to the written material.

### Video By Subject

- Gradient Descent Gradient Descent, Step-by-Step (https://www.youtube.com/watch?v=sDv4f4s2SB8)
  - Mathematics of Gradient Descent Intelligence and Learning (https://www.youtube.com/watch?v=jc2lthslyzM)
- Stochastic Gradient Descent Stochastic Gradient Descent, Clearly Explained (https://www.youtube.com/watch?v=vMh0zPT0tLl)
- Momentum Gradient Descent With Momentum (C2W2L06) (https://www.youtube.com/watch?v=k8fTYJPd3\_l)
- RMSProp RMSProp (C2W2L07) (https://www.youtube.com/watch?v=\_e-LFe\_igno)
- Adam Adam Optimization Algorithm (C2W2L08) (https://www.youtube.com/watch?v=JXQT\_vxqwls)
- Learning Rate Decay Learning Rate Decay (C2W2L09) (https://www.youtube.com/watch?v=QzulmoOg2JE)
- Momentum, Adagrad, RMSProp, Adam <u>UC Berkeley, STAT 157 Momentum, Adagrad, RMSProp, Adam (https://www.youtube.com/watch?v=gmwxUy7NYpA)</u>



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- Datasets from Kaggle (https://www.kaggle.com/) https://www.kaggle.com/ (https://www.kaggle.com/)
- Examples and code snippets were taken from "Hands-On Machine Learning with Scikit-Learn and TensorFlow"
   (<a href="http://shop.oreilly.com/product/0636920052289.do">http://shop.oreilly.com/product/0636920052289.do</a>)
- CS231n: Convolutional Neural Networks for Visual Recognition (https://cs231n.github.io/neural-networks-3/)
- <u>Deep Learning Wizard -Learning Rate Scheduling</u>
   (<a href="https://www.deeplearningwizard.com/deep\_learning/boosting\_models\_pytorch/lr\_scheduling/">https://www.deeplearningwizard.com/deep\_learning/boosting\_models\_pytorch/lr\_scheduling/</a>)
- Understanding Nesterov Momentum (NAG) (https://dominikschmidt.xyz/nesterov-momentum)
- · Sebastian Ruder An overview of gradient descent optimization algorithms (https://ruder.io/optimizing-gradient-descent/)