# LECTURE 12: ROOT-FINDING AND MINIMIZATION

STAT 545: INTRO. TO COMPUTATIONAL STATISTICS

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## **ROOT-FINDING IN ONE-DIMENSION**

Given some nonlinear function  $f: \mathbb{R} \to \mathbb{R}$ , solve

$$f(x) = 0$$

Invariably need iterative methods.

Assume *f* is continuous (else things are really messy).

More we know about f (e.g. gradients), better we can do.

Better: faster (asymptotic) convergence.

## **ROOT BRACKETING**

f(a) and f(b) have opposite signs  $\rightarrow$  root lies in (a,b).

a and b bracket the root.

Finding an initial bracketing can be non-trivial.

Typically, start with an initial interval and expand or contract.

Below, we assume we have an initial bracketing.

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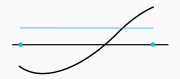
Below, we assume we have an initial bracketing.

Not always possible e.g.  $f(x) = (x - a)^2$  (in general, multiple roots/nearby roots lead to trouble).

Simplest root-finding algorithm.

Given an initial bracketing, cannot fail.

But is slower than other methods.



- Current interval = (a,b)
- Set  $C = \frac{a+b}{2}$
- New interval = (a, c) or (c, b)(whichever is a valid bracketing)

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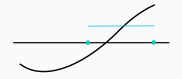


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- every (fixed) k iterations reduces error by one digit.
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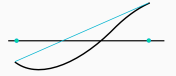
Superlinear convergence:

$$\lim_{n \to \infty} |\epsilon_{n+1}| = C \times |\epsilon_n|^m \qquad (m > 1)$$

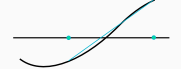
Quadratic convergence:

Number of significant figures doubles every iteration.

Linearly approximate f to find new approximation to root.



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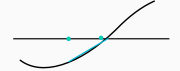
#### Secant method:

- · always keep the newest point
- Superlinear convergence (m = 1.618, the golden ratio)

$$\lim_{n\to\infty} |\epsilon_{n+1}| = C \times |\epsilon_n|^{1.618}$$

· Bracketing (and thus convergence) not guaranteed.

Linearly approximate *f* to find new approximation to root.



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## False position:

- · Can choose an old point that guarantees bracketing.
- · Convergence analysis is harder.

#### PRACTICAL ROOT-FINDING

In practice, people use more sophiticated algorithms.

Most popular is Brent's method.

Maintains bracketing by combining bisection method with a quadratic approximation.

Lots of book-keeping.

At any point uses both function evaluation as well as derivative to form a linear approximation.

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Taylor expansion: 
$$f(x + \delta) = f(x) + \delta f'(x) + \frac{\delta^2}{2} f''(x) + \cdots$$

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Assume second- and higher-order terms are negligible. Given  $x_i$ , choose  $x_{i+1} = x_i + \delta$  so that  $f(x_{i+1}) = 0$ :

# Newton's method (a.k.a. Newton-Raphson)

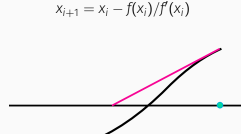
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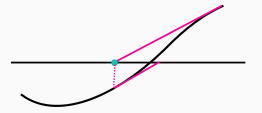
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$$0 = f(x_i) + \delta f'(x_i)$$

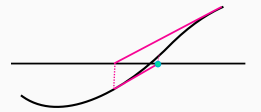
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Letting  $x^*$  be the root, we have

$$X_{i+1} - X^* = X_i - X^* - f(X_i)/f'(X_i)$$
  

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Also since  $x_i = x^* + \epsilon_i$ ,

$$f(x_i) \approx f(x^*) + \epsilon_i f'(x^*) + \frac{\epsilon_i^2}{2} f''(x^*)$$

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This gives

$$\epsilon_{i+1} = -\frac{f'(x_i)}{2f''(x_i)}\epsilon_i^2$$

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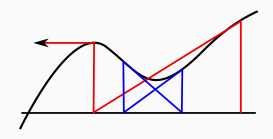
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Quadratic convergence (assuming f'(x) is non-zero at the root)

# PITFALLS OF NEWTON'S METHOD



Away from the root the linear approximation can be bad.

Can give crazy results (go off to infinity, cycles etc.)

However, once we have a decent solution can be used to rapidly 'polish the root'.

Often used in combination with some bracketing method.

# ROOT-FINDING FOR SYSTEMS OF NONLINEAR EQUATIONS

Now have N functions  $F_1, F_2, \dots, F_N$  of N variables  $x_1, x_2, \dots, x_N$ Find  $(x_1, \dots, x_N)$  such that:

$$F_i(x_1, \cdots, x_N) = 0$$
  $i = 1 \text{ to } N$ 

Much harder than the 1-d case.

Much harder than optimization.

# **NEWTON'S METHOD**

Again, consider a Taylor expansion:

$$F(x + \delta x) = F(x) + J(x) \cdot \delta x + O(\delta x^{2})$$

Here,  $J(\mathbf{x})$  is the Jacobian matrix at  $\mathbf{x}$ , with  $J_{ij} = \frac{\partial F_i}{\partial x_j}$ .

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Again, Newton's method finds  $\delta \mathbf{x}$  by solving  $\mathbf{F}(\mathbf{x} + \delta \mathbf{x}) = 0$ 

$$J(x) \cdot \delta x = -F(x)$$

Solve  $\delta \mathbf{x} = -\mathbf{J}(\mathbf{x})^{-1} \cdot \mathbf{F}(\mathbf{x})$  (e.g. by LU decomposition)

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Iterate  $\mathbf{x}_{new} = \mathbf{x}_{old} + \delta \mathbf{x}$  until convergence.

Can wildly careen through space if not careful.

# GLOBAL METHODS VIA OPTIMIZATION

Recall, we want to solve  $\mathbf{F}(\mathbf{x}) = 0$   $(F_i(\mathbf{x}) = 0, i = 1 \cdots N)$ .

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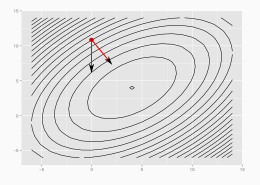
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.

Note: It is NOT sufficient to find a local minimum of f.

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We move along  $\delta \mathbf{x}$  instead of  $\nabla f = \mathbf{F}(\mathbf{x})\mathbf{J}(\mathbf{x})$ .

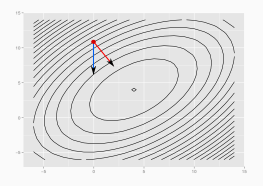
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Note: 
$$\nabla f \cdot \delta \mathbf{x} = (\mathbf{F}(\mathbf{x})\mathbf{J}(\mathbf{x})) \cdot (-\mathbf{J}^{-1}(\mathbf{x})\mathbf{F}(\mathbf{x})) = -\mathbf{F}(\mathbf{x})\mathbf{F}(\mathbf{x}) < 0$$

## NEWTON'S METHOD WITH BACKTRACKING

A full Newton step sets  $\mathbf{x}_{new} = \mathbf{x}_{old} + \delta \mathbf{x}$ .

This can cause f to increase i.e.  $f(\mathbf{x}_{new} > f(\mathbf{x}_{old})$ .

In this case, backtrack and set  $\mathbf{x}_{new} = \mathbf{x}_{old} + \lambda \delta \mathbf{x}, \quad \lambda \in (0, 1).$ 

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Finding best  $\lambda$ : too much work usually.

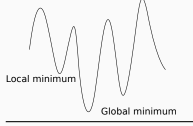
However, just causing f to decrease is not sufficient.

Can use Wolfe conditions (later)

#### GLOBAL AND LOCAL MINIMUM

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Finding a global minimum is hard! Usually settle for finding a local minimum (like the EM algorithm).

Conceptually (deceptively?) simpler than EM.

Let  $x_{old}$  be our current value.

Update 
$$x_{new}$$
 as  $x_{new} = x_{old} - \eta \left. \frac{\mathrm{d}f}{\mathrm{d}x} \right|_{x_{old}}$ 

The steeper the slope, the bigger the move.

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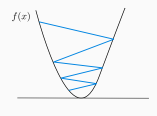
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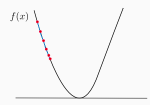
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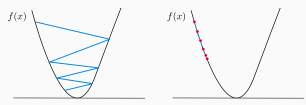
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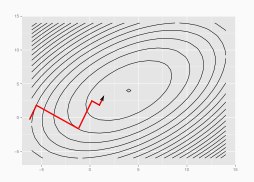
Better methods adapt step-size according to the curvature of f.

#### GRADIENT DESCENT IN HIGHER-DIMENSIONS

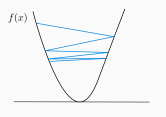
Steepest descent also applies to higher dimensions too:

$$x_{new} = x_{old} - \eta |\nabla f|_{x_{old}}$$

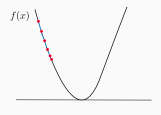
Now, even using the optimal  $\eta$  can be inefficient:



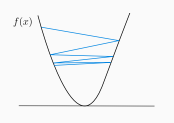
More on this later.



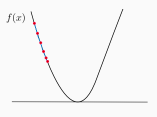
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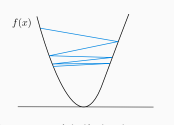
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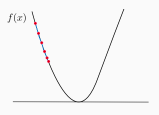
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Avg. decrease at least some fraction of initial rate:

$$f(\mathbf{x} + \lambda \delta \mathbf{x}) \le f(\mathbf{x}) + c_1 \lambda (\nabla f \cdot \delta \mathbf{x}), \qquad c_1 \in (0, 1) \text{ e.g. } 0.9$$



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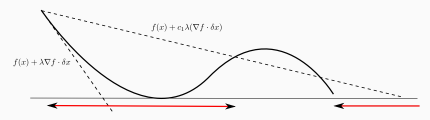
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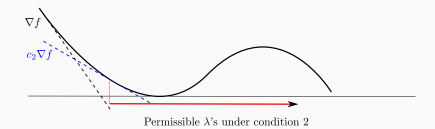
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Final rate is greater than some fraction of initial rate:

$$\nabla f(\mathbf{x} + \lambda \delta \mathbf{x}) \cdot \delta \mathbf{x} \ge c_2 \nabla f(\mathbf{x}) \delta \mathbf{x},$$
  $c_2 \in (0, 1) \text{ e.g. } 0.1$ 



Permissible  $\lambda$ 's under condition 1



A simple way to satisfy Wolfe conditions:

Set 
$$\delta x = -\nabla f, c_1 = c_2 = .5$$

Start with  $\lambda = 1$ , and while condition i is not satisfied, set

$$\lambda = \beta_i t \text{ (for } \beta_1 \in (0,1), \beta_2 > 1 \text{ and } \beta_1 * \beta_2 < 1$$

### **ESTIMATING MLE**

Consider a set of observations  $Y = (y_1, \dots, y_N)$ .

Assume 
$$y_i \sim p(y|\theta)$$

$$\theta_{MLE} = \operatorname{argmax} \ell(\theta) = \operatorname{argmax} \sum_{i=1}^{N} \log p(x_i | \theta)$$

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Starting with an initial  $\theta_0$ , iterate:

$$\theta_{i+1} = \theta_i + \eta_i \nabla \ell(\theta_i)$$

# GRADIENT DESCENT (CONTD.)

$$\nabla \ell(\theta) = \sum_{i=1}^{N} \nabla \log p(x_i | \theta)$$

#### Cons:

- Calculating the gradient is O(N). (Each iteration must cycle through all datapoints.)
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#### Pros:

- Convergence is better understood.
- Accelerated methods are available (e.g. Newton's method, conjugate gradient)

### STOCHASTIC GRADIENT DESCENT

Use a noisy gradient  $\widehat{\nabla \ell}$ .

Typically split data into N/B batches of size B. Each iteration, calculate gradient on one of the batches  $B_i$ :

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#### Pros:

- Calculating the gradient is O(B).
   (Often, each batch is just a single datapoint)
- Much faster convergence (just one sweep through the data can get you a decent solution).
- Often, you get better solutions.
- $\cdot$  Useful for online systems, tracking heta that varies over time .

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#### Cons:

- · Convergence analysis is harder.
- Noisy gradients mean the algorithm will never converge.
   Typically need to reduce the step size every iteration.
   We want

$$\eta_i \to 0, \quad \sum_{i=1}^{\infty} \eta_i = \infty$$

E.g. 
$$\eta_i = \frac{a}{b+i}$$

#### STOCHASTIC GRADIENT DESCENT WITH MOMENTUM

One way to accelarate convergence is to include a momentum term:

$$\theta_{i+1} = \theta_i + \eta_i \widehat{\nabla \ell}(\theta_i) + \beta_i \underbrace{(\theta_i - \theta_{i-1})}_{\text{momentum}}$$

### STOCHASTIC GRADIENT DESCENT WITH MOMENTUM

One way to accelarate convergence is to include a momentum term:

$$\theta_{i+1} = \theta_i + \eta_i \widehat{\nabla \ell}(\theta_i) + \beta_i \underbrace{(\theta_i - \theta_{i-1})}_{\text{momentum}}$$

More generally,

$$\theta_{i+1} = \theta_i + \eta_i \widehat{\nabla \ell}(\theta_i + \gamma(\theta_i - \theta_{i-1})) + \beta_i(\theta_i - \theta_{i-1})$$

Include many popular algorithms:

- Polyak's heavy ball method (HB):  $\gamma = 0$
- · Nesterov's accelerated gradient (NAG):  $\gamma_i = \beta_i$

#### **ADAPTIVE METHODS**

Adaptive methods accelerate convergence by using the entire history of iterates to determine step-sizes.

Often take the general form

$$\theta_{i+1} = \theta_i + \eta_i H_i^{-1} \widehat{\nabla \ell} (\theta_i + \gamma (\theta_i - \theta_{i-1})) + \beta_i H_i^{-1} H_{i-1} (\theta_i - \theta_{i-1})$$

where  $H_i$  is some combination of all previous gradients. E.g.

$$H_i = \operatorname{diag}\left(\sum_{j=1}^i g_j \circ g_j\right),$$

with  $g_j = \widehat{\nabla \ell}(\theta_j + \gamma(\theta_j - \theta_{j-1}))$ , and  $\circ$  element-wise product.

Examples are AdaGrad, Adam etc.