# SciencesPo Computational Economics Spring 2017

Florian Oswald

January 17, 2019

# 1 Optimization 2: Algorithms and Constraints

Florian Oswald Sciences Po, 2018

### 1.1 Bracketing

- A derivative-free method for *univariate f*
- works only on **unimodal** *f*
- (Draw choosing initial points and where to move next)

### 1.2 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
    - \*  $f(c) \le f(d) \implies \min \max \text{ lie in } [a,d]. \text{ replace } b \text{ with } d, \text{ start again with } [a,d]$
    - \*  $f(c) > f(d) \implies$  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.

```
In [1]: using Plots
    using Optim
    plotlyjs()
    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_{pper}|, |x - x_{per}|) \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_{pper}|, |x - x_{lower}|) \le 2*(1.5e-08*|x|+2.2e-16): true
 * Objective Function Calls: 38
1.2.1 Bisection Methods
  • Root finding: Roots.jl
  • Root finding in multivariate functions: IntervalRootFinding.jl
In [2]: using Roots
        #ăfind the zeros of this function:
        f(x) = \exp(x) - x^4
        ## bracketing
        fzero(f, 8, 9) # 8.613169456441398
        fzero(f, -10, 0) # -0.8155534188089606
Out[2]: -0.8155534188089606
In [3]: using IntervalRootFinding, IntervalArithmetic
        -10..10
        ArgumentError: Module IntervalRootFinding not found in current path.
    Run `Pkg.add("IntervalRootFinding")` to install the IntervalRootFinding package.
        Stacktrace:
         [1] _require(::Symbol) at ./loading.jl:435
```

```
[2] require(::Symbol) at ./loading.jl:405
         [3] include_string(::String, ::String) at ./loading.jl:522
In [4]: X = IntervalBox(1...3, 2...4)
        UndefVarError: IntervalBox not defined
        Stacktrace:
         [1] include_string(::String, ::String) at ./loading.jl:522
In [5]: a = @interval(0.1, 0.3)
        b = 0interval(0.3, 0.6)
        a + b
        UndefVarError: @interval not defined
        Stacktrace:
         [1] include_string(::String, ::String) at ./loading.jl:522
In [6]: rts = IntervalRootFinding.roots(x->x^2 - 2, -10..10, Bisection)
        UndefVarError: IntervalRootFinding not defined
        Stacktrace:
         [1] include_string(::String, ::String) at ./loading.jl:522
```

### 1.3 Rosenbrock Banana and Optim.jl

• We can supply the objective function and - depending on the solution algorithm - the gradient and hessian as well.

```
In [7]: using Optim
        using OptimTestProblems
        for (name, prob) in MultivariateProblems.UnconstrainedProblems.examples
           println(name)
        end
Rosenbrock
Quadratic Diagonal
Hosaki
Large Polynomial
Penalty Function I
Beale
Extended Rosenbrock
Polynomial
Powell
Exponential
Paraboloid Diagonal
Paraboloid Random Matrix
Extended Powell
Trigonometric
Fletcher-Powell
Parabola
Himmelblau
In [8]: rosenbrock = MultivariateProblems.UnconstrainedProblems.examples["Rosenbrock"]
Out[8]: OptimTestProblems.MultivariateProblems.OptimizationProblem{Void, Void, Float64, String, Vo
1.4 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
       - This is very slow.
       - It requires large N.
       – But it's robust (will find global optimizer for large enough N)
In [9]: # 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]

#### 1.5 Local Descent Methods

- Applicable to multivariate problems
- We are searching for a *local model* that provides some guidance in a certain region of *f* over **where to go to next**.
- Gradient and Hessian are informative about this.

#### 1.5.1 Local Descent Outline

All descent methods follow more or less this structure. At iteration *k*,

- 1. Check if candidate  $\mathbf{x}^{(k)}$  satisfies stopping criterion:
  - if yes: stop
  - if no: continue
- 2. Get the local *descent direction*  $\mathbf{d}^{(k)}$ , using gradient, hessian, or both.
- 3. Set the *step size*, i.e. the length of the next step,  $\alpha^k$
- 4. Get the next candidate via

$$\mathbf{x}^{(k+1)} \longleftarrow \alpha^k \mathbf{d}^{(k)}$$

#### 1.5.2 The Line Search Strategy

- An algorithm from the line search class chooses a direction  $\mathbf{d}^{(k)} \in \mathbb{R}^n$  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  $\mathbf{d}^{(k)}$ , one needs to decide the *step length*  $\alpha$  to travel by solving

$$\min_{\alpha>0} f(x_k + \alpha \mathbf{d}^{(k)})$$

• 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.

```
* |g(x)| 1.0e-08: true |g(x)| = 6.66e-15
```

\* Stopped by an increasing objective: false

\* Reached Maximum Number of Iterations: false

\* Objective Calls: 71 \* Gradient Calls: 71

\* Hessian Calls: 23

#### 1.5.3 The Trust Region Strategy

- First choose max step size, then the direction
- Finds the next step  $\mathbf{x}^{(k+1)}$  by minimizing a model of  $\hat{f}$  over a *trust region*, centered on  $\mathbf{x}^{(k)}$ 
  - 2nd order Tayloer approx of f is common.
- Radius  $\delta$  of trust region is changed based on how well  $\hat{f}$  fits f in trust region.
- Get x' via

$$\label{eq:linear_problem} \begin{aligned} \min_{\mathbf{x}'} \quad \hat{f}(\mathbf{x}') \\ \text{subject to} \quad \|\mathbf{x} - \mathbf{x}' \leq \delta\| \end{aligned}$$

```
In [11]: # Optim.jl has a TrustRegion for Newton (see below for Newton's Method)
         NewtonTrustRegion(; initial_delta = 1.0, # The starting trust region radius
                             delta_hat = 100.0, # The largest allowable trust region radius
                             eta = 0.1, #When rho is at least eta, accept the step.
                             rho_lower = 0.25, # When rho is less than rho_lower, shrink the t
                             rho_upper = 0.75) # When rho is greater than rho_upper, grow the
         res = Optim.optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, rosenbrock.initial_x
Out[11]: Results of Optimization Algorithm
          * Algorithm: Newton's Method (Trust Region)
          * Starting Point: [-1.2,1.0]
          * Minimizer: [0.999999994405535,0.9999999988644926]
          * Minimum: 3.405841e-19
          * Iterations: 25
          * Convergence: true
            * |x - x'| 1.0e-32: false
              |x - x'| = 8.84e-06
            * |f(x) - f(x')| 1.0e-32 |f(x)|: false
              |f(x) - f(x')| = 1.87e + 08 |f(x)|
            * |g(x)| 1.0e-08: true
              |g(x)| = 5.53e-09
            * Stopped by an increasing objective: false
            * Reached Maximum Number of Iterations: false
          * Objective Calls: 26
          * Gradient Calls: 26
```

#### 1.5.4 Stopping criteria

1. maximum number of iterations reached

\* Hessian Calls: 22

- 2. absolute improvement  $|f(x) f(x')| \le \epsilon$
- 3. relative improvement  $|f(x) f(x')|/|f(x)| \le \epsilon$
- 4. Gradient close to zero  $|g(x)| \approx 0$

#### 1.5.5 Gradient Descent

Here we define

$$\mathbf{g}^{(k)} = \nabla f(\mathbf{d}^{(k)})$$

And our descent becomes

$$\mathbf{d}^{(k)} = -\nabla \frac{\mathbf{g}^{(k)}}{\|\mathbf{g}^{(k)}\|}$$

 Minimizing wrt step size results in a jagged path (each direction is orthogonal to previous direction!)

$$\alpha^{(k)} = \arg\min\alpha f(\mathbf{x}^{(k)} + \alpha \mathbf{d}^{(k)})$$

• Conjugate Gradient avoids this issue.

```
In [12]: # Optim.jl again
         GradientDescent(; alphaguess = LineSearches.InitialPrevious(),
                            linesearch = LineSearches.HagerZhang(),
                            P = nothing,
                            precondprep = (P, x) -> nothing)
Out[12]: Optim.GradientDescent{LineSearches.InitialPrevious{Float64},LineSearches.HagerZhang{Float64},
           alpha: Float64 1.0
           alphamin: Float64 0.0
           alphamax: Float64 Inf
         , LineSearches.HagerZhang{Float64}
           delta: Float64 0.1
           sigma: Float64 0.9
           alphamax: Float64 Inf
           rho: Float64 5.0
           epsilon: Float64 1.0e-6
           gamma: Float64 0.66
           linesearchmax: Int64 50
           psi3: Float64 0.1
           display: Int64 0
```

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

, nothing, #5, Optim.Flat())

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
 * Convergence: false
   * |x - x'| 1.0e-32: false
     |x - x'| = 1.82e-04
   * |f(x) - f(x')| 1.0e-32 |f(x)|: false
     |f(x) - f(x')| = 1.97e-03 |f(x)|
   * |g(x)| 1.0e-08: false
     |g(x)| = 8.21e-02
   * Stopped by an increasing objective: false
   * Reached Maximum Number of Iterations: true
 * Objective Calls: 2532
 * Gradient Calls: 2532
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
     |x - x'| = 5.08e-06
   * |f(x) - f(x')| 1.0e-32 |f(x)|: false
     |f(x) - f(x')| = 1.62e-03 |f(x)|
  * |g(x)| 1.0e-08: false
     |g(x)| = 2.53e-03
   * Stopped by an increasing objective: false
   * Reached Maximum Number of Iterations: true
 * Objective Calls: 12532
 * Gradient Calls: 12532
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
     |x - x'| = 2.00e-11
  * |f(x) - f(x')| 1.0e-32 |f(x)|: false
```

|f(x) - f(x')| = 1.61e-03 |f(x)|

\* |g(x)| 1.0e-08: true |g(x)| = 9.99e-09

\* Stopped by an increasing objective: false

\* Reached Maximum Number of Iterations: false

\* Objective Calls: 51177 \* Gradient Calls: 51177

#### 1.6 Second Order Methods

#### 1.6.1 Newton's Method

• We start with a 2nd order Taylor approx over x at step *k*:

$$q(x) = f(x^{(k)}) + (x - x^{(k)})f'(x^{(k)}) + \frac{(x - x^{(k)})^2}{2}f''(x^{(k)})$$

• We set find it's root and rearrange to find the next step k + 1:

$$\frac{\partial q(x)}{\partial x} = f'(x^{(k)}) + (x - x^{(k)})f''(x^{(k)}) = 0$$
$$x^{(k+1)} = x^{(k)} - \frac{f'(x^{(k)})}{f''(x^{(k)})}$$

- The same argument works for multidimensional functions by using Hessian and Gradient
- We would get a descent  $\mathbf{d}^k$  by setting:

$$\mathbf{d}^k = -\frac{\mathbf{g}^k}{\mathbf{H}^k}$$

- There are several options to avoid (often costly) computation of the Hessian H:
- 1. Quasi-Newton updates **H** starting from identity matrix
- 2. Broyden-Fletcher-Goldfarb-Shanno (BFGS) does better with approx linesearch
- 3. L-BFGS is the limited memory version for large problems

In [14]: optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [0.0, 0.0], Newton(),Optim.Option

rm
+00
+00
+00
+00
+00
+00
+00
+00
+00
+00
-01

```
7.814556e-02
    11
         4.951399e-06
         9.065070e-10
                          6.017046e-04
    12
         9.337686e-18 1.059738e-07
3.081488e-31 1.110223e-15
    13
    14
Out[14]: 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
              |x - x'| = 3.06e-09
            * |f(x) - f(x')| 1.0e-32 |f(x)|: false
              |f(x) - f(x')| = 3.03e+13 |f(x)|
            * |g(x)| 1.0e-08: true
              |g(x)| = 1.11e-15
            * Stopped by an increasing objective: false
            * Reached Maximum Number of Iterations: false
          * Objective Calls: 44
          * Gradient Calls: 44
          * Hessian Calls: 14
In [15]: @show optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [-1.0, 3.0], BFGS());
optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [-1.0, 3.0], BFGS()) = Results of Optimize
 * 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
     |x - x'| = 1.54e-08
   * |f(x) - f(x')| 1.0e-32 |f(x)|: false
     |f(x) - f(x')| = 3.55e+10 |f(x)|
  * |g(x)| 1.0e-08: true
     |g(x)| = 1.63e-12
   * Stopped by an increasing objective: false
   * Reached Maximum Number of Iterations: false
 * Objective Calls: 137
 * Gradient Calls: 137
In [16]: # low memory BFGS
         @show optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [0.0, 0.0], LBFGS());
```

```
optimize(rosenbrock.f, rosenbrock.g!, rosenbrock.h!, [0.0, 0.0], LBFGS()) = Results of Optimize
 * Algorithm: L-BFGS
 * Starting Point: [0.0,0.0]
 * Minimizer: [0.9999999999928,0.999999999998559]
 * Minimum: 5.191703e-27
 * Iterations: 24
 * Convergence: true
   * |x - x'| 1.0e-32: false
     |x - x'| = 4.58e-11
   * |f(x) - f(x')| 1.0e-32 |f(x)|: false
     |f(x) - f(x')| = 8.50e+07 |f(x)|
   * |g(x)| 1.0e-08: true
     |g(x)| = 1.44e-13
   * Stopped by an increasing objective: false
   * Reached Maximum Number of Iterations: false
 * Objective Calls: 67
 * Gradient Calls: 67
```

#### ## Direct Methods

- No derivative information is used *derivative free*
- If it's very hard / impossible to provide gradient information, this is our only chance.
- Direct methods use other criteria than the gradient to inform the next step (and ulimtately convergence).

#### 1.6.2 Cyclic Coordinate Descent – Taxicab search

- We do a line search over each dimension, one after the other
- taxicab because the path looks like a NYC taxi changing direction at each block.
- given  $\mathbf{x}^{(1)}$ , we proceed

$$\mathbf{x}^{(2)} = \arg\min_{x_1} f(x_1, x_2^{(1)}, \dots, x_n^{(1)})$$
  
$$\mathbf{x}^{(3)} = \arg\min_{x_2} f(x_1^{(2)}, x_2, x_3^{(2)}, \dots, x_n^{(2)})$$

• unfortunately this can easily get stuck because it can only move in 2 directions.

```
return x end
```

Out[17]: cyclic\_coordinate\_descent (generic function with 1 method)

#### 1.6.3 General Pattern Search

- We search according to an arbitrary pattern P of candidate points, anchored at current guess
   x.
- With step size  $\alpha$  and set  $\mathcal{D}$  of directions

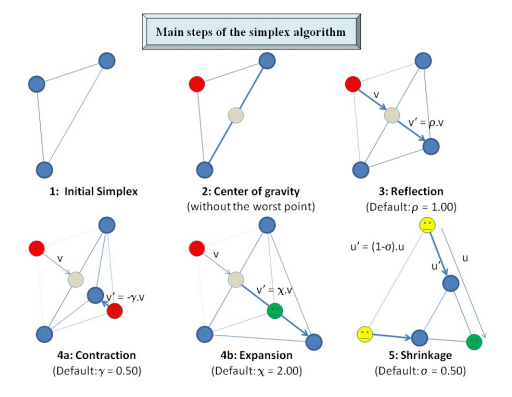
$$\mathcal{P} = \mathbf{x} + \alpha \mathbf{d}$$
 for  $\mathbf{d} \in \mathcal{D}$ 

- Convergence is guaranteed under conditions:
  - $\mathcal{D}$  must be a positive spanning set: at least one  $\mathbf{d} \in \mathcal{D}$  has a non-zero gradient.

```
In [18]: function generalized_pattern_search(f, x, , D, , =0.5)
             y, n = f(x), length(x)
             evals = 0
             while >
                 improved = false
                 for (i,d) in enumerate(D)
                     x = x + *d
                     y = f(x)
                     evals += 1
                     if y < y
                         x, y, improved = x, y, true
                         D = unshift!(deleteat!(D, i), d)
                         break
                     end
                 end
                 if !improved
                      *=
                 end
             end
             println("$evals evaluations")
             return x
         end
Out[18]: generalized_pattern_search (generic function with 2 methods)
In [19]: D = [[1,0],[0,1],[-1,-0.5]]
         D = [[1,0],[0,1]]
         y=generalized_pattern_search(rosenbrock.f,zeros(2),0.8,D,1e-6)
11923 evaluations
Out[19]: 2-element Array{Float64,1}:
          0.999673
          0.999347
```

### 1.7 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 it.
- When it works, it works quite fast.
- No derivatives required.

• But.

### 1.8 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.

### 1.9 things to read up on

- Divided Rectangles (DIRECT)
- simulated annealing and other stochastic gradient methods

### 1.10 Stochastic Optimization Methods

- Gradient based methods like steepest descent may be susceptible to getting stuck at local minima.
- Randomly shocking the value of the descent direction may be a solution to this.
- For example, one could modify our gradient descent from before to become

$$\mathbf{x}^{(k+1)} \longleftarrow \mathbf{x}^{(k)} + \alpha^k \mathbf{g}^{(k)} + \mathbf{v}^{(k)}$$

- where  $\mathbf{v}^{(k)} \sim N(0, \sigma_k^2)$ , decreasing with k.
- This stochastic gradient descent is often used when training neural networks.

#### 1.10.1 Simulated Annealing

- We specify a temperature that controls the degree of randomness.
- At first the temperature is high, letting the search jump around widely. This is to escape local minima.
- The temperature is gradually decreased, reducing the step sizes. This is to find the local optimimum in the *best* region.
- At every iteration k, we accept new point  $\mathbf{x}'$  with

$$\Pr(\operatorname{accept} \mathbf{x}') = \begin{cases} 1 & \text{if } \Delta y \le 0\\ \min(e^{\Delta y/t}, 1) & \text{if } \Delta y > 0 \end{cases}$$

- here  $\Delta y = f(\mathbf{x}') f(\mathbf{x})$ , and t is the temperature.
- Pr(accept x') is called the **Metropolis Criterion**, building block of *Accept/Reject* algorithms.

```
x_best, y_best = x, y
             for k in 1 : k_max
                 x = x + rand(T)
                 y = f(x)
                 y = y - y
                 if y = 0 \mid | rand() < exp(-y/t(k))
                     x, y = x, y
                 end
                 if y < y_best</pre>
                     x_best, y_best = x, y
                 end
                 ytrace[k] = y_best
             return x_best,ytrace
         end
Out[21]: simulated_annealing (generic function with 1 method)
In [22]: function ackley(x, a=20, b=0.2, c=2)
             d = length(x)
             return -a*exp(-b*sqrt(sum(x.^2)/d)) - exp(sum(cos.(c*xi) for xi in x)/d) + a + e
         end
         using Plots
         plotlyjs()
         surface(-30:0.1:30,-30:0.1:30,(x,y)->ackley([x, y]))
In [23]: using Distributions
         d = Dict()
         for sig in (1,5,25), t1 in (1,10,25)
             tmp = [simulated_annealing(ackley,[15,15],MvNormal(2,sig),x->t1/x,100) for i in 1
             d[(sig,t1)] = Dict()
             d[(sig,t1)][:y] = mapslices(x->ackley(x),hcat([tmp[i][1] for i in 1:300]...),[1])
             d[(sig,t1)][:ytrace] = hcat([tmp[i][2] for i in 1:300]...)
         end
         d
         \# x = [simulated\_annealing(ackley, [15, 15], MvNormal(2, 1), x -> 1.0/x, 100) for i in 1:100]
         \# y = [simulated\_annealing(ackley, [15,15], MvNormal(2,5), x->10.0/x, 100) for i in 1:100]
         # map((x)->ackley([x[1],x[2]]),y)
Out[23]: Dict{Any,Any} with 9 entries:
           (5, 25) => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (5, 10) => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (1, 25) => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (5, 1)
                    => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (25, 1) => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (25, 25) \Rightarrow Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 19.0043 ])
                    => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (1, 1)
           (1, 10) => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
           (25, 10) => Dict{Any,Any}(Pair{Any,Any}(:ytrace, [19.0043 19.0043 19.0043 1
```

### 2 Constraints

Recall our core optimization problem:

$$\min_{x \in \mathbb{R}^n} f(x) \text{ s.t. } x \in \mathcal{X}$$

- Up to now, the feasible set was  $\mathcal{X} \in \mathbb{R}^n$ .
- In **constrained problems**  $\mathcal{X}$  is a subset thereof.
- We already encountered *box constraints*, e.g.  $x \in [a, b]$ .
- Sometimes the contrained solution coincides with the unconstrained one, sometimes it does not.
- There are equality constraints and inequality constraints.

### 2.1 Lagrange Multipliers

• Used to optimize a function subject to equality constraints.

$$\min_{x} f(x)$$
 subject to  $h(x) = 0$ 

where both f and h have continuous partial derivatives.

• We look for contour lines of f that are aligned to contours of h(x) = 0.

In other words, we want to find the best x s.t. h(x) = 0 and we have

$$\nabla f(x) = \lambda \nabla h(x)$$

for some *Lagrange Mutliplier*  $\lambda$  \* Notice that we need the scalar  $\lambda$  because the magnitudes of the gradients may be different. \* We therefore form the the **Lagrangian**:

$$\mathcal{L}(x,\lambda) = f(x) - \lambda h(x)$$

#### 2.1.1 Example

Suppose we have

$$\min_{x} - \exp\left(-\left(x_1 x_2 - \frac{3}{2}\right)^2 - \left(x_2 - \frac{3}{2}\right)^2\right)$$
subject to  $x_1 - x_2^2 = 0$ 

We form the Lagrangiagn:

$$\mathcal{L}(x_1, x_2, \lambda) = -\exp\left(-\left(x_1 x_2 - \frac{3}{2}\right)^2 - \left(x_2 - \frac{3}{2}\right)^2\right) - \lambda(x_1 - x_2^2)$$

Then we compute the gradient wrt to  $x_1, x_2, \lambda$ , set to zero and solve.

```
In [35]: gr()
    f(x1,x2) = -exp.(-(x1.*x2 - 3/2).^2 - (x2-3/2).^2)
    c(x1) = sqrt(x1)
    x=0:0.01:3.5
    contour(x,x,(x,y)->f(x,y),lw=1.5,levels=[collect(0:-0.1:-0.85)...,-0.887,-0.95,-1])
    plot!(c,0.01,3.5,label="",lw=2,color=:black)
    scatter!([1.358],[1.165],markersize=5,markercolor=:red,label="Constr. Optimum")
```

• If we had multiple constraints (l), we'd just add them up to get

$$\mathcal{L}(\mathbf{x}, \check{}) = f(\mathbf{x}) - \sum_{i=1}^{l} \lambda_i h_i(\mathbf{x})$$

### 2.2 Inequality Constraints

Suppose now we had

$$\min_{\mathbf{x}} f(\mathbf{x})$$
subject to  $g(\mathbf{x}) \le 0$ 

which, if the solution lies right on the constraint boundary, means that

$$\nabla f - \mu \nabla g = 0$$

for some scalar  $\mu$  - as before.

- In this case, we say the **constraint is active**.
- In the opposite case, i.e. the solution lies **inside** the contrained region, we way the contraint is **inactive**.
- In that case, we are back to an *unconstrained* problem, look for  $\nabla f = 0$ , and set  $\mu = 0$ .

scatter!([1],[1.5],markersize=5,markercolor=:red,label="Unconstr. Optimum")

### 2.3 Infinity Step

• We could do an **infinite step** to avoid *infeasible points*:

$$f_{\infty\text{-step}} = \begin{cases} f(\mathbf{x}) & \text{if } g(\mathbf{x}) \le 0 \\ \infty & \text{else.} \end{cases}$$
$$= f(\mathbf{x}) + \infty(g(\mathbf{x} > 0))$$

- Unfortunately, this is discontinous and non-differentiable, i.e. hard to handle for algorithms.
- Instead, we use a *linear penalty*  $\mu g(\mathbf{x})$  on the objective if the constraint is violated.
- The penalty provides a lower bound to ∞:

$$\mathcal{L}(\mathbf{x}, \mu) = f(\mathbf{x}) + \mu g(\mathbf{x})$$

• We can get back the infinite step by maximizing the penalty:

$$f_{\infty ext{-step}} = \max_{\mu \geq 0} \mathcal{L}(\mathbf{x}, \mu)$$

• Every infeasible **x** returns  $\infty$ , all others return  $f(\mathbf{x})$ 

#### 2.4 Kuhn-Karush-Tucker (KKT)

• Our problem thus becomes

$$\min_{\mathbf{x}} \max_{\mu \geq 0} \mathcal{L}(\mathbf{x}, \mu)$$

- This is called the **primal problem**. Optimizing this requires:
- 1.  $g(\mathbf{x}^*) \leq 0$ . Point is feasible.
- 2.  $\mu \ge 0$ . Penalty goes into the right direction. *Dual feasibility*.
- 3.  $\mu g(\mathbf{x}^*) = 0$ . Feasible point on the boundary has  $g(\mathbf{x}) = 0$ , otherwise  $g(\mathbf{x}) < 0$  and  $\mu = 0$ .
- 4.  $\nabla f(\mathbf{x}^*) \mu \nabla g(\mathbf{x}^*) = 0$ . With an active constraint, we want parallel contours of objective and constraint. When inactive, our optimum just has  $\nabla f(\mathbf{x}^*) = 0$ , which means  $\mu = 0$ .

The preceding four conditions are called the Kuhn-Karush-Tucker (KKT) conditions. In the above order, and in general terms, they are:

- 1. Feasibility
- 2. Dual Feasibility
- 3. Complementary Slackness
- 4. Stationarity.

The KKT conditions are the FONCs for problems with smooth constraints.

### 2.5 Duality

We can combine equality and inequality constraints:

$$\mathcal{L}(\mathbf{x}, \ddot{\mathbf{x}}, \ddot{\mathbf{x}}) = f(\mathbf{x}) + \sum_{i} \lambda_{i} h_{i}(\mathbf{x}) + \sum_{j} \mu_{j} g_{j}(\mathbf{x})$$

where, notice, we reverted the sign of  $\lambda$  since this is unrestricted.

• The Primal problem is identical to the original problem and just as difficult to solve:

$$\min_{\mathbf{x}} \max_{\bar{z} \geq 0, \bar{z}} \mathcal{L}(\mathbf{x}, \bar{z}, \bar{z})$$

• The Dual problem reverses min and max:

$$\max_{\bar{z} \geq 0, \tilde{y}} \min_{\mathbf{x}} \mathcal{L}(\mathbf{x}, \bar{z}, \tilde{y})$$

#### 2.5.1 Dual Values

• The *max-min-inequality* states that for any function f(a, b)

$$\max_{\mathbf{a}} \min_{\mathbf{b}} f(\mathbf{a}, \mathbf{b}) \le \min_{\mathbf{b}} \max_{\mathbf{a}} f(\mathbf{a}, \mathbf{b})$$

- Hence, the solution to the dual is a lower bound to the solution of the primal problem.
- The solution to the *dual function*,  $\min_{\mathbf{x}} \mathcal{L}(\mathbf{x}, \vec{\ }, \vec{\ })$  is the min of a collection of linear functions, and thus always concave.
- It is easy to optimize this.
- In general, solving the dual is easy whenever minimizing  $\mathcal{L}$  wrt x is easy.

## Penalty Methods

- We can convert the constrained problem back to unconstrained by adding penalty terms for constraint violoations.
- A simple method could just count the number of violations:

$$p_{\text{count}}(\mathbf{x}) = \sum_{i} (h_i(\mathbf{x}) \neq 0) + \sum_{i} (g_i(\mathbf{x} > 0))$$

• and add this to the objective in an *unconstrained* problem with penalty  $\rho > 0$ 

$$\min_{\mathbf{x}} f(\mathbf{x}) + \rho p_{\text{count}}(\mathbf{x})$$

- One can choose the penalty function: for example, a quadratic penalty will produce a smooth objective function
- Notice that  $\rho$  needs to become very large sometimes here.

### ## Augmented Lagrange Method

• This is very similar, but specific to equality constraints.

#### ## Interior Point Method

- Also called *barrier method*.
- These methods make sure that the search point remains always feasible.
- As one approaches the constraint boundary, the barrier function goes to infinity. Properties:
- 1.  $p_{\text{barrier}}(\mathbf{x})$  is continuous
- 2.  $p_{\text{barrier}}(\mathbf{x})$  is non negative
- 3.  $p_{\text{barrier}}(\mathbf{x})$  goes to infinitely as one approaches the constraint boundary

#### 2.5.2 Barriers

• Inverse Barrier

$$p_{\text{barrier}}(\mathbf{x}) = -\sum_{i} \frac{1}{g_i(\mathbf{x})}$$

• Log Barrier

$$p_{\text{barrier}}(\mathbf{x}) = -\sum_{i} \begin{cases} \log(-g_i(\mathbf{x})) & \text{if } g_i(\mathbf{x}) \geq -1 \\ 0 & \text{else.} \end{cases}$$

• The approach is as before, one transforms the problem to an unconstrained one and increases  $\rho$  until convergence:

$$\min_{\mathbf{x}} f(\mathbf{x}) + \frac{1}{\rho} p_{\text{barrier}}(\mathbf{x})$$

#### 2.5.3 Examples

$$\min_{x \in \mathbb{R}^2} \sqrt{x_2} \text{ subject to } \begin{cases} x_2 \ge 0 \\ x_2 \ge (a_1 x_1 + b_1)^3 \\ x_2 \ge (a_2 x_1 + b_2)^3 \end{cases}$$

## 2.6 Constrained Optimisation with NLopt.jl

- We need to specify one function for each objective and constraint.
- Both of those functions need to compute the function value (i.e. objective or constraint) *and* it's respective gradient.
- NLopt expects contraints always to be formulated in the format

$$g(x) \leq 0$$

where *g* is your constraint function

- The constraint function is formulated for each constraint at x. it returns a number (the value of the constraint at x), and it fills out the gradient vector, which is the partial derivative of the current constraint wrt x.
- There is also the option to have vector valued constraints, see the documentation.
- We set this up as follows:

```
In [27]: using NLopt
         count = 0 # keep track of # function evaluations
         function myfunc(x::Vector, grad::Vector)
             if length(grad) > 0
                 grad[1] = 0
                 grad[2] = 0.5/sqrt(x[2])
             end
             global count
             count::Int += 1
             println("f_$count($x)")
             sqrt(x[2])
         end
         function myconstraint(x::Vector, grad::Vector, a, b)
             if length(grad) > 0
                 grad[1] = 3a * (a*x[1] + b)^2
                 grad[2] = -1
             end
             (a*x[1] + b)^3 - x[2]
         end
         opt = Opt(:LD_MMA, 2)
         lower_bounds!(opt, [-Inf, 0.])
         xtol_rel!(opt,1e-4)
         min_objective!(opt, myfunc)
         inequality_constraint!(opt, (x,g) -> myconstraint(x,g,2,0), 1e-8)
         inequality_constraint!(opt, (x,g) \rightarrow myconstraint(x,g,-1,1), 1e-8)
         (minfunc, minx, ret) = NLopt.optimize(opt, [1.234, 5.678])
         println("got $minfunc at $minx after $count iterations (returned $ret)")
f_1([1.234, 5.678])
f_2([0.878739, 5.55137])
f_3([0.826216, 5.0439])
f_4([0.473944, 4.07677])
f_5([0.353898, 3.03085])
f_6([0.333873, 1.97179])
```

```
f_7([0.333334, 1.04509])
f_8([0.333334, 0.469503])
f_9([0.333333, 0.305792])
f_10([0.333333, 0.296322])
f_11([0.333333, 0.296296])
got 0.5443310477213124 at [0.333333, 0.296296] after 11 iterations (returned XTOL_REACHED)
```

WARNING: using NLopt.optimize in module Main conflicts with an existing identifier.

### 2.7 NLopt: Rosenbrock

- Let's tackle the rosenbrock example again.
- To make it more interesting, let's add an inequality constraint.

$$\min_{x \in \mathbb{R}^2} (1 - x_1)^2 + 100(x_2 - x_1^2)^2 \text{ subject to } 0.8 - x_1^2 - x_2^2 \ge 0$$

• in NLopt format, the constraint is  $x_1 + x_2 - 0.8 \le 0$ 

```
In [28]: function rosenbrockf(x::Vector,grad::Vector)
             if length(grad) > 0
                     grad[1] = -2.0 * (1.0 - x[1]) - 400.0 * (x[2] - x[1]^2) * x[1]
                     grad[2] = 200.0 * (x[2] - x[1]^2)
             return (1.0 - x[1])^2 + 100.0 * (x[2] - x[1]^2)^2
         end
         function r_constraint(x::Vector, grad::Vector)
             if length(grad) > 0
                 grad[1] = 2*x[1]
                 grad[2] = 2*x[2]
                 return x[1]^2 + x[2]^2 - 0.8
         end
         opt = Opt(:LD_MMA, 2)
         lower_bounds!(opt, [-5, -5.0])
         min_objective!(opt,(x,g) -> rosenbrockf(x,g))
         inequality_constraint!(opt, (x,g) -> r_constraint(x,g))
         ftol_rel!(opt,1e-9)
         NLopt.optimize(opt, [-1.0,0.0])
```

# 2.8 JuMP.jl

- Introduce JuMP.jl
- JuMP is a mathematical programming interface for Julia. It is like AMPL, but for free and with a decent programming language.

Out [28]: (0.07588358473630112, [0.724702, 0.524221], :FTOL\_REACHED)

• The main highlights are:

- It uses automatic differentiation to compute derivatives from your expression.
- It supplies this information, as well as the sparsity structure of the Hessian to your preferred solver.
- It decouples your problem completely from the type of solver you are using. This is great, since you don't have to worry about different solvers having different interfaces.
- In order to achieve this, JuMP uses MathProgBase.jl, which converts your problem formulation into a standard representation of an optimization problem.
- Let's look at the readme
- The technical citation is Lubin et al [?]

### 2.9 JuMP: Quick start guide

- this is form the quick start guide
- please check the docs, they are excellent.

#### 2.9.1 Step 1: create a model

- A model collects variables, objective function and constraints.
- it defines a solver to be used.

```
using Clp
m = Model(solver=ClpSolver()) # provide a solver
#ăDefine variables
@variable(m, x )
                         # No bounds
@variable(m, x <= ub )  # Upper bound only</pre>
@variable(m, lb <= x <= ub ) # Lower and upper bounds</pre>
# we can create arrays of a variable
N = 2
Ovariable(m, x[1:M,1:N] >= 0)
# or put them in a block
Ovariables m begin
   X
   y >= 0
   Z[1:10], Bin
   X[1:3,1:3], SDP
   q[i=1:2], (lowerbound = i, start = 2i, upperbound = 3i)
   t[j=1:3], (Int, start = j)
end
# Equivalent to:
@variable(m, x)
@variable(m, y >= 0)
@variable(m, Z[1:10], Bin)
```

```
@variable(m, X[1:3,1:3], SDP)
@variable(m, q[i=1:2], lowerbound = i, start = 2i, upperbound = 3i)
@variable(m, t[j=1:3], Int, start = j)
# bounds can depend on indices
@variable(m, x[i=1:10] >= i )
```

### 2.10 Objective and Constraints

y = 0.2

• We can easily add objective and constraint functions:

```
@constraint(m, x[i] - s[i] <= 0) # Other options: == and >=
@constraint(m, sum(x[i] for i=1:numLocation) == 1)
@objective(m, Max, 5x + 22y + (x+y)/2) # or Min
```

• This is fully integrated with Julia. you can use the generator syntax for sums:

```
Oobjective(sum(x[i] + y[i]/pi for i = I1, j = I2 if i+j < some_val))
In [29]: ##ăSimple example
        using JuMP
         using Clp
         let
             m = Model(solver = ClpSolver())
             @variable(m, 0 \le x \le 2)
             Ovariable(m, 0 \le y \le 30)
             @objective(m, Max, 5x + 3*y)
             @constraint(m, 1x + 5y \le 3.0)
             print(m)
             status = solve(m)
             println("Objective value: ", getobjectivevalue(m))
             println("x = ", getvalue(x))
             println("y = ", getvalue(y))
         end
Max 5 x + 3 y
Subject to
x + 5 y 3
0 x 2
0 v 30
Objective value: 10.6
x = 2.0
```

```
In [30]: # JuMP: Rosenbrock Example
        # Instead of hand-coding first and second derivatives, you only have to give `JuMP` e
        # Here is an example.
        using Ipopt
        let
            m = Model(solver=IpoptSolver())
            @variable(m, x)
            @variable(m, y)
            @NLobjective(m, Min, (1-x)^2 + 100(y-x^2)^2)
            solve(m)
            println("x = ", getvalue(x), " y = ", getvalue(y))
        end
*************************************
This program contains Ipopt, a library for large-scale nonlinear optimization.
Ipopt is released as open source code under the Eclipse Public License (EPL).
        For more information visit http://projects.coin-or.org/Ipopt
************************************
This is Ipopt version 3.12.8, running with linear solver mumps.
NOTE: Other linear solvers might be more efficient (see Ipopt documentation).
Number of nonzeros in equality constraint Jacobian...:
                                                           0
Number of nonzeros in inequality constraint Jacobian.:
                                                           0
Number of nonzeros in Lagrangian Hessian...:
Total number of variables...:
                   variables with only lower bounds:
                                                           0
               variables with lower and upper bounds:
                                                           0
                    variables with only upper bounds:
                                                           0
Total number of equality constraints...:
Total number of inequality constraints...:
       inequality constraints with only lower bounds:
                                                           0
  inequality constraints with lower and upper bounds:
                                                           0
       inequality constraints with only upper bounds:
                                                           0
iter
       objective
                   inf_pr
                            inf_du lg(mu) ||d|| lg(rg) alpha_du alpha_pr ls
  0 1.0000000e+00 0.00e+00 2.00e+00 -1.0 0.00e+00 - 0.00e+00 0.00e+00
                                                     - 1.00e+00 2.50e-01f 3
  1 9.5312500e-01 0.00e+00 1.25e+01 -1.0 1.00e+00
```

```
2 4.8320569e-01 0.00e+00 1.01e+00 -1.0 9.03e-02
                                                     - 1.00e+00 1.00e+00f
  3 4.5708829e-01 0.00e+00 9.53e+00 -1.0 4.29e-01
                                                     - 1.00e+00 5.00e-01f
  4 1.8894205e-01 0.00e+00 4.15e-01 -1.0 9.51e-02
                                                     - 1.00e+00 1.00e+00f
  5 1.3918726e-01 0.00e+00 6.51e+00 -1.7 3.49e-01
                                                     - 1.00e+00 5.00e-01f
   6 5.4940990e-02 0.00e+00 4.51e-01 -1.7 9.29e-02
                                                     - 1.00e+00 1.00e+00f
    2.9144630e-02 0.00e+00 2.27e+00 -1.7 2.49e-01
                                                     - 1.00e+00 5.00e-01f
  8 9.8586451e-03 0.00e+00 1.15e+00 -1.7 1.10e-01
                                                     - 1.00e+00 1.00e+00f
                                                     - 1.00e+00 1.00e+00f
     2.3237475e-03 0.00e+00 1.00e+00 -1.7 1.00e-01
                            inf du lg(mu) ||d|| lg(rg) alpha du alpha pr ls
iter
       objective
                    inf pr
  10 2.3797236e-04 0.00e+00 2.19e-01 -1.7 5.09e-02
                                                     - 1.00e+00 1.00e+00f
  11 4.9267371e-06 0.00e+00 5.95e-02 -1.7 2.53e-02
                                                     - 1.00e+00 1.00e+00f
  12 2.8189505e-09 0.00e+00 8.31e-04 -2.5 3.20e-03
                                                     - 1.00e+00 1.00e+00f
  13 1.0095040e-15 0.00e+00 8.68e-07 -5.7 9.78e-05
                                                     - 1.00e+00 1.00e+00f
  14 1.3288608e-28 0.00e+00 2.02e-13 -8.6 4.65e-08
                                                     - 1.00e+00 1.00e+00f
Number of Iterations...: 14
                                                          (unscaled)
                                 (scaled)
               1.3288608467480825e-28
                                        1.3288608467480825e-28
Objective...:
                                                 2.0183854587685121e-13
Dual infeasibility...:
                       2.0183854587685121e-13
Constraint violation...:
                         0.000000000000000e+00
                                                  0.000000000000000e+00
Complementarity...:
                     0.000000000000000e+00 0.0000000000000e+00
Overall NLP error...:
                      2.0183854587685121e-13
                                               2.0183854587685121e-13
Number of objective function evaluations
                                                   = 36
Number of objective gradient evaluations
                                                   = 15
Number of equality constraint evaluations
                                                   = 0
Number of inequality constraint evaluations
                                                   = 0
Number of equality constraint Jacobian evaluations
Number of inequality constraint Jacobian evaluations = 0
Number of Lagrangian Hessian evaluations
Total CPU secs in IPOPT (w/o function evaluations)
                                                         0.158
Total CPU secs in NLP function evaluations
                                                         0.024
EXIT: Optimal Solution Found.
In [31]: # not bad, right?
        # adding the constraint from before:
        let
            m = Model(solver=IpoptSolver())
            @variable(m, x)
            @variable(m, y)
```

```
@NLobjective(m, Min, (1-x)^2 + 100(y-x^2)^2)
            ONLconstraint(m, x^2 + y^2 \le 0.8)
            solve(m)
            println("x = ", getvalue(x), " y = ", getvalue(y))
        end
This is Ipopt version 3.12.8, running with linear solver mumps.
NOTE: Other linear solvers might be more efficient (see Ipopt documentation).
Number of nonzeros in equality constraint Jacobian...:
                                                             0
Number of nonzeros in inequality constraint Jacobian.:
                                                             2
Number of nonzeros in Lagrangian Hessian...:
Total number of variables...:
                    variables with only lower bounds:
                                                             0
               variables with lower and upper bounds:
                                                             0
                    variables with only upper bounds:
                                                             0
Total number of equality constraints...:
Total number of inequality constraints...:
        inequality constraints with only lower bounds:
                                                             0
   inequality constraints with lower and upper bounds:
                                                             0
        inequality constraints with only upper bounds:
                                                             1
iter
        objective
                    inf_pr
                             inf_du lg(mu) ||d|| lg(rg) alpha_du alpha_pr
  0 1.0000000e+00 0.00e+00 2.00e+00 -1.0 0.00e+00
                                                       - 0.00e+00 0.00e+00
   1 9.5312500e-01 0.00e+00 1.25e+01 -1.0 5.00e-01
                                                       - 1.00e+00 5.00e-01f
   2 4.9204994e-01 0.00e+00 9.72e-01 -1.0 8.71e-02
                                                       - 1.00e+00 1.00e+00f
   3 2.0451702e+00 0.00e+00 3.69e+01 -1.7 3.80e-01
                                                       - 1.00e+00 1.00e+00H
   4 1.0409466e-01 0.00e+00 3.10e-01 -1.7 1.46e-01
                                                       - 1.00e+00 1.00e+00f
   5 8.5804626e-02 0.00e+00 2.71e-01 -1.7 9.98e-02
                                                       - 1.00e+00 1.00e+00h
  6 9.4244879e-02 0.00e+00 6.24e-02 -1.7 3.74e-02
                                                       - 1.00e+00 1.00e+00h
  7 8.0582034e-02 0.00e+00 1.51e-01 -2.5 6.41e-02
                                                       - 1.00e+00 1.00e+00h
  8 7.8681242e-02 0.00e+00 2.12e-03 -2.5 1.12e-02
                                                       - 1.00e+00 1.00e+00h 1
  9 7.6095770e-02 0.00e+00 6.16e-03 -3.8 1.37e-02
                                                       - 1.00e+00 1.00e+00h 1
                    inf_pr
                             inf_du lg(mu) ||d|| lg(rg) alpha_du alpha_pr ls
iter
       objective
  10 7.6033892e-02 0.00e+00 2.23e-06 -3.8 3.99e-04
                                                       - 1.00e+00 1.00e+00h 1
  11 7.5885642e-02 0.00e+00 2.07e-05 -5.7 7.99e-04
                                                       - 1.00e+00 1.00e+00h 1
  12 7.5885428e-02 0.00e+00 2.74e-11 -5.7 1.38e-06
                                                       - 1.00e+00 1.00e+00h 1
  13 7.5883585e-02 0.00e+00 3.19e-09 -8.6 9.93e-06
                                                       - 1.00e+00 1.00e+00f
Number of Iterations...: 13
                                                           (unscaled)
                                  (scaled)
Objective...:
               7.5883585442440671e-02 7.5883585442440671e-02
```

Dual infeasibility...: 3.1949178858070582e-09 3.1949178858070582e-09
Constraint violation...: 0.00000000000000e+00 0.000000000000000e+00
Complementarity...: 2.5454985882932001e-09 2.5454985882932001e-09
Overall NLP error...: 3.1949178858070582e-09 3.1949178858070582e-09

Number of objective function evaluations = 20Number of objective gradient evaluations = 14Number of equality constraint evaluations = 0Number of inequality constraint evaluations = 20Number of equality constraint Jacobian evaluations = 0Number of inequality constraint Jacobian evaluations = 14 Number of Lagrangian Hessian evaluations Total CPU secs in IPOPT (w/o function evaluations) 0.005 Total CPU secs in NLP function evaluations 0.003

EXIT: Optimal Solution Found. x = 0.7247018392092258 y = 0.5242206029480763

### 2.11 JuMP: Maximium Likelihood

- Let's redo the maximum likelihood example in JuMP.
- Let  $\mu$ ,  $\sigma^2$  be the unknown mean and variance of a random sample generated from the normal distribution.
- Find the maximum likelihood estimator for those parameters!
- density:

$$f(x_i|\mu,\sigma^2) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(x_i-\mu)^2}{2\sigma^2}\right)$$

• Likelihood Function

$$L(\mu, \sigma^2) = \prod_{i=1}^{N} f(x_i | \mu, \sigma^2) = \frac{1}{(\sigma \sqrt{2\pi})^n} \exp\left(-\frac{1}{2\sigma^2} \sum_{i=1}^{N} (x_i - \mu)^2\right)$$
$$= (\sigma^2 2\pi)^{-\frac{n}{2}} \exp\left(-\frac{1}{2\sigma^2} \sum_{i=1}^{N} (x_i - \mu)^2\right)$$

- Constraints:  $\mu \in \mathbb{R}, \sigma > 0$
- log-likelihood:

$$\log L = l = -\frac{n}{2} \log (2\pi\sigma^2) - \frac{1}{2\sigma^2} \sum_{i=1}^{N} (x_i - \mu)^2$$

• Let's do this in Jump.

```
In [32]: # Copyright 2015, Iain Dunning, Joey Huchette, Miles Lubin, and contributors
        # example modified
        using Distributions
        let
            distrib = Normal(4.5, 3.5)
            n = 10000
            data = rand(distrib,n);
            m = Model(solver=IpoptSolver())
            @variable(m, mu, start = 0.0)
            @variable(m, sigma >= 0.0, start = 1.0)
            solve(m)
            println(" = ", getvalue(mu),", mean(data) = ", mean(data))
            println("^2 = ", getvalue(sigma)^2, ", var(data) = ", var(data))
        end
This is Ipopt version 3.12.8, running with linear solver mumps.
NOTE: Other linear solvers might be more efficient (see Ipopt documentation).
Number of nonzeros in equality constraint Jacobian...:
Number of nonzeros in inequality constraint Jacobian.:
                                                          0
Number of nonzeros in Lagrangian Hessian...:
Total number of variables...:
                   variables with only lower bounds:
                                                          1
               variables with lower and upper bounds:
                                                          0
                   variables with only upper bounds:
                                                          0
Total number of equality constraints...:
Total number of inequality constraints...:
       inequality constraints with only lower bounds:
                                                          0
  inequality constraints with lower and upper bounds:
                                                          0
       inequality constraints with only upper bounds:
                                                          0
                   inf_pr inf_du lg(mu) ||d|| lg(rg) alpha_du alpha_pr ls
iter
       objective
  0 1.7210445e+05 0.00e+00 1.01e+02 -1.0 0.00e+00
                                                    - 0.00e+00 0.00e+00
  1 1.2939451e+05 0.00e+00 1.07e+02 -1.0 9.80e+00
                                                     - 1.00e+00 5.00e-01f
  2 7.8864616e+04 0.00e+00 4.43e+01 -1.0 2.27e-01
                                                    - 8.88e-01 1.00e+00f
  3 5.1720095e+04 0.00e+00 1.84e+01 -1.0 2.96e-01
                                                    - 1.00e+00 1.00e+00f
  4 3.7648943e+04 0.00e+00 7.44e+00 -1.0 3.75e-01
                                                    - 1.00e+00 1.00e+00f
  5 3.0796009e+04 0.00e+00 2.93e+00 -1.0 4.57e-01
                                                    - 1.00e+00 1.00e+00f
  6 2.7858621e+04 0.00e+00 1.07e+00 -1.0 5.13e-01
                                                    - 1.00e+00 1.00e+00f
                                                                          1
  7 2.6886136e+04 0.00e+00 3.39e-01 -1.0 4.94e-01
                                                    - 1.00e+00 1.00e+00f 1
```

```
8 2.6711082e+04 0.00e+00 6.36e-02 -1.7 3.19e-01 - 1.00e+00 1.00e+00f 1 9 2.6701110e+04 0.00e+00 4.25e-03 -2.5 1.01e-01 - 1.00e+00 1.00e+00f 1 iter objective inf_pr inf_du lg(mu) ||d|| lg(rg) alpha_du alpha_pr ls 10 2.6701072e+04 0.00e+00 1.72e-05 -3.8 6.78e-03 - 1.00e+00 1.00e+00f 1 11 2.6701072e+04 0.00e+00 8.12e-10 -5.7 4.62e-05 - 1.00e+00 1.00e+00f 1 12 2.6701072e+04 0.00e+00 4.04e-13 -8.6 1.02e-06 - 1.00e+00 1.00e+00f 1
```

Number of Iterations...: 12

(scaled) (unscaled) Objective...: 8.4542511167531504e+00 2.6701072346234618e+04 Dual infeasibility...: 4.0444162230182163e-13 1.2773485040573046e-09 0.000000000000000e+00 0.000000000000000e+00 Constraint violation...: Complementarity...: 2.5064398796454131e-09 7.9160923461665668e-06 Overall NLP error...: 2.5064398796454131e-09 7.9160923461665668e-06

```
Number of objective function evaluations = 18

Number of objective gradient evaluations = 13

Number of equality constraint evaluations = 0

Number of inequality constraint evaluations = 0

Number of equality constraint Jacobian evaluations = 0

Number of inequality constraint Jacobian evaluations = 0

Number of Lagrangian Hessian evaluations = 12

Total CPU secs in IPOPT (w/o function evaluations) = 0.007

Total CPU secs in NLP function evaluations = 0.026
```

EXIT: Optimal Solution Found.

- = 4.513536352090578, mean(data) = 4.513536352090578
- ^2 = 12.211002671496686, var(data) = 12.212223884224223

### 3 Linear Constrained Problems (LPs)

• Very similar to before, just that both objective and constraints are *linear*.

$$\begin{aligned} \min_{\mathbf{x}} \mathbf{c}^T \mathbf{x} \\ \text{subject to } \mathbf{w}_{LE}^{(i)T} \mathbf{x} &\leq b_i \text{ for } i \in 1, 2, 3, \dots \\ \mathbf{w}_{GE}^{(j)T} \mathbf{x} &\geq b_j \text{ for } j \in 1, 2, 3, \dots \\ \mathbf{w}_{EQ}^{(k)T} \mathbf{x} &= b_k \text{ for } k \in 1, 2, 3, \dots \end{aligned}$$

#### 3.0.1 Standard Form

- Usually LPs are given in standard form
- All constraints are less-than inequalities

• All choice variables are non-negative.

$$\min_{\mathbf{x}} \mathbf{c}^{T} \mathbf{x}$$
subject to  $\mathbf{A}\mathbf{x} \leq b$ 

$$\mathbf{x} > 0$$

- Greater-than inequality constraints are inverted
- equality constraints are split into two
- $\mathbf{x} = \mathbf{x}^+ \mathbf{x}^-$  and we constrain both components to be positive.

#### 3.0.2 Equality Form

$$\min_{\mathbf{x}} \mathbf{c}^{T} \mathbf{x}$$
subject to  $\mathbf{A}\mathbf{x} = b$ 

$$\mathbf{x} > 0$$

• Can transform standard into equality form

$$\mathbf{A}\mathbf{x} \leq b \rightarrow \mathbf{A}\mathbf{x} + \mathbf{s} = b, \mathbf{s} \geq 0$$

- equality constraints are split into two
- $x = x^+ x^-$  and we constrain both components to be positive.

### 3.0.3 Solving LPs

- Simplex Algorithm operates on Equality Form
- Moving from one vertex to the next of the feasible set, this is guaranteed to find the optimal solution if the problem is bounded.

# 4 Discrete Optimization / Integer Programming

- Here the choice variable is contrained to come from a discrete set  $\mathcal{X}$ .
- If this set is  $\mathcal{X} = \mathbb{N}$ , we have an **integer program**
- If only *some x* have to be discrete, this is a **mixed integer program**

### 4.1 Example

$$\min_{\mathbf{x}} x_1 + x_2$$
subject to  $||\mathbf{x}|| \le 2$ 

$$\mathbf{x} \in \mathbb{N}$$

- continuous optimum is  $(-\sqrt{2}, -\sqrt{2})$  and objective is  $y = -2\sqrt{2} = -2.828$
- Integer constrained problem is only delivering y = -2, and  $\mathbf{x}^* \in (-2,0), (-1,-1), (0,-2)$

### 4.2 Rounding

- One solution is to just round the continuous solution to the nearest integer
- We compute the **relaxed** problem, i.e. the one where *x* is continuous.
- Then we round up or down.
- Can go terribly wrong.

### 4.3 Cutting Planes

- This is an exact method
- We solve the relaxed problem first.
- Then we add linear constraints that result in the solution becoming integral.

#### 4.4 Branch and Bound

- This enumerates all possible soultions.
- Branch and bound does this, without having to compute all of them.

#### 4.5 Example: The Knapsack Problem

- We are packing our knapsack for a trip but only have space for the most valuable items.
- We have  $x_i = 0$  if item i is not in the sack, 1 else.

$$\min_{x} - \sum_{i=1}^{n} v_{i} x_{i}$$
s.t. 
$$\sum_{i=1}^{n} w_{i} x_{i} \leq w_{max}$$

$$w_{i} \in \mathbb{N}_{+}, v_{i} \in \mathbb{R}$$

- If ther are n items, we have  $2^n$  possible design vectors.
- But there is a useful recursive relationship.
- If we solved n-1 knapsack problems so far and deliberate about item n
  - If it's not worth including item n, then the solution **is** the knapsack problem for n-1 items and capacity  $w_{\text{max}}$
  - If it IS worth including it: solution will have value of knapsack with n-1 items and reduced capacity, plus the value of the new item

```
In [34]: # Copyright 2017, Iain Dunning, Joey Huchette, Miles Lubin, and contributors
       # This Source Code Form is subject to the terms of the Mozilla Public
       # License, v. 2.0. If a copy of the MPL was not distributed with this
       # file, You can obtain one at http://mozilla.org/MPL/2.0/.
       # JuMP
       # An algebraic modeling language for Julia
       # See http://github.com/JuliaOpt/JuMP.jl
       # knapsack.jl
       # Solves a simple knapsack problem:
       # max sum(p_j x_j)
       # st sum(w_j x_j) <= C
            x binary
       using JuMP, Cbc
       let
          # Maximization problem
          m = Model(solver=CbcSolver())
          Ovariable(m, x[1:5], Bin)
          profit = [ 5, 3, 2, 7, 4 ]
          weight = [2, 8, 4, 2, 5]
          capacity = 10
          # Objective: maximize profit
          @objective(m, Max, dot(profit, x))
          # Constraint: can carry all
          @constraint(m, dot(weight, x) <= capacity)</pre>
          # Solve problem using MIP solver
          status = solve(m)
          println("Objective is: ", getobjectivevalue(m))
          println("Solution is:")
          for i = 1:5
             print("x[$i] = ", getvalue(x[i]))
             println(", p[$i]/w[$i] = ", profit[i]/weight[i])
          end
       end
```

Objective is: 16.0

```
Solution is:
```

```
x[1] = 1.0, p[1]/w[1] = 2.5

x[2] = 0.0, p[2]/w[2] = 0.375

x[3] = 0.0, p[3]/w[3] = 0.5

x[4] = 1.0, p[4]/w[4] = 3.5

x[5] = 1.0, p[5]/w[5] = 0.8
```

### In []: