# SciencesPo Computational Economics Spring 2017

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# 1 Computational Economics: Unconstrained Optimization

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## 1.1 Some Taxonomy and Initial Examples

- In most of the examples to follow, we talk about *minimization* of a function f. Everything we do also applies to maximization, since  $\min_x f(x) = \max_x -f(x)$ .
- Here is a generic optimization problem:

$$\min_{x \in \mathbb{R}^n} f(x) \text{ s.t. } c_i(x) = 0, \quad i \in E$$
$$c_i(x) \ge 0, \quad i \in I$$

- This is a general way of writing an optimization problem. E are all indices as equality constraints, I are all inequality constraints.
- An example of such a problem might be

$$\min(x_1 - 2)^2 + (x_2 - 1)^2 \text{ s.t. } \begin{array}{l} x_1^2 - x_2 \le 0 \\ x_1 + x_2 \le 2 \end{array}$$

• Here is a picture of that problem taken from the textbook [@nocedal-wright] [2]

# 1.2 Kinds of problems considered

- Don't talk about stochastic optimization methods:
  - Simluated Annealing
  - MCMC
  - other Stochastic Search Methods
  - A gentle introduction is [@casella-R] [3]

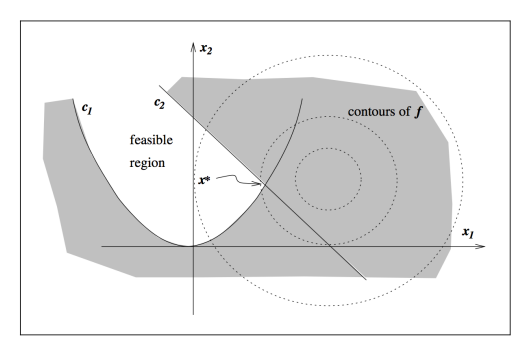


Figure 1.1 in [@nocedal-wright]

# 1.3 Transportation Problem

A chemical company has two factories  $F_1$ ,  $F_2$  and a dozen retail outlets  $R_1$ , ...,  $R_{12}$ . Each factory i can produce at most  $a_i$  tons of output each week. Each retail outlet j has a weekly demand of  $b_j$  tons per week. The cost of shipping from  $F_i$  to  $R_j$  is given by  $c_{ij}$ . How much of the product to ship from each factory to each outlet, minimize cost, and satisfy all constraints? let's call  $x_{ij}$  the number of tons shipped from i to j.

A mathematical formulation of this problem is

$$\min \sum_{i,j} c_{ij} x_{ij}$$
subject to 
$$\sum_{j=1}^{12} x_{ij} \le a_i, \quad i = 1, 2$$

$$\sum_{i=1}^{2} x_{ij} \ge b_j, \quad j = 1, \dots, 12$$

$$x_{ij} \ge 0, \quad i = 1, 2, j = 1, \dots, 12$$

- This is called a *linear programming* problem, because both objective function and all constrains are linear.
- With any of those being nonlinear, we would call this a non-linear problem.

#### 1.4 Constrained vs Unconstrained

• There are many applications of both in economics.

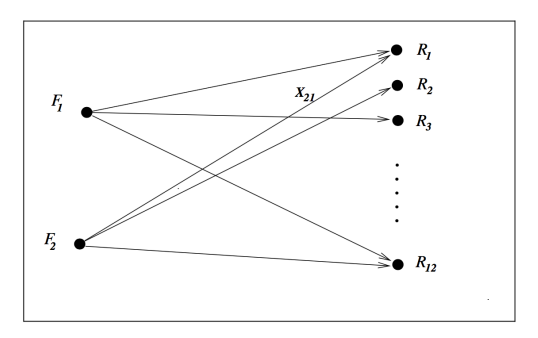


Figure 1.2 in [2]

- Unconstrained: maximimum likelihood
- Constrained: MPEC
- It is sometimes possible to transform a constrained problem into an unconstrained one.

# 1.5 Convexity

- Convex problems are easier to solve.
- What is convex?

A set  $S \in \mathbb{R}^n$  is convex if the straight line segment connecting any two points in S lies entirely inside S. A function f is a convex function, if its domain S is a convex set, and for any two points  $x, y \in S$ , we have that

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

for all  $\alpha \in [0, 1]$ 

- Simple instances of convex sets are the unit ball  $\{y \in \mathbb{R}^n, \|y\|_2 \le 1\}$ , and any set defined by linear equalities and inequalities.
- convex Programming describes a special case of the introductory minimizatin problem where
  - the objective function is convex,
  - the equality constrains are linear, and
  - the inequality constraints are concave.

# 1.6 Optimization Algorithms

- All of the algorithms we are going to see employ some kind of iterative proceedure.
- They try to improve the value of the objective function over successive steps.

- The way the algorithm goes about generating the next step is what distinguishes algorithms from one another.
  - Some algos only use the objective function
  - Some use both objective and gradients
  - Some add the Hessian
  - and many variants more

## 1.7 Desirable Features of any Algorithm

- Robustness: We want good performance on a wide variety of problems in their class, and starting from *all* reasonable starting points.
- Efficiency: They should be fast and not use an excessive amount of memory.
- Accuracy: They should identify the solution with high precision.

#### 1.8 A Word of Caution

- You should **not** normally attempt to write a numerical optimizer for yourself.
- Entire generations of Applied Mathematicians and other numerical pro's have worked on those topics before you, so you should use their work.
  - Any optimizer you could come up with is probably going to perform below par, and be highly likely to contain mistakes.
  - Don't reinvent the wheel.
- That said, it's very important that we understand some basics about the main algorithms, because your task is to choose from the wide array of available ones.

# 2 Unconstrained Optimization: What is a solution?

A typical unconstrained optimization problem will look something like this:

$$\min_{x} f(x), \quad x \in \mathbb{R}^n$$

and where  $f : \mathbb{R}^n \to \mathbb{R}$  is a smooth function.

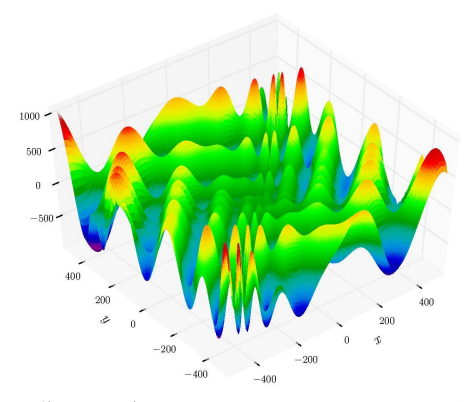
• In general, we would always like to find a *global* minimizer, i.e. a point

$$x^*$$
 where  $f(x^*) \le f(x) \quad \forall x$ 

- Since our algorithm is not going to visit many points in  $\mathbb{R}^n$  (or so we hope), we can never be totally sure that we find a global optimizer.
- Most optimizers can only find a *local* minimizer. That is a point

$$x^*$$
 where  $f(x^*) \le f(x) \quad \forall x \in \mathcal{N}$ 

where  $\mathcal{N}$  is a neighborhood around  $x^*$ .



Global min at f(512,404.2319). By Gaortizg GFDL or CC BY-SA 3.0, via Wikimedia Commons

# 2.1 Global minization can be very hard sometimes.

# 2.2 (Unconstrained) Optimization in Julia

- Umbrella Organisation: http://www.juliaopt.org
  - We will make ample use of this when we talk about constrained optimisation.
  - The Julia Interface to the very well established C-Library NLopt is called NLopt.jl. One could use NLopt without constraints in an unconstrained problem.
- Roots. jl: Simple algorithms that find the zeros of a univariate function.
- Baseline Collection of unconstrained optimization algorithms: Optim.jl

#### 2.3 Introducing Optim.jl

- Multipurpose unconstrained optimization package
  - provides 8 different algorithms with/without derivatives
  - univariate optimization without derivatives

#### 2.4 The Golden Ratio or Bracketing Search for 1D problems

- A derivative-free method
- a Bracketing method
  - find the local minimum of f on [a, b]
  - select 2 interior points c, d such that a < c < d < b

```
* f(c) \le f(d) \implies min must lie in [a,d]. replace b with d, start again with [a,d] * f(c) > f(d) \implies min must lie in [c,b]. replace a with c, start again with [c,b]
```

- how to choose *b*, *d* though?
- we want the length of the interval to be independent of whether we replace upper or lower bound
- we want to reuse the non-replaced point from the previous iteration.
- these imply the golden rule:
- new point  $x_i = a + \alpha_i(b-a)$ , where  $\alpha_1 = \frac{3-\sqrt{5}}{2}$ ,  $\alpha_2 = \frac{\sqrt{5}-1}{2}$
- $-\alpha_2$  is known as the *golden ratio*, well known for it's role in renaissance art.

#### 2.4.1 Bracketing Search in Julia

• The package Optim. jl provides an implementation of "Brent's Method" as well as the golden section search:

```
In [1]: using Plots
        using Optim
        f(x) = \exp(x) - x^4
        minf(x) = -f(x)
        brent = optimize(minf,0,2,Brent())
        golden = optimize(minf,0,2,GoldenSection())
        println("brent = $brent")
        println("golden = $golden")
        plot(f,0,2)
brent = Results of Optimization Algorithm
 * Algorithm: Brent's Method
 * Search Interval: [0.000000, 2.000000]
 * Minimizer: 8.310315e-01
 * Minimum: -1.818739e+00
* Iterations: 12
 * Convergence: max(|x - x_upper|, |x - x_lower|) \le 2*(1.5e-08*|x|+2.2e-16): true
 * Objective Function Calls: 13
golden = Results of Optimization Algorithm
 * Algorithm: Golden Section Search
 * Search Interval: [0.000000, 2.000000]
 * Minimizer: 8.310315e-01
 * Minimum: -1.818739e+00
 * Iterations: 37
 * Convergence: \max(|x - x_{upper}|, |x - x_{lower}|) \le 2*(1.5e-08*|x|+2.2e-16): true
 * Objective Function Calls: 38
In [2]: # how well does this do with many local minima?
        fun(x) = exp(x) - x^4 + sin(40*x)
        minf(x) = -fun(x)
        grid = collect(0:0.000001:2);
        # fun.(grid)
```

```
# v,idx = findmax(fun.(grid))
v,idx = findmax(Float64[fun.(x) for x in grid])
println("grid maximizer is $(grid[idx])")
golden = optimize(minf,0,2,GoldenSection())
brent = optimize(minf,0,2,Brent())
using Base.Test
println("brent minimizer = $(brent.minimizer)")
println("golden minimizer = $(golden.minimizer)")
plot(fun,0,2)

grid maximizer is 0.824692
brent minimizer = 0.8310314519658002
golden minimizer = 0.8310314579502819
```

WARNING: Method definition minf(Any) in module Main at In[1]:4 overwritten at In[2]:3.

#### 2.5 Beyond One Dimension

#### 2.5.1 Introducing Rosenbrock's Banana function

The Banana function is defined by

$$f(x,y) = (a-x)^2 + b(y-x^2)^2$$

#### 2.5.2 What is the minimum of that function?

- For a = 1, b = 100, what is the global minimum of that function?
- What are the inputs one needs to supply to an algorithm in a more general example?

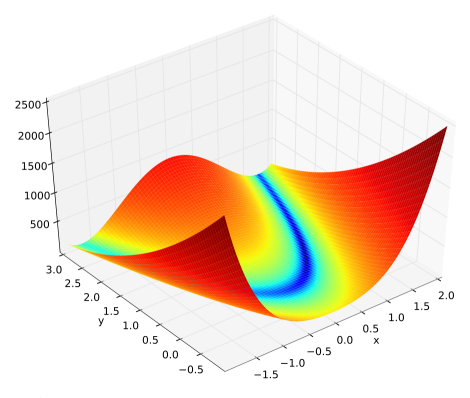
#### 2.6 Rosenbrock Banana and Optim.jl

- We will use Optim for the rest of this lecture.
- We need to supply the objective function and depending on the solution algorithm the gradient and hessian as well.

```
In [3]: using Optim
    rosenbrock = Optim.UnconstrainedProblems.examples["Rosenbrock"]

# contains:
# function rosenbrock(x::Vector)
# return (1.0 - x[1])^2 + 100.0 * (x[2] - x[1]^2)^2
# end

# function rosenbrock_gradient!(x::Vector, storage::Vector)
# storage[1] = -2.0 * (1.0 - x[1]) - 400.0 * (x[2] - x[1]^2) * x[1]
# storage[2] = 200.0 * (x[2] - x[1]^2)
# end
```



Banana for a = 0. By Gaortizg GFDL or CC BY-SA 3.0, via Wikimedia Commons

```
# function rosenbrock_hessian!(x::Vector, storage::Matrix)
# storage[1, 1] = 2.0 - 400.0 * x[2] + 1200.0 * x[1]^2
# storage[1, 2] = -400.0 * x[1]
# storage[2, 1] = -400.0 * x[1]
# storage[2, 2] = 200.0
# end
```

# there are many other examples on Optim. UnconstrainedProblems

Out[3]: Optim.UnconstrainedProblems.OptimizationProblem("Rosenbrock",Optim.UnconstrainedProblems

#### 2.7 Comparison Methods

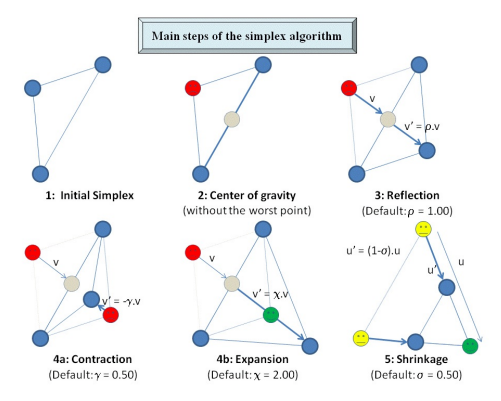
- We will now look at a first class of algorithms, which are very simple, but sometimes a good starting point.
- They just *compare* function values.
- *Grid Search* : Compute the objective function at  $G = \{x_1, \dots, x_N\}$  and pick the highest value of f.
  - This is very slow.
  - It requires large *N*.
  - But it's robust (will find global optimizer for large enough *N*)

```
In [4]: # grid search on rosenbrock
    grid = collect(-1.0:0.1:3);
```

```
grid2D = [[i;j] for i in grid,j in grid];
val2D = map(rosenbrock.f,grid2D);
r = findmin(val2D);
println("grid search results in minimizer = $(grid2D[r[2]])")
grid search results in minimizer = [1.0,1.0]
```

## 2.8 Bracketing for Multidimensional Problems: Nelder-Mead

- The Goal here is to find the simplex containing the local minimizer  $x^*$
- In the case where f is n-D, this simplex has n + 1 vertices
- In the case where f is 2-D, this simplex has 2 + 1 vertices, i.e. it's a triangle.
- The method proceeds by evaluating the function at all n + 1 vertices, and by replacing the worst function value with a new guess.
- this can be achieved by a sequence of moves:
  - reflect
  - expand
  - contract
  - shrink movements.



- this is a very popular method. The matlab functions fmincon and fminsearch implements
- When it works, it works quite fast.
- No derivatives required.

In [5]: optimize(rosenbrock, [0.0, 0.0], NelderMead())

Out[5]: Results of Optimization Algorithm

\* Algorithm: Nelder-Mead \* Starting Point: [0.0,0.0]

\* Minimizer: [0.9999710322210338,0.9999438685860869]

\* Minimum: 1.164323e-09

\* Iterations: 74 \* Convergence: true

\* ((y-)š)/n < 1.0e-08: true

\* Reached Maximum Number of Iterations: false

\* Objective Function Calls: 108

• But.

## 2.9 Bracketing for Multidimensional Problems: Comment on Nelder-Mead

Lagarias et al. (SIOPT, 1999): At present there is no function in any dimension greater than one, for which the original Nelder-Mead algorithm has been proved to converge to a minimizer.

Given all the known inefficiencies and failures of the Nelder-Mead algorithm [...], one might wonder why it is used at all, let alone why it is so extraordinarily popular.

## 2.10 Reminder: Optimality Conditions

#### 2.10.1 Notation

- Unless otherwise noted, we have  $x \in \mathbb{R}^n$  as an n element vector.
- The **gradient** of a function  $f: \mathbb{R}^n \to \mathbb{R}$  is denoted  $\nabla f: \mathbb{R}^n \to \mathbb{R}^n$  a it returns a vector

$$\nabla f(x) = \left(\frac{\partial f}{\partial x_1}(x), \frac{\partial f}{\partial x_2}(x), \dots, \frac{\partial f}{\partial x_n}(x)\right)$$

• It's **hessian** is a function denoted  $\nabla^2 f(x)$  or  $H_f : \mathbb{R}^n \to \mathbb{R}^{n \times n}$  and returns an (n, n) matrix given by

$$H_f(x) = \begin{pmatrix} \frac{\partial^2 f}{\partial x_1 \partial x_1}(x) & \frac{\partial^2 f}{\partial x_2 \partial x_1}(x) & \dots & \frac{\partial^2 f}{\partial x_n \partial x_1}(x) \\ \frac{\partial^2 f}{\partial x_1 \partial x_2}(x) & \frac{\partial^2 f}{\partial x_2 \partial x_2}(x) & \dots & \frac{\partial^2 f}{\partial x_n \partial x_2}(x) \\ \vdots & \vdots & \dots & \vdots \\ \frac{\partial^2 f}{\partial x_1 \partial x_n}(x) & \frac{\partial^2 f}{\partial x_2 \partial x_n}(x) & \dots & \frac{\partial^2 f}{\partial x_n \partial x_n}(x) \end{pmatrix}$$

#### 2.10.2 Optimality Conditions

- **First Order Necessary Conditions**: If f is continously differentiable and  $x^*$  is a local minimizer of f, then  $\nabla f(x^*) = 0$ .
- Second Order Necessary Conditions: If f is twice continuously differentiable and  $x^*$  is a local minimizer of f, then  $\nabla f(x^*) = 0$  and  $H_f(x^*)$  is positive semi-definite, i.e. we have  $s^T H_f(x^*) s \ge 0$  for all  $s \in \mathbb{R}^n$ .

- Second Order Sufficient Conditions: If f is twice continuously differentiable and at  $x^*$  we have that  $\nabla f(x^*) = 0$  and  $H_f(x^*)$  is positive definite, i.e.  $s^T H_f(x^*) s > 0$ ,  $s \neq 0$  then  $x^*$  is a local minimizer of f.
- More sophisticated solvers make ample use of those.

# 2.11 Reminder: Sufficient Conditions for Global Optimality

- If f is convex, then any local minimizer  $x^*$  is a global minimizer.
- If f is convex and differentiable on  $\mathbb{R}^n$ , then any point  $x^*$  is a global minimizer if and only if it is a stationary point, i.e. if  $f \nabla f(x^*) = 0$ .

# 2.12 Reminder: Taylor's Theorem

- Many of the ensuing methods are based on Taylor's theorem, so let's remind ourselves of it:
- Suppose that  $f \in C^{n+1}[a,b]$ , and  $x, x^0 \in [a,b]$ . Then

$$f(x) = f(x^{0}) + f'(x^{0})(x - x^{0}) + f''(x^{0})\frac{(x - x^{0})^{2}}{2} + \dots + f^{(n)}(x^{0})\frac{(x - x^{0})^{n}}{n!} + R_{n+1}(x)$$

where  $R_{n+1}(x) = \ell(\|(x-x^0)\|^{n+1})$  is reminder term that converges at a rate n+1 to zero, i.e. \* we say a function f is  $\ell(\|x\|^n)$  if  $\lim_{x\to 0} \|f(x)\|/\|x\|^n = 0$ . \* we say a function f is  $\mathcal{O}(\|x\|^n)$  if  $\lim_{x\to 0} \|f(x)\|/\|x\|^n < \infty$ 

# 2.13 A quick Note on Computing Derivatives

- Finite Differences
- Automatic Differentiation
- We have talked about this in a separate session.
- For now just remember that if we don't supply analytic gradients, and the algorithm requires them, this often triggers a numerical approximation of the gradient known as finite differences. This is most of the times a slow proceedure.

## 2.14 Two Strategies: Line Search and Trust Region

• We only provide an overview of methods here. If you want to *really* know the details about those algorithms, I invite you to consult [@nocedal-wright].

#### 2.14.1 The Line Search Strategy

- An algorithm from the line search class chooses a direction  $p_k \in \mathbb{R}^n$  and searches along that direction starting from the current iterate  $x_k \in \mathbb{R}^n$  for a new iterate  $x_{k+1} \in \mathbb{R}^n$  with a lower function value.
- After deciding on a direction  $p_k$ , one needs to decide the step length  $\alpha$  to travel by solving

$$\min_{\alpha>0} f(x_k + \alpha p_k)$$

• In practice, solving this exactly is too costly, so algos usually generate a sequence of trial values  $\alpha$  and pick the one with the lowest f.

## 2.14.2 The Trust Region Strategy

- Here we construct a model function  $m_k$  that is similar to f around  $x_k$ .
- We acknowledge that  $m_k$  is decent approximation of f only in some *region*.
- The problem is then to find a candidate step length p by solving

$$\min_{p} m_k(x_k + p)$$

where  $x_k + p$  lies inside the trust region.

- If candidate p does not produce a value lower than  $f(x_k)$ , we must have had a too large trust region, shrink it, and do it again.
- Usually the trust region is a ball  $||p||_2 \le \Delta$ , where  $\Delta$  is called the *trust region radius*, but elliptical and box regions are possible.
- A common definition of the model function is a quadratic of the form

$$m_k(x_k + p) = f(x_k) + p^T \nabla f(x_k) + \frac{1}{2} p^T B(x_k) p$$

where gradient and matrix  $B(x_k)$  are evaluated at the current iterate, so the model function is in agreement to first order with f at the current guess.

• The matrix  $B(x_k)$  is either the Hessian  $H_f$  or some approximation to it.

## 2.15 Trust Region Example

- Suppose we have  $f(x) = 10(x_2 x_1^2)^2 + (1 x_1)^2$ . At point  $x_k = (0, 1)$  gradient and hessian are:
- ?

$$\nabla f(x_k) = \begin{bmatrix} -2 \\ 20 \end{bmatrix}, \qquad \nabla^2 f(x_k) = \begin{bmatrix} -38 & 0 \\ 0 & 20 \end{bmatrix}$$

- In this figure, we use  $B(x_k) = \nabla^2 f(x_k)$ .
- After each unsuccessful step, the new candidate step will be shorter, and will go in a different direction
- This is the main difference to line search methods.
- Main difference between the two methods: order in which they change *direction* and *step length* 
  - line search fixes direction  $p_k$  and finds right distance  $\alpha_k$
  - trust region fixes an appropriate radius  $\Delta_k$

# 2.16 Line Search Methods: Which Direction to go?

#### 2.16.1 Steepest Descent

- The direction  $-\nabla f(x_k)$  is an obvious choice: Among all possible directions, along this one f decreases most rapidly.
- This claim can be verified using Taylor's theorem.

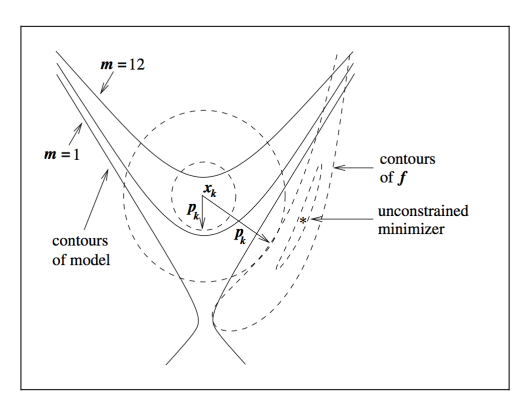


Figure 2.4 of [2]

• The **steepest descent method** is a line search method that moves along

$$p_k = -\nabla f(x_k)$$

• The step length  $\alpha_k$  can be chosen in many different ways.

\* Convergence: false

• There are many other *descent* directions, steepest descent is but one of them.

```
In [6]: # there is a dedicated LineSearch package: https://github.com/JuliaNLSolvers/LineSearche
GD = optimize(rosenbrock.f, rosenbrock.g!,[0.0, 0.0],GradientDescent())
GD1 = optimize(rosenbrock.f, rosenbrock.g!,[0.0, 0.0],GradientDescent(),Optim.Options(it
GD2 = optimize(rosenbrock.f, rosenbrock.g!,[0.0, 0.0],GradientDescent(),Optim.Options(it

println("gradient descent = $GD")
println("\n")
println("gradient descent 2 = $GD1")
println("gradient descent 3 = $GD2")

gradient descent = Results of Optimization Algorithm
* Algorithm: Gradient Descent
* Starting Point: [0.0,0.0]
* Minimizer: [0.9356732500354086,0.875073922357589]
* Minimum: 4.154782e-03
* Iterations: 1000
```

```
* |x - x'| < 1.0e-32: false
```

\* |f(x) - f(x')| / |f(x)| < 1.0e-32: false

\* |g(x)| < 1.0e-08: false

\* f(x) > f(x'): false

\* Reached Maximum Number of Iterations: true

\* Objective Function Calls: 3532

\* Gradient Calls: 3532

## gradient descent 2 = Results of Optimization Algorithm

\* Algorithm: Gradient Descent

\* Starting Point: [0.0,0.0]

\* Minimizer: [0.9978398797724763,0.9956717950747302]

\* Minimum: 4.682073e-06

\* Iterations: 5000

\* Convergence: false

\* |x - x'| < 1.0e-32: false

\* |f(x) - f(x')| / |f(x)| < 1.0e-32: false

\* |g(x)| < 1.0e-08: false

\* f(x) > f(x'): false

\* Reached Maximum Number of Iterations: true

\* Objective Function Calls: 17532

\* Gradient Calls: 17532

#### gradient descent 3 = Results of Optimization Algorithm

\* Algorithm: Gradient Descent

\* Starting Point: [0.0,0.0]

\* Minimizer: [0.9999999914304203,0.9999999828109042]

\* Minimum: 7.368706e-17

\* Iterations: 20458

\* Convergence: true

\* |x - x'| < 1.0e-32: false

\* |f(x) - f(x')| / |f(x)| < 1.0e-32: false

\* |g(x)| < 1.0e-08: true

\* f(x) > f(x'): false

\* Reached Maximum Number of Iterations: false

\* Objective Function Calls: 71635

\* Gradient Calls: 71635

#### 2.17 Line Search Methods: The Newton Direction

- Probably the most important descent direction.
- In vector notation, the 2nd order taylor series approximation to  $f(x_k + p)$  is

$$f(x_k + p) \approx f(x_k) + p^T \nabla f(x_k) + \frac{1}{2} p^T \nabla^2 f(x_k) p \equiv m_k(p)$$

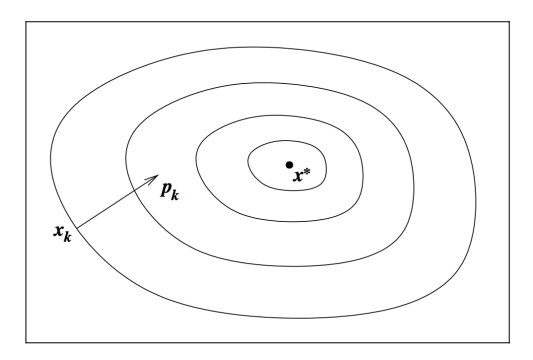


Figure 2.5 of Nocedal-Wright [2]

- the Newton direction is obtained by finding the vector p that minimizes  $m_k(p)$ , i.e. by setting the derivative of  $m_k(p)$  to zero.
- We obtain

$$p_k^N = -(\nabla^2 f(x_k))^{-1} \nabla f(x_k)$$

- The newton direction is reliable if the discrepancy between truth and model m is not too large at  $x_k$ .
- The biggest drawback is the need to compute the Hessian. This can be difficult analytically at times, and overly expensive numerically.

In [7]: optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [0.0, 0.0], Newton(),Optim.Options(), (0.0, 0.0))

Iter	Function value	Gradient norm
0	1.000000e+00	2.000000e+00
1	8.431140e-01	1.588830e+00
2	6.776980e-01	3.453340e+00
3	4.954645e-01	4.862093e+00
4	3.041921e-01	2.590086e+00
5	1.991512e-01	3.780900e+00
6	9.531907e-02	1.299090e+00
7	5.657827e-02	2.445401e+00
8	2.257807e-02	1.839332e+00
9	6.626125e-03	1.314236e+00
10	8.689753e-04	5.438279e-01
11	4.951399e-06	7.814556e-02
12	9.065070e-10	6.017046e-04
13	9.337686e-18	1.059738e-07

Out[7]: Results of Optimization Algorithm

\* Algorithm: Newton's Method

\* Starting Point: [0.0,0.0]

\* Minimizer: [0.9999999999994,0.99999999999999]

\* Minimum: 3.081488e-31

\* Iterations: 14

\* Convergence: true

\* |x - x'| < 1.0e-32: false

\* |f(x) - f(x')| / |f(x)| < 1.0e-32: false

\* |g(x)| < 1.0e-08: true

\* f(x) > f(x'): false

\* Reached Maximum Number of Iterations: false

\* Objective Function Calls: 58

\* Gradient Calls: 58

#### 2.18 Quasi-Newton Methods

- In response to the difficulties of getting the Hessian, quasi-newton methods propose to approximate  $B(x_k)$  with something *similar* to the hessian.
- Taylors Theorem implies that

$$\nabla^2 f(x_{k+1} - x_k) \approx \nabla f(x_{k+1}) - \nabla f(x_k)$$

and so we choose a *B* matrix that mimics this property.

This leads to the secant condition

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

- There are different ways to update the hessian in this way.
- One of the best known is the BFGS method (after Broydon, Fletcher, Goldfarb and Shanno).
- Those methods get the search direction by using  $B_k$  instead of the exact Hessian, i.e.

$$p_k = -B_k^{-1} \nabla f(x_k)$$

In [8]: @show optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [-1.0, 3.0], BFGS());
# low memory BFGS

@show optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [0.0, 0.0], LBFGS());

optimize(rosenbrock.f,rosenbrock.g!,rosenbrock.h!,[-1.0,3.0],BFGS()) = Results of Optimization A

- \* Algorithm: BFGS
- \* Starting Point: [-1.0,3.0]
- \* Minimizer: [0.9999999999956,0.999999999997]
- \* Minimum: 1.707144e-27
- \* Iterations: 39
- \* Convergence: true
  - \* |x x'| < 1.0e-32: false
  - \* |f(x) f(x')| / |f(x)| < 1.0e-32: false

```
* |g(x)| < 1.0e-08: true
  * f(x) > f(x'): false
  * Reached Maximum Number of Iterations: false
* Objective Function Calls: 176
* Gradient Calls: 176
optimize(rosenbrock.f,rosenbrock.g!,rosenbrock.h!,[0.0,0.0],LBFGS()) = Results of Optimization A
* Algorithm: L-BFGS
* Starting Point: [0.0,0.0]
* Minimizer: [1.00000000000007,1.00000000000001]
* Minimum: 5.374115e-30
* Iterations: 21
* Convergence: true
  * |x - x'| < 1.0e-32: false
  * |f(x) - f(x')| / |f(x)| < 1.0e-32: false
  * |g(x)| < 1.0e-08: true
  * f(x) > f(x'): false
  * Reached Maximum Number of Iterations: false
* Objective Function Calls: 90
* Gradient Calls: 90
```

#### 2.19 Practical Considerations

# 2.19.1 Stopping criteria

- In all of the above examples, we did not alter the default values for stopping criteria.
- There are different things you could focus on as a stopping criterion with Optim, and similarly in most solver packages.
  - xtol: changes in *x* from one iterate to the next
  - f\_tol: perentage changes in *f* from one iterate to the next
  - g\_tol: absolute value of the gradient beign smaller than this number.

#### 2.20 Some Applications in Economics

- Maximum Likelihood Estimation
- The Nested Fixed Point Algorithm
  - [@rust-bus] [4] is a maximum likelihood estimation with an inner loop that solves a dynamic programming problem.
  - [@BLP] [1] is a GMM estimation with an inner fixed point problem.

# References

- [1] S. Berry, J. Levinsohn, and A. Pakes. Automobile prices in market equilibrium. *Econometrica: Journal of the Econometric Society*, pages 841–890, 1995.
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- [3] Christian Robert and George Casella. *Introducing Monte Carlo Methods with R*. Springer Science & Business Media, 2009.
- [4] John Rust. Optimal replacement of gmc bus engines: An empirical model of harold zurcher. *Econometrica*, 55(5):999–1033, 1987.