# Optimization Machine Learning and Pattern Recognition

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(These slides have been adapted from previous versions by Charles Sutton, Amos Storkey, David Barber, and from Sam Roweis (1972-2010))

# Outline

Unconstrained Optimization Problems

- Gradient descent
- Second order methods
- Constrained Optimization Problems
  - Linear programming
  - Quadratic programming
- Non-convexity
- Reading: Murphy 8.3.2, 8.3.3, 8.5.2.3, 7.3.3.
   Barber A.3, A.4, A.5 up to end A.5.1, A.5.7, 17.4.1 pp 379-381.

# Why Numerical Optimization?

- Logistic regression and neural networks both result in likelihoods that we cannot maximize in closed form.
- ► End result: an "error function" E(w) which we want to minimize.
- Note  $\operatorname{argmin} f(\mathbf{x}) = \operatorname{argmax} f(\mathbf{x})$
- $\blacktriangleright$  e.g.,  $E(\mathbf{w})$  can be the negative of the log likelihood.
- Consider a fixed training set; think in weight (not input) space. At each setting of the weights there is some error (given the fixed training set): this defines an error surface in weight space.
- Learning  $\equiv$  descending the error surface.



# Role of Smoothness

If E completely unconstrained, minimization is impossible.



All we could do is search through all possible values  $\mathbf{w}$ .

Key idea: If E is continuous, then measuring  $E(\mathbf{w})$  gives information about E at many nearby values.

#### Role of Derivatives

Another powerful tool that we have is the gradient

$$\nabla E = \left(\frac{\partial E}{\partial w_1}, \frac{\partial E}{\partial w_2}, \dots, \frac{\partial E}{\partial w_D}\right)^T.$$

- Two ways to think of this:
  - ► Each  $\frac{\partial E}{\partial w_k}$  says: If we wiggle  $w_k$  and keep everything else the same, does the error get better or worse?
  - The function

$$f(\mathbf{w}) = E(\mathbf{w}_0) + (\mathbf{w} - \mathbf{w}_0)^\top \nabla E|_{\mathbf{w}_0}$$

is a linear function of w that approximates E well in a neighbourhood around  $w_0$ . (Taylor's theorem)

 Gradient points in the direction of steepest error ascent in weight space.

# Numerical Optimization Algorithms

 Numerical optimization algorithms try to solve the general problem

# $\min_{\mathbf{w}} E(\mathbf{w})$

- Different types of optimization algorithms expect different inputs
  - Zero-th order: Requires only a procedure that computes E(w).
     These are basically search algorithms.
  - First order: Also requires the gradient  $\nabla E$
  - Second order: Also requires the Hessian matrix  $\nabla \nabla E$
  - ▶ High order: Uses higher order derivatives. Rarely useful.
  - ► Constrained optimization: Only a subset of w values are legal.
- Today we'll discuss first order, second order, and constrained optimization

# **Optimization Algorithm Cartoon**

 Basically, numerical optimization algorithms are iterative. They generate a sequence of points

```
\mathbf{w}_0, \mathbf{w}_1, \mathbf{w}_2, \dots

E(\mathbf{w}_0), E(\mathbf{w}_1), E(\mathbf{w}_2), \dots

\nabla E(\mathbf{w}_0), \nabla E(\mathbf{w}_1), \nabla E(\mathbf{w}_2), \dots
```

Basic optimization algorithm is

```
initialize \mathbf{w}

while E(\mathbf{w}) is unacceptably high

calculate \mathbf{g} = \nabla E

Compute direction \mathbf{d} from \mathbf{w}, E(\mathbf{w}), \mathbf{g}

(can use previous gradients as well...)

\mathbf{w} \leftarrow \mathbf{w} - \eta \mathbf{d}

end while

return \mathbf{w}
```

## Gradient Descent

- Locally the direction of steepest descent is the gradient.
- Simple gradient descent algorithm:

```
initialize w

while E(\mathbf{w}) is unacceptably high

calculate \mathbf{g} \leftarrow \frac{\partial E}{\partial \mathbf{w}}

\mathbf{w} \leftarrow \mathbf{w} - \eta \mathbf{g}

end while

return \mathbf{w}
```

- $\eta$  is known as the step size (sometimes called *learning rate*)
  - We must choose  $\eta > 0$ .
  - $\eta$  too small  $\rightarrow$  too slow
  - $\eta$  too large  $\rightarrow$  instability

# Effect of Step Size



► Take  $\eta = 0.1$ . Works well.  $w_0 = 1.0$   $w_1 = \mathbf{w}_0 - 0.1 \cdot 2w_0 = 0.8$   $w_2 = \mathbf{w}_1 - 0.1 \cdot 2w_1 = 0.64$   $w_3 = \mathbf{w}_2 - 0.1 \cdot 2w_2 = 0.512$ ...  $w_{25} = 0.0047$ 

# Effect of Step Size



Take η = 1.1. Not so good. If you step too far, you can leap over the region that contains the minimum

$$w_{0} = 1.0$$
  

$$w_{1} = \mathbf{w}_{0} - 1.1 \cdot 2w_{0} = -1.2$$
  

$$w_{2} = \mathbf{w}_{1} - 1.1 \cdot 2w_{1} = 1.44$$
  

$$w_{3} = \mathbf{w}_{2} - 1.1 \cdot 2w_{2} = -1.72$$
  
...  

$$w_{25} = 79.50$$

Finally, take η = 0.000001. What happens here?

#### Batch vs online

So far all the objective functions we have seen look like:

$$E(\mathbf{w}; D) = \sum_{n=1}^{n} E^{n}(\mathbf{w}; y^{n}, \mathbf{x}^{n}).$$

 $D=\{(\mathbf{x}^1,y^1),(\mathbf{x}^2,y^2),\ldots(\mathbf{x}^n,y^n)\}$  is the training set.

- Each term sum depends on only one training instance
- The gradient in this case is always

$$\frac{\partial E}{\partial \mathbf{w}} = \sum_{n=1}^{N} \frac{\partial E^n}{\partial \mathbf{w}}$$

- The algorithm on slide 8 scans all the training instances before changing the parameters.
- Seems dumb if we have millions of training instances. Surely we can get a gradient that is "good enough" from fewer instances, e.g., a couple thousand? Or maybe even from just one?

#### Batch vs online

 Batch learning: use all patterns in training set, and update weights after calculating

$$\frac{\partial E}{\partial \mathbf{w}} = \sum_{n=1}^{N} \frac{\partial E^n}{\partial \mathbf{w}}$$

- ► On-line learning: adapt weights after each pattern presentation, using <u>∂E<sup>n</sup></u>/∂w
- Batch more powerful optimization methods
- Batch easier to analyze
- **On-line** more feasible for huge or continually growing datasets
- On-line may have ability to jump over local optima

# Algorithms for Batch Gradient Descent

Here is batch gradient descent.

```
 \begin{array}{l} \text{initialize } \mathbf{w} \\ \text{while } E(\mathbf{w}) \text{ is unacceptably high} \\ \quad \text{calculate } \mathbf{g} \leftarrow \sum_{n=1}^{N} \frac{\partial E^n}{\partial \mathbf{w}} \\ \quad \mathbf{w} \leftarrow \mathbf{w} - \eta \ \mathbf{g} \\ \text{end while} \\ \end{array}
```

return  $\mathbf{w}$ 

▶ This is just the algorithm we have seen before. We have just "substituted in" the fact that  $E = \sum_{n=1}^{N} E^n$ .

Algorithms for Online Gradient Descent

- Here is (a particular type of) online gradient descent algorithm initialize w
   while E(w) is unacceptably high

   Pick j as uniform random integer in 1...N
   calculate g ← ∂E<sup>j</sup>/∂w
   w ← w η g
   end while
   return w
- This version is also called "stochastic gradient ascent" because we have picked the training instance randomly.
- There are other variants of online gradient descent.

# Problems With Gradient Descent

- Setting the step size  $\eta$
- Shallow valleys
- Highly curved error surfaces
- Local minima

# Shallow Valleys

Typical gradient descent can be fooled in several ways, which is why more sophisticated methods are used when possible. One problem:



- Gradient descent goes very slowly once it hits the shallow valley.
- One hack to deal with this is momentum

$$\mathbf{d}_t = \beta \mathbf{d}_{t-1} + (1-\beta)\eta \nabla E(\mathbf{w}_t)$$

Now you have to set both η and β. Can be difficult and irritating.

# Curved Error Surfaces

A second problem with gradient descent is that the gradient might not point towards the optimum. This is because of curvature



- Note: gradient is the *locally* steepest direction. Need not directly point toward local optimum.
- Local curvature is measured by the Hessian matrix:  $H_{ij} = \partial^2 E / \partial w_i w_j.$
- By the way, do these ellipses remind you of anything?

#### Second Order Information

Taylor expansion

$$E(\mathbf{w} + \boldsymbol{\delta}) \simeq E(\mathbf{w}) + \boldsymbol{\delta}^T \nabla_{\mathbf{w}} E + \frac{1}{2} \boldsymbol{\delta}^T H \boldsymbol{\delta}$$

where

$$H_{ij} = \frac{\partial^2 E}{\partial w_i \partial w_j}$$

- H is called the Hessian.
- If H is positive definite, this models the error surface as a quadratic bowl.

## Quadratic Bowl



# **Direct Optimization**

A quadratic function

$$E(\mathbf{w}) = \frac{1}{2}\mathbf{w}^T H\mathbf{w} + \mathbf{b}^T \mathbf{w}$$

can be minimised directly using

$$\mathbf{w} = -H^{-1}\mathbf{b}$$

but this requires

- ► Knowing/computing H, which has size O(D<sup>2</sup>) for a D-dimensional parameter space
- Inverting H,  $O(D^3)$

#### Newton's Method

Use the second order Taylor expansion

$$E(\mathbf{w} + \boldsymbol{\delta}) \simeq E(\mathbf{w}) + \boldsymbol{\delta}^T \nabla_{\mathbf{w}} E + \frac{1}{2} \boldsymbol{\delta}^T H \boldsymbol{\delta}$$

- From the last slide, the minimum of the approximation is  $\delta^* = -H^{-1} \nabla_{\mathbf{w}} E$
- Use that as the direction in steepest descent
- ▶ This is called *Newton's method*.
- You may have heard of Newton's method for finding a root, i.e., a point x such that f(x) = 0. Similar thing, we are finding zeros of ∇f.
- Compare Newton step to gradient descent  $\boldsymbol{\delta} = -\eta \nabla_{\mathbf{w}} E$

## Advanced First Order Methods

- Newton's method is fast in that once you are close enough to a minimum.
- What we mean by this is that it needs very few iterations to get close to the optimum (You can actually prove this if you take an optimization course)
- If you have a not-too-large number of parameters and instances, this is probably method of choice.
- But for most ML problems, it is slow. Why? How many second derivatives are there?
- Instead we use "fancy" first-order methods that try to approximate second order information using only gradients.
- These are the state of the art for batch methods
  - One type: Quasi-Newton methods (I like one called *limited* memory BFGS).
  - Conjugate gradient
  - We won't discuss how these work, but you should know that they exist so that you can use them.

#### Constrained problems

• Constraints: e.g.  $f(\mathbf{w}) < 0$ .

- Example: Observe the points {0.5, 1.0} from a Gaussian with known mean μ = 0.8 and unknown standard deviation σ.
   Want to estimate σ by maximum likelihood.
- Constraint: σ must be positive.
- In this case to find the maximum likelihood solution, the optimization problem is

$$\max_\sigma \sum_{i=1}^2 [-\frac{1}{2\sigma^2}(x^i-\mu)^2 - \frac{1}{2}\log(2\pi\sigma^2)]$$
 subject to  $\sigma>0$ 

In this case: solution can be done analytically. More complex cases require a numerical method for constrained optimization.

#### Constrained problems

- Either remove constraints by re-parameterization. E.g.  $\mathbf{w} > 0$ . Set  $\phi = \log(\mathbf{w})$ . Now  $\phi$  unconstrained.
  - Or use a constrained optimization method, e.g. for linear programming, quadratic programming.

# Linear Programming

- Find optimum, within a (potentially unbounded) polytope, of a linear function
- Polytope = polygon or higher dimensional generalization thereof.
- Easy: maximum (if it exists) must be at vertex of polytope (or on a convex set containing such a vertex). Hill climb on vertices using an adjacency walk (Simplex algorithm)

# Quadratic Programming

- Find optimum, within a (potentially unbounded) polytope, of a quadratic form
- Interior point methods, Active set methods.
- Second order methods for convex quadratic functions Newton-Raphson, Conjugate Gradient variants.
- A number of machine learning methods are cast as quadratic programming problems (e.g. Support Vector Machines).

## Non-convexity and local minima

If you follow the gradient, where will you end up? Once you hit a local minimum, gradient is 0, so you stop.



Certain nice functions, such as the likelihood for linear and logistic regression are *convex*, meaning that the second derivative is always positive. This implies that any local minimum is global.

- Dealing with local minima: Train multiple models from different starting places, and then choose best (or combine in some way).
- No guarantees. Unrealistic to believe this will find global mimimum.
- Local minima occur, e.g. for neural networks
- ▶ Bayesian interpretation, where  $E(\mathbf{w}) = -\log p(\mathbf{w}|D)$
- ► Finding local minima of E(w) as a way of approximating integration over the posterior by finding local maxima of p(w|D)

#### **Convex Functions**

• A function  $f : \mathbb{R}^d \to \mathbb{R}$  is convex if for  $\alpha \in [0, 1]$ 

$$f(\alpha \mathbf{x} + (1 - \alpha)\mathbf{y}) \le \alpha f(\mathbf{x}) + (1 - \alpha)f(\mathbf{y})$$

Essentially "bowl shaped"

Examples:

$$f(x) = x^2$$
  $f(x) = -\log x$   $f(\mathbf{x}) = \log\left(\sum_d \exp\{x_d\}\right)$ 

If f differentiable, this implies

$$f(\mathbf{x}_0) + (\mathbf{x} - \mathbf{x}_0)^\top \nabla f|_{\mathbf{x}_0} \le f(\mathbf{x})$$

for all x and  $x_0$ . (To see this: take limit of above as  $x \rightarrow y$ .) This implies that any local minimum is a global one!

# **Convex Optimization Problems**

A convex optimization problem is one that can be written as

 $\min f_0(\mathbf{x})$  subject to  $f_i(\mathbf{x}) \leq 0$   $i \in \{1 \dots N\}$ 

for some choice of functions  $f_0 \dots f_N$  where each  $f_i$  is convex

- Optimise convex function over a convex set...
- Unconstrained problems: Use methods from before. You'll find a global optimum!
- Convexity means any local optimum is also global optimum.
- Constrained convex problems: Interior point methods, Active set methods.
- Most convex optimization problems can be solved efficiently in practice.
- (How high a scale you can reach depends on the type of problem you have)

# Optimization: Summary

- Complex mathematical area. Do not implement your own optimization algorithms if you can help it!
- My advice: For unconstrained problems
  - Batch is less hassle than online. But if you have big data, you must use online. Batch is too slow
  - (For neural networks, typically online methods are method of choice.)
  - If online, you use gradient descent. Forget about second order stuff.
  - If batch, use one of the fancy first-order methods (quasi-Newton or conjugate gradients). DO NOT implement either of these yourself!
- If you have a constrained problem
  - Linear programs are easy. Use off the shelf tools.
  - More than that: Try to convert into unconstrained problem.
- Convex problems: Global minimum. Non-convex: Local optima.

#### What you should take away

- Complex mathematical area. Do not implement your own optimization algorithms if you can help it!
- Stuff you should understand:
  - How and why we convert learning problems into optimization problems
  - Modularity between modelling and optimization
  - Gradient descent
  - Why gradient descent can run into problems
  - Especially local minima
- Methods of choice: Fancy first-order methods (e.g., quasi-Newton, CG) for moderate amounts of data. Stochastic gradient for large amounts of data.