Optimization

Brooks Paige

Week 2

Overview

- Gradient Descent
 - ► Convergence analysis.
 - ► What can go wrong?
 - ▶ Batch and stochastic gradient descent.
- Momentum and other add-ons
 - ► Nesterov, Adagrad, RMSprop, Adam...
- Second-order methods

The need for optimization

Machine learning often requires fitting a model to data. Often this means finding the parameters θ of the model that 'best' fit the data, i.e. given some data-dependent loss $\mathcal{L}(\theta)$, we want

$$\underset{\boldsymbol{\theta}}{\arg\min} \, \mathcal{L}(\boldsymbol{\theta}).$$

In all but very simple cases, it is extremely difficult to find an algorithm that will guarantee to find the optimal θ .

How can we minimize a function f?

... particularly if we don't know much about it?

The trick is to use a Taylor expansion. For a small value δ , we can approximate $f(\mathbf{x} + \boldsymbol{\delta})$ as

$$f(\mathbf{x} + \boldsymbol{\delta}) \approx f(\mathbf{x}) + \boldsymbol{\delta}^{\top} \nabla f(\mathbf{x}).$$

This is a **first-order** approximation, which uses the function value and its gradient at x.

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$$f(\mathbf{x} + \boldsymbol{\delta}) \approx f(\mathbf{x}) + \boldsymbol{\delta}^{\top} \nabla f(\mathbf{x}) + \frac{1}{2} \boldsymbol{\delta}^{\top} H(\mathbf{x}) \boldsymbol{\delta},$$

where $H(\mathbf{x}) = \nabla^2 f(\mathbf{x})$ is the Hessian of f, is more accurate but also more expensive to compute.

First order methods

Gradient descent

We wish to find x that minimizes f(x). For general f there is no closed-form solution to this problem and we typically resort to iterative methods.

Gradient descent is a first-order method; it only uses the gradient. We assume our function is roughly linear around a starting point x, and

$$f(\mathbf{x} + \mathbf{s}) \approx f(\mathbf{x}) + \mathbf{s}^{\top} \nabla f(\mathbf{x}).$$

Our goal is to choose a good step direction s.

Gradient descent

As part of an iterative algorithm, for $\mathbf{x}_{t+1} \approx \mathbf{x}_t$, we have

$$f(\mathbf{x}_{t+1}) \approx f(\mathbf{x}_t) + \underbrace{(\mathbf{x}_{t+1} - \mathbf{x}_t)^{\top}}_{\mathbf{s}} \nabla f(\mathbf{x}_t).$$

The simplest steepest descent approach is to choose a step direction s (or equivalently, a next location x_{t+1}), as

$$\mathbf{s} \equiv \mathbf{x}_{t+1} - \mathbf{x}_t = -\epsilon \nabla f(\mathbf{x}_t)$$

for small **learning rate** ϵ .

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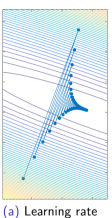
$$\mathbf{s} \equiv \mathbf{x}_{t+1} - \mathbf{x}_t = -\epsilon \nabla f(\mathbf{x}_t)$$

for small **learning rate** ϵ . Assuming our approximation holds, this decreases f, since

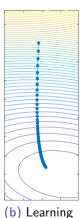
$$f(\mathbf{x}_{t+1}) \approx f(\mathbf{x}_t) - \underbrace{\epsilon \nabla f(\mathbf{x}_t)^{\top} \nabla f(\mathbf{x}_t)}_{>0} \leq f(\mathbf{x}_t).$$

Learning rates for gradient descent

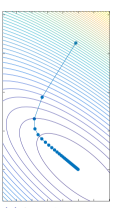
Note the difference between converging to the minimal function value, and the optimum parameters; we might have almost converged to the minimal value but still be a long way from the optimum parameters.



(a) Learning rate too large



(b) Learning rate too small

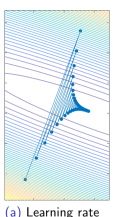


(c) Learning rate OK

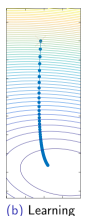
Learning rates for gradient descent

Common to consider adapting the learning rate.

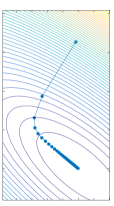
This is usually determined by experimenting with different values or learning schedules.



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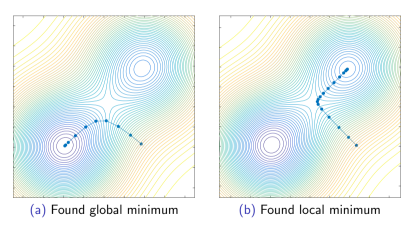


(b) Learni rate too small



(c) Learning rate OK

Gradient descent: local minima



For non-convex functions, depending on the initial point, even for the same function and small learning rate, we can converge to different solutions.

Stochastic gradient descent

• In machine learning, the loss function is often the sum (average) of losses for individual examples:

$$L(\boldsymbol{\theta}) = \frac{1}{N} \sum_{i=1}^{N} L_i(\boldsymbol{\theta}) = \frac{1}{N} \sum_{i=1}^{N} \ell(f(x_i | \boldsymbol{\theta}), y_i)$$

- If the dataset is big, one step of gradient descent could be very expensive.
- Instead, we can use stochastic gradient descent, iterating over the training set examples:

$$\boldsymbol{\theta}_{t+1} = \boldsymbol{\theta}_t - \epsilon \nabla L_i(\boldsymbol{\theta})$$

We can make progress even before we see all the data!

Mini-batch stochastic gradient descent

- The problem: gradient estimates are noisy
- The compromise: use mini-batches of training examples and compute the gradient estimate based on those. For $j=1,\ldots,M$ subsampled from $i=1,\ldots,N$.

$$\nabla_{\boldsymbol{\theta}} L(\boldsymbol{\theta}) = \frac{1}{N} \sum_{i=1}^{N} \nabla_{\boldsymbol{\theta}} L_i(\boldsymbol{\theta}) = \frac{1}{M} \sum_{i=1}^{M} \nabla_{\boldsymbol{\theta}} L_j(\boldsymbol{\theta}).$$

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- How should we choose the batch size?
 - ► Large batch size: less noisy estimates, convergence might be faster (in number of iterations)
 - ► Smaller batch size: faster parameter updates.
 - ► GPUs usually allow for larger batch sizes.

Fluctuations

- Stochastic gradient descent introduces fluctuations in the training process.
 The gradient estimate is noisy, so each update does not necessarily point toward the direction of true gradient.
- As such, it may even increase the loss.
- Reducing learning rate reduces fluctuations (but may slow down convergence).

Momentum

One simple idea to limit the zig-zag behavior is to make an update in the average direction of the previous updates.

Moving average: Consider a set of numbers x_1, \ldots, x_t . Then the average a_t is given by

$$a_t = \frac{1}{t} \sum_{\tau=1}^{t} x_{\tau} = \frac{1}{t} (x_t + (t-1)a_{t-1}) = \epsilon_t x_t + \mu_t a_{t-1}$$

for suitably chosen ϵ_t and $0 \le \mu_t \le 1$. If μ_t is small then the more recent x contribute more strongly to the moving average.

Momentum

The idea of momentum is to use a form of moving average to the updates:

$$\tilde{\mathbf{g}}_{t+1} = \mu_t \tilde{\mathbf{g}}_t - \epsilon \nabla f(\mathbf{x}_t)$$

$$\mathbf{x}_{t+1} = \mathbf{x}_t + \tilde{\mathbf{g}}_{t+1}$$

Hence, instead of using the update $-\epsilon \nabla f(\mathbf{x})$, we use the moving average of the update $\tilde{\mathbf{g}}_{t+1}$ to form the new update.

Momentum

- Momentum can increase the speed of convergence since, for smooth objectives, as we get close to the minimum the gradient decreases and standard gradient descent starts to slow down.
- If the learning rate is too large, standard gradient descent may oscillate, but momentum may reduce oscillations by going in the average direction.
- However, the momentum parameter μ may need to be reduced with the iteration count to ensure convergence.
- Particularly useful when the gradient is noisy. By averaging over previous gradients, the noise 'averages' out and the moving average direction can be much less noisy.
- Momentum is also useful to avoid saddles (a point where the gradient is zero, but the objective function is not a minimum, such as the function x^3 at the origin) since typically the momentum will carry you over the saddle.

Nesterov's Accelerated Gradient

This looks similar to momentum but has a slightly different update

$$\tilde{\mathbf{g}}_{t+1} = \mu_t \tilde{\mathbf{g}}_t - \epsilon \nabla f(\mathbf{x}_t + \mu_t \tilde{\mathbf{g}}_t)
\mathbf{x}_{t+1} = \mathbf{x}_t + \tilde{\mathbf{g}}_{t+1}$$

That is, we use the gradient of the point we will move to, rather than the current point.

- Need to choose a schedule for μ_t . Nesterov suggests $\mu_t = 1 3/(t+5)$.
- For convex (and not stochastic) functions, has rate of convergence to the optimum

$$f(\mathbf{x}_t) - f^* \le \frac{c}{t^2}$$

for some constant c, compared to 1/t convergence for gradient descent.

Let's imagine that we've arrived at our current parameter \mathbf{x}_t based on an update

$$\mathbf{x}_t = \mathbf{x}_{t-1} + \mathbf{s}_{t-1}.$$

We now (retrospectively) ask: 'What would have been a better update than the one we actually made?'. Well, using the update we arrived at a function value

$$f(\mathbf{x}_t) = f(\mathbf{x}_{t-1} + \mathbf{s}_{t-1}).$$

We could have got to a better value if we had gone to $\mathbf{x}_t' = \mathbf{x}_t - \epsilon \nabla f(\mathbf{x}_t)$ instead, with a step direction

$$\mathbf{s}_t = \mathbf{x}_t' - \mathbf{x}_{t-1}$$

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Hence, we define

$$\mathbf{s}_t = \mathbf{s}_{t-1} - \epsilon \nabla f(\mathbf{x}_{t-1} + \mathbf{s}_{t-1})$$

and make the update

$$\mathbf{x}_t = \mathbf{x}_{t-1} + \mathbf{s}_t.$$

(Note the difference from the original update $\mathbf{x}_t = \mathbf{x}_{t-1} + \mathbf{s}_{t-1}!$)

• The basic update

$$\mathbf{s}_t = \mathbf{s}_{t-1} - \epsilon \nabla f(\mathbf{x}_{t-1} + \mathbf{s}_{t-1})$$

is a form of momentum that pushes x along the direction it was previously going.

- Even when we reach a point where the gradient is zero, then s_t will be the same as s_{t-1} .
- We therefore introduce a term μ to dampen oscillations and ensure convergence, giving the standard Nesterov update

$$\mathbf{s}_{t} = \mu_{t-1}\mathbf{s}_{t-1} - \epsilon \nabla f(\mathbf{x}_{t-1} + \mu_{t-1}\mathbf{s}_{t-1})$$
$$\mathbf{x}_{t} = \mathbf{x}_{t-1} + \mathbf{s}_{t}$$

Since $\mu < 1$, ${\bf s}$ will quickly converge to zero around the minimum.

Adaptive learning rates

Adagrad

Suppose that your data is sparse: some features are rare (and thus relevant weights rarely get updates). Using the same learning rate for all of them seems detrimental. Adagrad fixes that:

$$g_{t,i} = (\nabla L(\boldsymbol{\theta}^t))_i \qquad G_{t,ii} = \sum_{\tau=1}^t g_{\tau,i}^2$$

$$\theta_i^{t+1} = \theta_i^t - \frac{\eta}{\sqrt{G_{t,ii} + \varepsilon}} g_{t,i}$$

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- + Adaptive learning rates. Using a default choice of $\eta=1.0$ or maybe $\eta=0.1$ works pretty often.
- The accumulated gradients eventually make learning very slow and the learning rate never recovers.

RMSprop

RMSprop fixes the problem with Adagrad vanishing learning rates by using moving average of past squared gradients instead (like momentum is doing for the gradients themselves):

$$s_{t,i} = (1 - \gamma)s_{t-1,i} + \gamma g_{t,i}^2$$
$$\theta_i^{t+1} = \theta_i^t - \frac{\eta}{\sqrt{s_{t,i} + \varepsilon}} g_{t,i}$$

The algorithm was first presented in Geoff Hinton Coursera course (Lecture 6e).

Adam

Adam is a very popular optimizer, basically combining RMSprop ideas with momentum.

$$m_{t,i} = (1 - \beta)m_{t-1,i} + \beta g_{t,i}$$

$$s_{t,i} = (1 - \gamma)s_{t-1,i} + \gamma g_{t,i}^2$$

$$\theta_i^{t+1} = \theta_i^t - \frac{\eta}{\sqrt{s_{t,i} + \varepsilon}} m_{t,i}$$

All of those algorithms are implemented in the popular machine learning libraries. A bunch of tweaks appeared since (AdamW, NAdam, Amsgrad, ...).

Higher order methods

Second-order methods

Instead of a first-order Taylor expansion, we can use a second-order one, incorporating the Hessian. This is like assuming the function is roughly quadratic around the current $\mathbf{x} \in \mathbb{R}^D$, with

$$f(\mathbf{x} + \mathbf{s}) \approx f(\mathbf{x}) + \mathbf{s}^{\top} \nabla f(\mathbf{x}) + \frac{1}{2} \mathbf{s}^{\top} \mathbf{H}(\mathbf{x}) \mathbf{s}$$

where $\mathbf{H}(\mathbf{x})$ is the matrix of all partial derivatives of $f(\mathbf{x})$, i.e.

$$\mathbf{H}(\mathbf{x}) = \left[egin{array}{cccc} rac{\partial^2 f}{\partial x_1^2} & \cdots & rac{\partial^2 f}{\partial x_1 \partial x_D} \ dots & \ddots & dots \ rac{\partial^2 f}{\partial x_D \partial x_1} & \cdots & rac{\partial^2 f}{\partial x_2^2} \end{array}
ight].$$

Newton's method

For **Newton's method**, we consider

$$f(\mathbf{x} + \mathbf{s}) \approx f(\mathbf{x}) + \mathbf{s}^{\top} \nabla f(\mathbf{x}) + \frac{1}{2} \mathbf{s}^{\top} \mathbf{H}(\mathbf{x}) \mathbf{s}$$

and find the step direction by minimizing it with respect to s. Differentiating the right hand side yields

$$\nabla_{\mathbf{s}} f(\mathbf{x} + \mathbf{s}) \approx \nabla f(\mathbf{x}) + \mathbf{H}(\mathbf{x})\mathbf{s};$$

setting equal to zero and solving for s, we find the minimum to be at

$$\mathbf{s} = -\mathbf{H}(\mathbf{x})^{-1} \nabla f(\mathbf{x}).$$

Often, a learning rate $\epsilon \in (0,1)$ is included to dampen the update,

$$\mathbf{x}_{t+1} = \mathbf{x}_t - \epsilon \mathbf{H}^{-1} \nabla f(\mathbf{x}_t).$$

Newton's method

When we ran gradient descent, we made a linear approximation to $f(\mathbf{x})$ and then stepped a short distance ϵ .

Here, we instead make a quadratic approximation to $f(\mathbf{x})$, and move *directly to its minimum*. The rough idea (Newton's method):

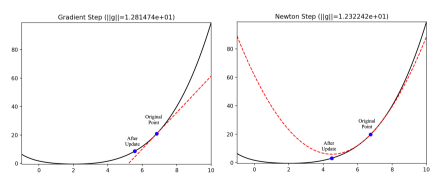


Figure: Kilian Weinberger

Comments on Newton's method

- Storing the Hessian and solving the linear system $\mathbf{H}^{-1}\nabla f$ is very expensive. One way to make it cheaper can be to use only a diagonal approximation to the Hessian.
- Newton's method is not guaranteed to produce a downhill step!
- This only happens (for sure) if ϵ is small and **H** is positive definite.
- If **H** is positive definite, one can use a line search in the direction $\mathbf{H}^{-1}\nabla f$ to ensure we go downhill and make a non-trivial jump.
- Not very useful in general for stochastic objectives.

Quasi-Newton methods

Quasi-Newton methods avoid computing the Hessian, using an approximation instead, finding a matrix ${\bf B}$ which satisfies

$$\nabla f(\mathbf{x}_{t+1}) = \nabla f(\mathbf{x}_t) + \mathbf{B}(\mathbf{x}_{t+1} - \mathbf{x}_t),$$

corresponding to the Taylor expansion of the gradient. This leads to an optimization step

$$\mathbf{x}_{t+1} = \mathbf{x}_t - \epsilon \mathbf{B}_t^{-1} \nabla f(\mathbf{x}_t).$$

- In Broyden-Fletcher-Goldfarb-Shannon (BFGS), an approximate inverse Hessian \mathbf{B}_{t}^{-1} is formed iteratively.
- L-BFGS is a popular practical method that limits the memory requirement.
- This update sort of looks like the Adagrad / RMSProp updates. . . !

Outer product approximations

Suppose we are optimizing a function that decomposes into a sum of squared errors, e.g.

$$L(\boldsymbol{\theta}) = \sum_{n=1}^{N} (y_n - f(\boldsymbol{\theta}, \mathbf{x}_n))^2.$$

The Hessian of this overall loss function has entries

$$\frac{\partial L}{\partial \theta_i \partial \theta_j} = 2 \sum_{n=1}^{N} \left((f(\boldsymbol{\theta}, \mathbf{x}_n) - y_n) \frac{\partial f_n}{\partial \theta_i \partial \theta_j} + \frac{\partial f_n}{\partial \theta_i} \frac{\partial f_n}{\partial \theta_j} \right).$$

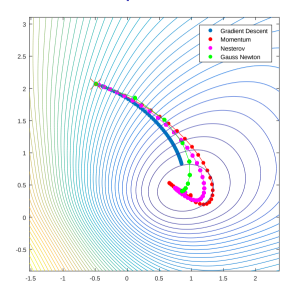
Near the minimum, the first term will be small, as $f(\theta, \mathbf{x}_n) \approx y_n$. This suggests that we can construct a Hessian approximation by using the outer product of the gradients,

$$\mathbf{B} = 2\sum_{i=1}^{N} \nabla f_n(\boldsymbol{\theta}) \nabla f_n(\boldsymbol{\theta})^{\top}.$$

Gauss-Newton on a sum of squares loss

Gauss-Newton using a learning rate of $\epsilon=0.95$ and regularizer $\lambda=0.05$.

The Gauss Newton method converges in fewer iterations, but each iteration is more costly.



Summary

- Gradient descent: general-purpose optimization of real-valued scalar functions
- Stochastic / mini-batch methods can speed up runtime by fast parallel estimation of an approximate step direction
- Momentum and adaptive learning rates can speed up rate of optimization
- Second order methods are memory-intensive and not typically helpful on stochastic objectives, but can converge in far fewer iterations