CSC411: Optimization for Machine Learning

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Overview of Optimization

An informal definition of optimization

Minimize (or maximize) some quantity.

Applications

- ▶ Engineering: Minimize fuel consumption of an automobile
- Economics: Maximize returns on an investment
- Supply Chain Logistics: Minimize time taken to fulfill an order
- Life: Maximize happiness

More formally

Goal: find $\theta^* = \operatorname{argmin}_{\theta} f(\theta)$, (possibly subject to constraints on θ).

- $\theta \in \mathbb{R}^n$: optimization variable
- $f: \mathbb{R}^n \to \mathbb{R}$: objective function

Maximizing $f(\theta)$ is equivalent to minimizing $-f(\theta)$, so we can treat everything as a minimization problem.

Optimization is a large area of research

The best method for solving the optimization problem depends on which assumptions we want to make:

- ▶ Is θ discrete or continuous?
- ▶ What form do constraints on θ take? (if any)
- ▶ Is f "well-behaved"? (linear, differentiable, convex, submodular, etc.)

Optimization for Machine Learning

Often in machine learning we are interested in learning the parameters $\boldsymbol{\theta}$ of a model.

Goal: minimize some loss function

- ► For example, if we have some data (x, y), we may want to maximize $P(y|x, \theta)$.
- ▶ Equivalently, we can minimize $-\log P(y|x,\theta)$.
- We can also minimize other sorts of loss functions

log can help for numerical reasons

Gradient Descent

Gradient Descent: Motivation

From calculus, we know that the minimum of f must lie at a point where $\frac{\partial f(\theta^*)}{\partial \theta} = 0$.

- ▶ Sometimes, we can solve this equation analytically for θ .
- Most of the time, we are not so lucky and must resort to iterative methods.

Review

▶ Gradient: $\nabla_{\theta} f = (\frac{\partial f}{\partial \theta_1}, \frac{\partial f}{\partial \theta_2}, ..., \frac{\partial f}{\partial \theta_k})$

Outline of Gradient Descent Algorithm

Where η is the learning rate and T is the number of iterations:

- ▶ Initialize θ_0 randomly
- ▶ for t = 1 : T:

The learning rate shouldn't be too big (objective function will blow up) or too small (will take a long time to converge)

Gradient Descent with Line-Search

Where η is the learning rate and T is the number of iterations:

- ▶ Initialize θ_0 randomly
- for t = 1 : T:
 - Finding a step size η_t such that $f(\theta_t \eta_t \nabla_{\theta_{t-1}}) < f(\theta_t)$

 - $\blacktriangleright \ \theta_t \leftarrow \theta_{t-1} + \delta_t$

Require a line-search step in each iteration.

Gradient Descent with Momentum

We can introduce a momentum coefficient $\alpha \in [0,1)$ so that the updates have "memory":

- ▶ Initialize θ_0 randomly
- ▶ Initialize δ_0 to the zero vector
- for t = 1 : T:

 - $\theta_t \leftarrow \theta_{t-1} + \delta_t$

Momentum is a nice trick that can help speed up convergence. Generally we choose α between 0.8 and 0.95, but this is problem dependent

Outline of Gradient Descent Algorithm

Where η is the learning rate and T is the number of iterations:

- ▶ Initialize θ_0 randomly
- Do:
- Until convergence

Setting a convergence criteria.

Some convergence criteria

- ► Change in objective function value is close to zero: $|f(\theta_{t+1}) f(\theta_t)| < \epsilon$
- ▶ Gradient norm is close to zero: $\|\nabla_{\theta} f\| < \epsilon$
- Validation error starts to increase (this is called early stopping)

Checkgrad

- ▶ When implementing the gradient computation for machine learning models, it's often difficult to know if our implementation of f and ∇f is correct.
- ► We can use finite-differences approximation to the gradient to help:

$$\frac{\partial f}{\partial \theta_i} \approx \frac{f((\theta_1, \dots, \theta_i + \epsilon, \dots, \theta_n)) - f((\theta_1, \dots, \theta_i - \epsilon, \dots, \theta_n))}{2\epsilon}$$

Why don't we always just use the finite differences approximation?

- slow: we need to recompute f twice for each parameter in our model.
- numerical issues

Stochastic Gradient Descent

- Any iteration of a gradient descent (or quasi-Newton) method requires that we sum over the entire dataset to compute the gradient.
- SGD idea: at each iteration, sub-sample a small amount of data (even just 1 point can work) and use that to estimate the gradient.
- Each update is noisy, but very fast!
- ▶ It can be shown that this method produces an unbiased estimator of the true gradient.
- ► This is the basis of optimizing ML algorithms with huge datasets (e.g., recent deep learning).
- Computing gradients using the full dataset is called batch learning, using subsets of data is called mini-batch learning.

Stochastic Gradient Descent

- The reason SGD works is because similar data yields similar gradients, so if there is enough redundancy in the data, the noise from subsampling won't be so bad.
- SGD is very easy to implement compared to other methods, but the step sizes need to be tuned to different problems, whereas batch learning typically "just works".
- Tip 1: divide the log-likelihood estimate by the size of your mini-batches. This makes the learning rate invariant to mini-batch size.
- ▶ Tip 2: subsample without replacement so that you visit each point on each pass through the dataset (this is known as an epoch).

Convexity

Definition of Convexity

A function f is **convex** if for any two points θ_1 and θ_2 and any $t \in [0,1]$,

$$f(t\theta_1+(1-t)\theta_2)\leq tf(\theta_1)+(1-t)f(\theta_2)$$

We can *compose* convex functions such that the resulting function is also convex:

- ▶ If f is convex, then so is αf for $\alpha \geq 0$
- ▶ If f_1 and f_2 are both convex, then so is $f_1 + f_2$
- etc., see http://www.ee.ucla.edu/ee236b/lectures/functions.pdf for more

Why do we care about convexity?

- Any local minimum is a global minimum.
- ► This makes optimization a lot easier because we don't have to worry about getting stuck in a local minimum.

Examples of Convex Functions

Quadratics

```
Slide Type
In [6]:
         import matplotlib.pyplot as plt
        plt.xkcd()
         theta = linspace(-5, 5)
         f = theta**2
        plt.plot(theta, f)
Out[6]: [<matplotlib.lines.Line2D at 0x3ceae90>]
         20 -
          15 -
          10
          5 -
```

Examples of Convex Functions

Negative logarithms

```
Slide Type
In [8]:
         import matplotlib.pyplot as plt
        plt.xkcd()
        theta = linspace(0.1, 5)
         f = -np.log(theta)
        plt.plot(theta, f)
Out[8]: [<matplotlib.lines.Line2D at 0x3ef4a10>]
          2.0
          1.5
          1.0
          0.5
          0.0
         -0.5
         -1.0
         -1.5
         -2.0L
```

Convexity for logistic regression

Cross-entropy objective function for logistic regression is also convex!

$$f(\theta) = -\sum_{n} t^{(n)} \log p(y = 1 | x^{(n)}, \theta) + (1 - t^{(n)}) \log p(y = 0 | x^{(n)}, \theta)$$

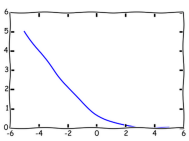
Plot of $-\log \sigma(\theta)$

```
In [15]:

def sigmoid(x):
    return 1 / (1 + np.exp(-x))

theta = linspace(-5, 5)
    f = -np.log(sigmoid(theta))
    plt.plot(theta, f)
```

Out[15]: [<matplotlib.lines.Line2D at 0x4c453d0>]



More on optimization

- ➤ Automatic Differentiation Modern technique (used in libraries like tensorflow, pytorch, etc) to efficiently compute the gradients required for optimization. A survey of these techniques can be found here: https://arxiv.org/pdf/1502.05767.pdf
- Convex Optimization by Boyd & Vandenberghe Book available for free online at http://www.stanford.edu/~boyd/cvxbook/
- Numerical Optimization by Nocedal & Wright Electronic version available from UofT Library

Cross-Validation

Cross-Validation: Why Validate?

So far:

Learning as Optimization

Goal: Optimize model complexity (for the task)

while minimizing under/overfitting

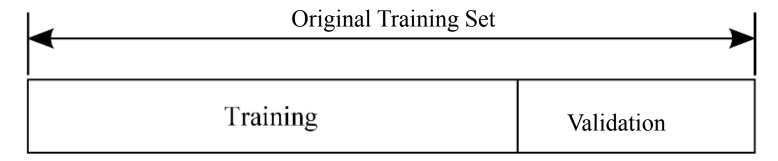
We want our model to **generalize well** without **overfitting**.

We can ensure this by validating the model.

Types of Validation

Hold-Out Validation: Split data into training and validation sets.

Usually 30% as hold-out set.



Problems:

- Waste of dataset
- Estimation of error rate might be misleading

Types of Validation

Cross-Validation: Random subsampling

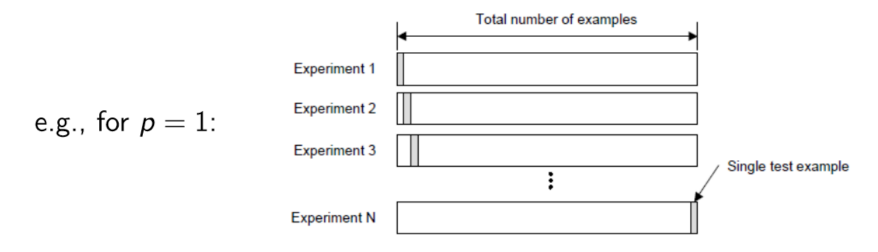


Problem:

 More computationally expensive than holdout validation.

Variants of Cross-Validation

Leave-*p***-out**: Use *p* examples as the validation set, and the rest as training; repeat for all configurations of examples.

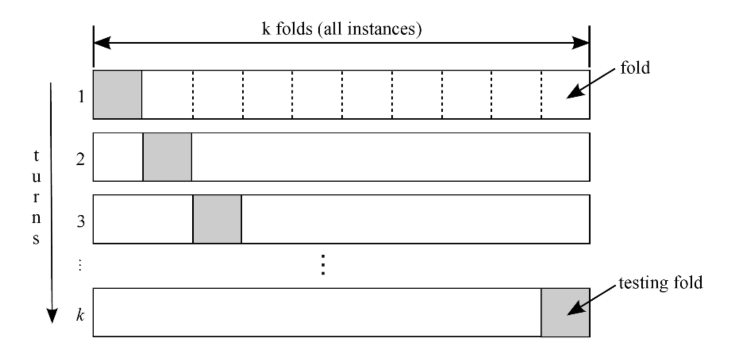


Problem:

• **Exhaustive**. We have to train and test $\binom{N}{p}$ times, where N is the # of training examples.

Variants of Cross-Validation

K-fold: Partition training data into K equally sized subsamples. For each fold, use the other K-1 subsamples as training data with the last subsample as validation.



K-fold Cross-Validation

 Think of it like leave-p-out but without combinatoric amounts of training/testing.

Advantages:

- All observations are used for both training and validation. Each observation is used for validation exactly once.
- Non-exhaustive: More tractable than leave-pout

K-fold Cross-Validation

Problems:

- Expensive for large N, K (since we train/test K models on N examples).
 - But there are some efficient hacks to save time...
- Can still overfit if we validate too many models!
 - Solution: Hold out an additional test set before doing any model selection, and check that the best model performs well on this additional set (*nested cross-validation*).
 Cross-Validception

Practical Tips for Using K-fold Cross-Val

Q: How many folds do we need?

A: With larger K, ...

- Error estimation tends to be more accurate
- But, computation time will be greater

In practice:

- Usually use $K \approx 10$
- BUT, larger dataset => choose smaller K