Optimization

To optimize (or minimize) $f(\mathbf{x}) : \mathbf{R}^n \to \mathbf{R}$ is to find $\mathbf{x}_0 \in \mathbf{R}^n$ so that $f(\mathbf{x}_0) \le f(\mathbf{x})$ for all $\mathbf{x} \in \mathbf{R}^n$.

Algorithms

- Use calculus to find an exact solution if you can.
- Golden section search does not require derivatives. See goldenSectionSearch.R to graph and minimize $f(x) = \frac{1}{10}x^2 2\sin(x)$ over (0,4). (Also try (-15,15).)
- Gradient descent requires first partial deriviatives.

Recall from calculus that for a function $y = f(x_1, ..., x_n)$, the the gradient of f is defined as $\nabla f(x_1, ..., x_n) = \left(\frac{\partial y}{\partial x_1}, ..., \frac{\partial y}{\partial x_n}\right)$. To minimize f by gradient descent, choose an initial point $(x_1, ..., x_n)$ and iteratively move opposite the gradient by iterating on

$$\mathbf{x}_{i+1} = \mathbf{x}_i - \gamma \nabla f(\mathbf{x}_i)$$

where γ is the *step size* parameter. Note that γ can be adjusted as the algorithm proceeds; a *line search* can be used to guarantee convergence for a well-behaved f. e.g. See gradientDescent.R.

- Newton's method requires first and second partial derivatives. It is an iterative method for approximating the roots of a function, finding x such that f(x) = 0. In optimization, Newton's method is applied to the derivative function f'(x) to find x such that f'(x) = 0, since such an x is a minimum, maximum, or inflection point. e.g. For the n = 1 case, see Newton.R. (The n > 1 case requires a Hessian matrix that isn't introduced in the prerequisites to STAT 327.)
- Nelder-Mead is a heuristic method that does not require derivatives. It evaluates f over a simplex, a set of n+1 vertices in n dimensions that is a generalization of a triangle (2+1) vertices in 2 dimensions, repeatedly replacing the worst vertex with one computed from the others. See NelderMead.R.

R functions

• n = 1

optimize(f, interval, ...) minimizes the continuous function f, whose opposite -f is unimodal, over its first argument over the interval (interval[1], interval[2]), where ... are additional arguments passed to f. It returns a list containing:

- minimum, the argument that minimizes f
- objective, the value f (minimum)

?optimize says it uses golden section search (with successive parabolic interpolation).

Note: If f is not unimodal, optimize may get stuck at a local minimum.

optim(par, fn, gr=NULL, ..., method=c("Nelder-Mead", "BFGS", "CG", "L-BFGS-B", "SANN", "Brent") minimizes the function fn over its first vector argument, a vector of parameters, starting at initial values in the vector par, where ... are additional arguments passed to fn. gr, the gradient of fn, is required for some values of method. optim() returns a list containing:

- par, the parameters that minimize fn
- value, which is f(par)
- convergence, a code with 0 indicating success, 1 indicating an iteration limit was reached, and other values indicating other trouble

Regarding method,

- "Nelder-Mead", the default, does not require gr.
- "BFGS" approximates Netwon's method; it requires gr
- (- "CG" uses a *conjugate gradient* method that may be more fragile than "BFGS" but useful for large problems; it requires gr
- "L-BFGS-B" is a limited-memory variant of "BFGS" that is suitable for large n
- "SANN" uses *simulated annealing*, a probabilistic heuristic for finding an approximately optimal solution; it does not require derivatives
- "Brent", useful only for n=1, just calls optimize; it does not use $\operatorname{\mathsf{gr}}$

Statistics uses optimization to estimate parameters

Optimization finds those parameter values that make the observed data most likely.

e.g. To find the optimal simple linear regression model of the form $\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x$ for the data $\{(x_i, y_i) : i = 1, \dots, n\}$, we find $\hat{\beta}_0$ and $\hat{\beta}_1$ that minimize the sum of squared errors,

$$SSE = SSE(\hat{\beta}_0, \hat{\beta}_1; \{(x_i, y_i)\}) = \sum_{i=1}^{n} e_i^2 = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2 = \sum_{i=1}^{n} (y_i - (\hat{\beta}_0 + \hat{\beta}_1 x_i))^2$$

We could use optimization to find the least-squares model, but we already have a closed form solution from calculus in lm(). In the homework, you'll use optimization to solve an important variant that, instead of using least squares, uses least absolute deviations. e.g.

```
SSE = function(beta, x, y) { # usual least squares
   return(sum((y - (beta[1] + beta[2] * x))^2))
}
out = optim(par=c(0, 0), fn=SSE, x=mtcars$wt, y=mtcars$mpg)
m = lm(mpg ~ wt, data=mtcars)
out2 = optim(par=c(0, 0), fn=SSE, x=mtcars$wt, y=mtcars$mpg, control=list(reltol=1e-12))
# Try constant model too.
```

Optimization is a big field. See http://cran.r-project.org/web/views/Optimization.html for much more.