Unconstrained optimization

Nick Henderson, AJ Friend (Stanford University) Kevin Carlberg (Meta)

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Unconstrained optimization

Theory, methods, and software for problems exihibiting the characteristics below

- Convexity:
 - convex: local solutions are global
 - non-convex: local solutions are not global
- Optimization-variable type:
 - continuous: gradients facilitate computing the solution
 - discrete: cannot compute gradients, NP-hard
- Constraints:
 - unconstrained : simpler algorithms
 - constrained: more complex algorithms; must consider feasibility
- Number of optimization variables:
 - low-dimensional: can solve even without gradients
 - high-dimensional: requires gradients to be solvable in practice

Theory

Theory 3

Outline

Theory

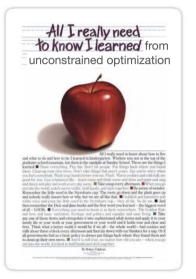
Algorithms

Gradient-based algorithms

Derivative-free algorithms

Theory

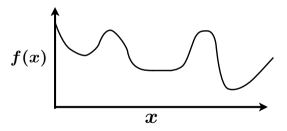
All I really need to know...



Unconstrained optimization in one variable

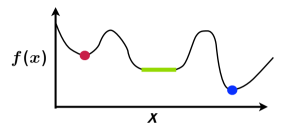
minimize
$$f(x)$$

- $x \in \mathbf{R}$ is a real-valued variable
- $ightharpoonup f(x) \in C^2: \mathbf{R} \to \mathbf{R}$ is the objective function, which returns a single real number



▶ What is a solution to this problem?

What is a solution?



- ▶ Global minumum: A point x^* satisfying $f(x^*) \le f(x)$ for all x in the domain of interest
- Strong local minumum: A point x^* satisfying $f(x^*) < f(x)$ for all x in a neighborhood of x^*
- ▶ Weak local minumum: A point x^* satisfying $f(x^*) \le f(x)$ for all x in a neighborhood of x^*

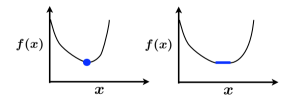
Theory

Convexity

For a convex objective function in one variable, we have

$$f(tx + (1-t)y) \le tf(x) + (1-t)f(y)$$

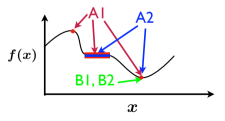
 $\text{ for } t \in [0,1]$



► Any local minimum is a global minimum!

Optimality conditions for single-variable minimization

- ▶ Necessary conditions for a weak local minimum:
 - **A1**: $f'(x^*) = 0$
 - ▶ **A2**: $f''(x^*) \ge 0$
- ► Sufficient conditions for a strong local minimum:
 - ▶ **B1**: $f'(x^*) = 0$, and
 - **B2**: $f''(x^*) > 0$
- **Stationary point**: a point x^* satisfying $f'(x^*) = 0$
- Saddle point: a stationary point that is not a local minimum or maximum



Unconstrained optimization in multiple variables

minimize f(x)

- $\mathbf{r} \in \mathbf{R}^n$ is an n-dimensional vector of real numbers
- $lackbox{}{} f(x) \in C^2: \mathbf{R}^n
 ightarrow \mathbf{R}$ is the objective function, which returns a single real number
- ► The same notions of weak local, strong local, and global minima, as well as convexity, extend to multiple dimensions

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Derivatives in multiple dimensions

Vector of optimization variables:

$$x = \begin{bmatrix} x_1 \\ x_2 \end{bmatrix}$$

Gradient (i.e., first derivative) of f:

$$\nabla f(x) = \begin{bmatrix} \frac{\partial f}{\partial x_1} \\ \frac{\partial f}{\partial x_2} \end{bmatrix}$$

Hessian (i.e., second derivative) of f:

$$\nabla^2 f(x) = \begin{bmatrix} \frac{\partial^2 f}{\partial x_1^2} & \frac{\partial^2 f}{\partial x_1 \partial x_2} \\ \frac{\partial^2 f}{\partial x_2 \partial x_1} & \frac{\partial^2 f}{\partial x_2^2} \end{bmatrix}$$

Stationary points

Stationary point: a point x^* satisfying $\nabla f(x^*) = 0$

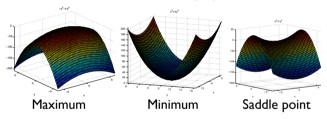


Figure 1: Types of stationary points in two dimensions

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Optimality conditions for multiple-variable minimization

Can simply extend the univariate conditions to multiple dimensions

- ▶ Necessary conditions for a weak local minimum:
 - ▶ **A1**: $\nabla f(x^*) = 0$
 - ▶ **A2**: $\nabla^2 f(x^*) \succeq 0$
 - $ightharpoonup
 abla^2 f(x) \succeq 0$ means that all the eigenvalues of $abla^2 f(x)$ are non-negative
- Sufficient conditions for a strong local minimum:
 - ▶ **B1**: $\nabla f(x^*) = 0$, and
 - ▶ **B2**: $\nabla^2 f(x^*) \succ 0$
 - $lackbox{ }
 abla^2 f(x) \succ 0$ means that all the eigenvalues of $abla^2 f(x)$ are strictly positive

Theory

Algorithms

Outline

Theory

Algorithms

Gradient-based algorithms

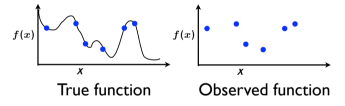
Derivative-free algorithms

Optimization algorithms

- ▶ We now know:
 - What an unconstrained optimization problem is
 - How to characterize local/global solutions using optimality conditions
- ► How do we *compute* these solutions?
 - Analytically: only possible for very simple problems (e.g., Brachistochrone problem)
 - Numerically: required for most practical problems
- ▶ **Numerical optimization** algorithms are used to numerically solve these problems with computers

Optimization algorithms

- ▶ In general, we are *mostly blind* to the function we are trying to minimize.
- We can only compute the function f at a finite number of points, and each evaluation may be computationally expensive



- ▶ Derivative information (gradient ∇f and Hessian $\nabla^2 f$) is sometimes available
 - generally more expensive to compute
 - can help a lot (determine optimality criteria)
 - ightharpoonup especially helpful in high dimensions (n large)

Optimization algorithms

- Goals
 - Practical: reasonable memory requirements
 - ▶ **Robust**: low failure rate, convergence conditions are met
 - ► **Fast**: convergence in a few iterations, low cost per iteration
 - Application typically dictates specific requirements
- Algorithm design involves tradeoffs to achieve these goals
 - Example: using derivatives may reduce number of iterations, but each iteration becomes more expensive
- Algorithms are iterative in nature
- Categorization
 - ► Gradient-based v. derivative-free
 - Global v. local: aims to converge to a global or local minimum
 - Gradient-based algorithms tend to be local, while derivative-free algorithms tend to be global

Gradient-based algorithms

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Gradient-based algorithms

Imagine you are lost on a mountain in extremely thick fog



- ► How would you get down (i.e., find the minimum)?
- ► Chances are, you would use the *slope* of the ground beneath you in some way to go downhill and descend the mountain
- ▶ This is the approach taken by gradient-based algorithms

Gradient-based algorithms: benefits and drawbacks

Benefits

- Efficient for many variables (i.e., in high dimensions)
- Well-suited for smooth objective and constraint functions

Drawbacks

- Requires computing the gradient (challenging in some cases)
- Convergence is only local (local optimization)
 - Mitigated by using multiple initial guesses to find multiple local minima
 - Can then choose the best local minimum
- Not well-suited for discrete optimization
- Not well-suited for noisy functions
- ► Second derivatives (Hessians) are also very valuable
 - ightharpoonup However, Hessians are $n \times n$ symmetric matrices, so expensive to construct and store
 - If Hessians are needed, they are often approximated using quasi-Newton methods

Gradient-based algorithms: framework

- ightharpoonup At each iteration k, gradient-based methods compute both
 - 1. a search direction p_k , and
 - 2. a **step length** α_k (referred to as the **learning rate** in machine learning)

Algorithm 1 Gradient-based framework

Choose initial guess x_0 , set $k \leftarrow 0$

while (not converged) do

Choose direction p_k and step length α_k

(This often involves computing local information, e.g., $\nabla f(x_k)$, $\nabla^2 f(x_k)$)

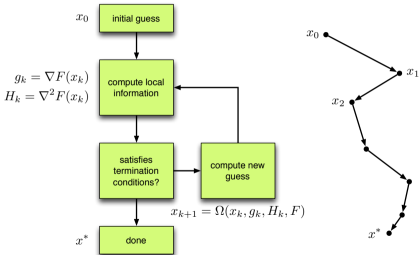
$$x_{k+1} = x_k + \alpha_k p_k.$$

$$k \leftarrow k + 1$$

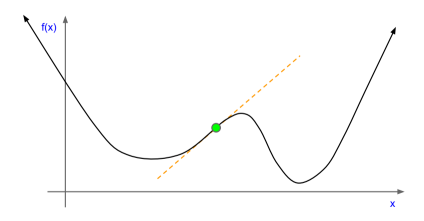
 $\nu \leftarrow \nu +$

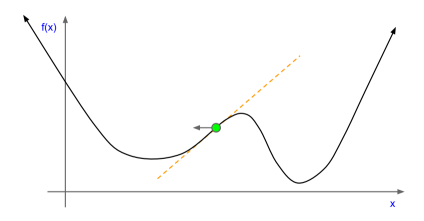
end while

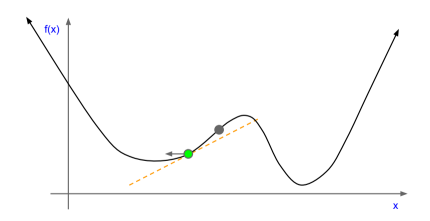
Gradient-based algorithms: overview

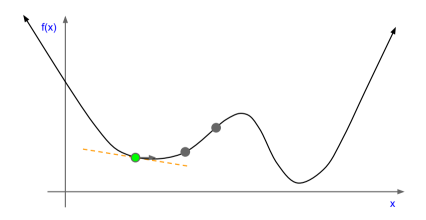


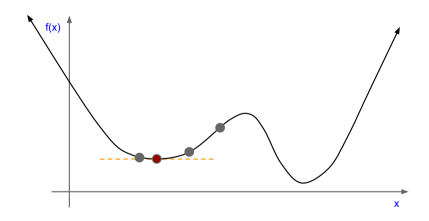
Gradient-based algorithms











Gradient-based algorithms: two classes

There are two classes of gradient-based algorithms.

- Line-search methods:
 - 1. compute p_k to be a descent direction
 - 2. compute α_k to produce a sufficient decrease in the objective function
- Trust-region methods:
 - 1. determine a maximum allowable step length (trust-region radius) Δ_k ,
 - 2. compute step p_k with $||p_k|| \leq \Delta$ using a model $m(p) \approx f(x_k + p)$
 - 3. **accept step** if actual objective-function reduction is close to (or better than) the model-preducted objective-function reduction, and set $x_{k+1} = x_k + p_k$ (note $\alpha_k = 1$)
 - 4. otherwise, **reject step**, set $x_{k+1}=x_k$, and shrink trust-region radius such that $\Delta_{k+1}<\Delta_k$

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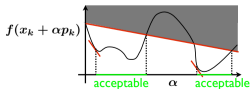
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Line-search methods: convergence

Theorem (Sufficient conditions for convergence)

For sufficiently smooth, well-defined problems, sufficient conditions for convergence $\lim_{k\to\infty}\|\nabla f_k\|=0$ of line search methods are:

- C1. p_k are descent directions $(p_k^T \nabla f(x_k) < 0)$
- C2. α_k produces a sufficient decrease (satisfy the Wolfe conditions)
- C2. Wolfe conditions $(0 < c_1 < c_2 < 1)$:
 - ▶ Decrease f: $f(x_k + \alpha_k p_k) \leq \underline{f}(x_k) + c_1 \alpha_k \nabla f_k^T p_k$
 - ▶ Increase ∇f : $\nabla f(x_k + \alpha_k p_k)^T p_k \ge c_2 \nabla f_k^T p_k$.



Line-search methods: key steps

- 1. Choose search direction p_k that is a descent direction (satisfy C1)
- 2. Choose step length α_k that satsifies the Wolfe conditions (satisfy C2)

Line-search methods: step 1 (gradient descent)

Choose search direction p_k that is a descent direction (satisfy C1)

- ▶ **Gradient descent** (i.e., steepest descent): $p_k = -\nabla f(x_k)$
 - Steepest direction downhill
 - ▶ Advantages: only first-order information, always a descent direction, low storage
 - Disadvantages: linear convergence rate, sensitive to variable scaling
 - Stochastic gradient descent is an approximation of this yielding sublinear convergence
- ► Conjugate gradient: $p_k = -\nabla f(x_k) + \beta_k p_{k-1}$
 - \triangleright β_k computed to ensure p_k and p_{k-1} are approximately conjugate (accounts for previous progress)
 - ► Advantages: only first-order information, low storage, more effective than steepest descent and almost as simple to implement
 - ▶ Disadvantages: linear convergence rate (but faster than steepest descent), sensitive to variable scaling

Line-search methods: step 1 (modified Newton's method)

Choose search direction p_k that is a descent direction (satisfy C1)

Recall that $\nabla f(x^*) = 0$ is a necessary condition for optimality

- ▶ This is just *n* nonlinear equations in *n* unknowns!
- Thus, we could apply Newton's method to solve it and obtain quadratic convergence!
- ▶ This would lead to $p_k = -(\nabla^2 f(x_k))^{-1} \nabla f(x_k)$
- ▶ If f is strongly convex quadratic and $\alpha_k = 1$, this converges in **one iteration**!
 - Scale invariant: this holds regardless of variable scaling
 - Natural step length: $\alpha_k = 1$
 - ► The Hessian overcomes issues with ill-conditioning/poorly scaled variables
- \blacktriangleright However, p_k is not guaranteed to be a descent direction (i.e., might not satisfy C1)
- lacktriangle So, we must *modify* Newton's method to ensure p_k is a descent direction

Line-search methods: step 1 (modified Newton's method)

- ▶ Modified Newton's method: $p_k = -(\nabla^2 f(x_k) + E_k)^{-1} \nabla f(x_k)$
 - lacktriangledown if $abla^2 f(x_k) + E_k$ is positive definite, then p_k is a descent direction
 - ▶ Thus, E_k is computed to ensure $\nabla^2 f(x_k) + E_k$ is positive definite
 - ▶ Advantages: quadratic convergence, scale invariant, natural step length
 - Disadvantage: second-order information (expensive), large storage
- ▶ Quasi-Newton methods: $p_k = -(B_k)^{-1} \nabla f(x_k)$
 - $lacktriangleright B_k$ updated each iteration using only the gradient to satisfy the secant condition

$$B_{k+1}(x_{k+1} - x_k) = \nabla f_{k+1} - \nabla f_k$$

- Popular updates:
 - Symmetric rank-one (SR1): enforces symmetry, rank 1
 - ▶ Broyden, Fletcher, Goldfarb, Shanno (BFGS): enforces positive definiteness, rank 2
- ► Advantages: only first-order information, superlinear convergence, scale invariant, natural step length, limited-memory variant L-BFGS ensures low storage
- ▶ **Disadvantages**: may not be a descent direction (e.g., if SR1), approximate Hessians may be inaccurate and dense

Line-search methods: step 2

Choose step length α_k that satisfies the Wolfe conditions (satisfy C2)

- Backtracking:
 - ▶ **Goal**: given p_k find α such that $f(x_k + \alpha p_k) < f(x_k)$.
 - **Procedure**: start with initial guess $\alpha > 0$ (use $\alpha = 1$ for Newton's method)
 - 1. if $f(x_k + \alpha p_k) < f(x_k)$, then return α , otherwise continue
 - 2. decrease α by some factor $0 < \delta < 1$: $\alpha \leftarrow \delta \alpha$
 - repeat

Gradient-based algorithms: two classes

There are two classes of gradient-based algorithms.

- Line-search methods:
 - 1. compute p_k to be a descent direction
 - 2. compute α_k to produce a sufficient decrease in the objective function
- ► Trust-region methods :
 - 1. determine a maximum allowable step length (trust-region radius) Δ_k ,
 - 2. compute step p_k with $\|p_k\| \leq \Delta$ using a model $m(p) \approx f(x_k + p)$
 - 3. accept step if actual objective-function reduction is close to (or better than) the model-preducted objective-function reduction, and set $x_{k+1} = x_k + p_k$ (note $\alpha_k = 1$)
 - 4. otherwise, **reject step**, set $x_{k+1}=x_k$, and shrink trust-region radius such that $\Delta_{k+1}<\Delta_k$

Trust-region methods: overview

- ► Trust-region methods sequentially minimize an approximate, easy-to-solve model within a local trust region
- ▶ The trust region is the region within which the approximate model is trusted
- ► The subproblem is often *convex* (**sequential convex programming**)



▶ If the step is unacceptable (inaccurate model), the size of the trust region is reduced (we trust the model less) and minimization is repeated around the same point

Trust-region methods

Trust-region methods often use a quadratic model $m_k(p)$ of the true function $f(x_k+p)$ at the point x_k

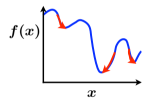
$$m_k(p) = f_k + g_k^T p + \frac{1}{2} p^T B_k p$$

- ▶ If B_k is the exact Hessian, the difference between $m_k(p)$ and $f(x_k + p)$ is $O(\|p\|^3)$
- At each trust-region step, the following constrained problem is approximately solved for p_k

minimize
$$m_k(p)$$
 subject to $||p|| \leq \Delta_k$

Gradient-based algorithms for global optimization

- ► Gradient-based algorithms are best-suited for finding local minima: they "go downhill" until local optimality conditions are satisfied
- ► To find multiple local minima (and hopefully the global minimum), gradient-based methods can be run multiple times using different initial guesses



- ► However even if we happen to find the global minimum, we cannot verify that we have done so! This means we do not know if we've solved the problem.
- ▶ This tuning/babysitting does not arise in convex optimization!

Computation of gradients

- ▶ To implement gradient-based algorithms, derivative information must be computed
- ▶ There are three main ways to compute these gradients
 - 1. Analytical (can use symbolic tools, e.g., Mathematica)
 - 2. Finite differences
 - 3. Automatic differentiation

Finite differences

- ► We can approximate the gradient by evaluating the function several times when the gradient is unavailable analytically
- **Forward-difference**: 1st-order accurate

$$\frac{\partial f}{\partial x_i}(x) = \frac{f(x + \epsilon e_i) - f(x)}{\epsilon} + O(\epsilon)$$

▶ **Central-difference**: 2nd-order accurate, but twice as expensive

$$\frac{\partial f}{\partial x_i}(x) = \frac{f(x + \epsilon e_i) - f(x - \epsilon e_i)}{2\epsilon} + O(\epsilon^2)$$

- Tradeoff:
 - $ightharpoonup \epsilon$ too large: inaccurate due to truncation error
 - ullet too small: inaccurate due to subtractive cancellation from round-off error arising in finite-precision calculations

Automatic differentiation

- Use computational representation of a function
- Key observations:
 - ▶ Any function is composed of a sequence of simple operations
 - ▶ The chain rule from calculus. For f(y(x(w))),

$$\frac{df}{dw} = \frac{df}{dy}\frac{dy}{dx}\frac{dx}{dw}$$

- Performs differentiation on only elemental operations
- Avoids subtractive cancellation
- ► Software tools (e.g. ADIFOR) do this automatically
- ▶ **Backpropagation** in deep learning is a specific case of (reverse mode) automatic differentiation

Derivative-free algorithms

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Derivative-free algorithms

Why derivative-free algorithms?

- Gradients may not be available
 - ightharpoonup f(x) from laboratory experiments
 - impractical or cumbersome to implement analytic gradinets
- ▶ Noise or non-smoothness in the objective function
 - this creates many local minima, so gradient information less useful
 - require global optimization
- ► May want to direct effort *globally* (less information at more points) rather than *locally* (more information at fewer points)
- ▶ Can use global optimization to define initial guesses for local optimization

Benefits and drawbacks of derivative-free algorithms

Benefits:

- Well-suited for discrete variables
- Often better at finding the global optimum (if non-convex)
- ▶ Robust with respect to noise in the function
- Useful for multi-objective optimization
- Amenable to parallelization

Drawbacks:

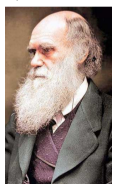
- **Extremely slow convergence in high dimensions (***n* large)
- Difficult to efficiently treat constraints
- Not typically used if gradients are available

Derivative-free algorithm categorization

- ▶ **Heuristic**: use techniques inspired by nature (global optimization)
 - Simulated annealing
 - Basin hopping (Monte Carlo)
 - Evolutionary techniques
 - Genetic algorithms
 - Differential evolution
 - Swarm intelligence (particle swarm optimization, ant colony optimization)
- ▶ **Direct search**: query a sequence of nearby points (local optimization)
 - Directional: coordinate search (e.g., Powell's method), pattern search
 - Simplicial: Nelder–Mead nonlinear simplex

Evolutionary Algorithms

- ► Evolutionary algorithms were invented in the 1960's by John Holland, who wanted to better understand the evolution of life by computer simulation
- ► The algorithm is based on **reproduction** (recombination and mutation) and **selection** (survival of the fittest)



Derivative-free algorithms

Figure 2: Charles Darwin

Evolutionary Algorithm

minimize f(x)

- ightharpoonup A population member is represented by a point x in the variable space (its DNA)
- ightharpoonup 'Fitness' is the objective function value f(x)
- At each iteration, rather than work with a *single point*, we consider an entire *population of points* across the entire space
- ▶ Benefit: more likely to find a global optimum and won't be "trapped" by local minima
- Drawback: very expensive in high dimensions

Overview of evolutionary algorithm

- 1. Initialize population
- 2. Determine mating pool
- 3. Generate children via crossover
 - ► Continuous variables: interpolate
 - Discrete variables: replace parts of their representing variables
- 4. Mutation (add randomness to the children's variables)
- 5. Evaluate fitness of children
- 6. Replace worst parents with the children