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Announcements

- PS3 is now late
- PS4 is posted, due in two weeks
 - This is all numerical optimization
 - The PCA activity, while cool, got punted to PS5
 - Happened because the optimization activities are plenty long on their own

Check-In

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Motivating Problem

Numerical Optimization

- lacktriangle We have a function, $f(\mathbf{X}, heta)$, where \mathbf{X} is data and heta are parameters
- Numerical optimization answers the question:

$$\operatorname*{argmin}_{ heta}fig(\mathbf{X}, hetaig)$$

- From last week, remember argmin (or argmax) says: what value of θ minimizes (or maximizes) the output of $f(\mathbf{X}, \theta)$?
- Note that now ${f X}$ is a matrix argument (that also contains our outcome, y) and heta is (often) a vector argument
- If we are using argmin, $f(\mathbf{X}, \theta)$ is often referred to as a *loss function*
 - Our goal is to write problems such that when we minimize the loss function, we've found the answer!

Just thinking about argmin

How could we use a computer to minimize this?

$$\operatorname*{argmin}_{x}x^{2}-2x-3$$

- Take the derivative with respect to x, set $\frac{df}{dx} = 0$, and solve
- lacktriangle Make a list of possible x values, plug them in, and check which one gives the lowest value
- Plot it and eyeball it
- Some secret fourth thing???

How do we optimize?

- Remember that analytic solutions can be hard/impossible and grid search can be inefficient/slow/imprecise
- Numerical optimization leverages an algorithm to find a solution faster, more efficiently, and more precisely than grid search
- There are LOTS of algorithms with different tradeoffs
- We will develop my favorite one today
 - It might actually be my all-time favorite algorithm?

Tools

Single Precision Floating Point Numbers

- In computers, numbers are typically stored using 32 bits of memory
 - Each bit can have a value of either 0 or 1
 - This means that there are only 4,294,967,296 possible values
 - This has to cover every possible positive number, negative number, decimal, or super huge value
- The way computers handle this is using floating point numbers
 - Think of this as scientific notation
 - ullet 2, 395, 423 ightarrow 2.395423 imes 10⁶
- How is this stored?
 - One bit stores the sign (+/-)
 - Eight bits store the exponent (256 values, ranging from -126 to 127 with all 0s or 1s held back)
 - 23 bits stores the mantissa (this is about 6-8 decimal places of precision)
 - Importantly, the more extreme the value of the exponent, the less precise the entire number
 - This means nearly all computations contain some amount of rounding error

Monotonic Functions and Transformations

- A function is monotonic if it preserves order
- For a monotonic function, *f* :
 - \bullet $a > b \implies f(a) > f(b)$
- We will use the *log transformation* all the time in numerical optimization
- Because log(x) is a monotonic function:

$$\operatorname*{argmin}_x f(x) = \operatorname*{argmin}_x \log f(x)$$

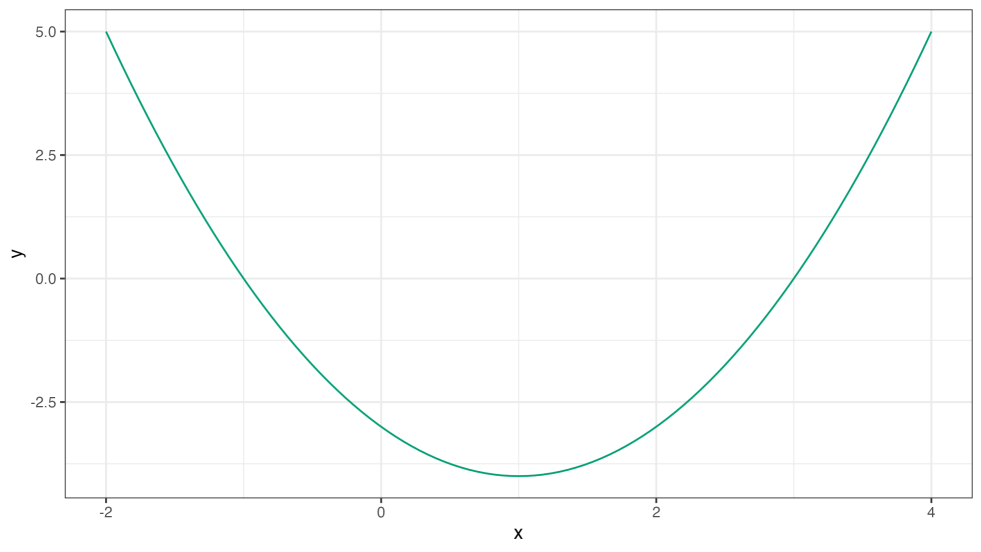
- You can turn multiplication into addition, leading to less extreme exponent values
 - $\bullet \ \log(ab) = \log(a) + \log(b)$
- The log transform maps positive numbers with negative exponents to negative numbers (with smaller exponents) and positive numbers with positive exponents into positive numbers (with smaller exponents)

Derivatives and Gradients

- Recall the derivative of a function with respect to a variable x is the slope of a line tangent to the function at a specific value of x
- The gradient is the multivariate (read: *vector*) generalization of the derivative
 - The gradient constructs a column vector of partial derivatives
- If you think of a multivariate function as a surface, the gradient is a vector that points "uphill"
- lacksquare We write the derivative of f with respect to x as: $rac{df}{dx}$
- If f takes a vector argument, \mathbf{x} , of dimension k, we write the gradient of f as: ∇f

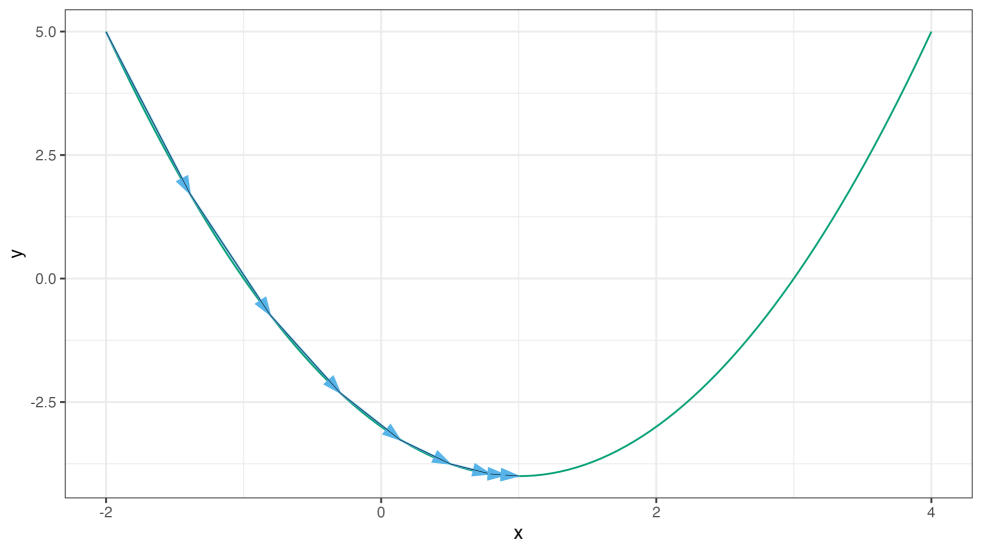
$$abla f = rac{\partial f}{\partial \mathbf{x}} = \left[rac{\partial f}{\partial x_1}rac{\partial f}{\partial x_2}\cdotsrac{\partial f}{\partial x_k}
ight]^ op$$

Algorithm: Gradient Descent



Gradient Descent

- Big idea: To find the minimum value of a function, we move downhill until we get to the bottom
- Algorithm: To find the argmin of a function f given parameter values θ_n at iteration n, and step size λ :
 - 1. Compute the gradient of f with respect to θ_n
 - 2. Find the next parameter values, θ_{n+1} , according to the update rule:
 - ullet $heta_{n+1} \leftarrow heta_n \lambda
 abla f$
 - 3. Repeat 1&2 until convergence
- What actually happens: Starting at some guess, use the gradient to repeatedly take steps "downhill" until you hit the bottom



- So how do we write a loss function?
- Frame your solution as a minimization problem
 - Note that if you want to maximize something, it's the same as minimizing the negative of that thing
- Recall that OLS attempts to minimize the sum of squared residuals
 - For OLS with a single covariate x, outcome y, and coefficients, β_0 , β_1 , write down the sum of squared residuals
 - Then how do we express that we want to minimize? What do we minimize with respect to?
 - What does this give us?

Set up the problem:

$$\sum_{x_i,y_i\in\mathbf{X}}ig(y_i-(eta_0+eta_1x_i)ig)^2$$

The sum of squared residuals $\,$

Set up the problem:

$$\sup_{eta} \sum_{x_i,y_i \in \mathbf{X}} ig(y_i - (eta_0 + eta_1 x_i)ig)^2$$
 Find eta that minimizes The sum of squared residuals

Set up the problem:

$$\hat{\beta} = \operatorname*{argmin}_{\beta \text{ that minimizes}} \sum_{x_i,y_i \in \mathbf{X}} \left(y_i - (\beta_0 + \beta_1 x_i)\right)^2$$

Gradient Descent with OLS

We start with our loss function:

$$\ell(\mathbf{X},eta) = \sum_{x_i,y_i \in \mathbf{X}} ig(y_i - (eta_0 + eta_1 x_i)ig)^2$$

• Now we need to take the gradient wrt the parameters we optimize over:

$$abla \ell(\mathbf{X},eta) = egin{bmatrix} rac{\partial \ell}{\partial eta_0} \ rac{\partial \ell}{\partial eta_1} \end{bmatrix}$$

The gradient:

$$abla \ell(\mathbf{X},eta) = egin{bmatrix} rac{\partial \ell}{\partial eta_0} \ rac{\partial \ell}{\partial eta_1} \end{bmatrix} = egin{bmatrix} \sum_{x_i,y_i \in \mathbf{X}} -2ig(y_i - (eta_0 + eta_1 x_i)ig) \ \sum_{x_i,y_i \in \mathbf{X}} -2x_iig(y_i - (eta_0 + eta_1 x_i)ig) \end{bmatrix}$$

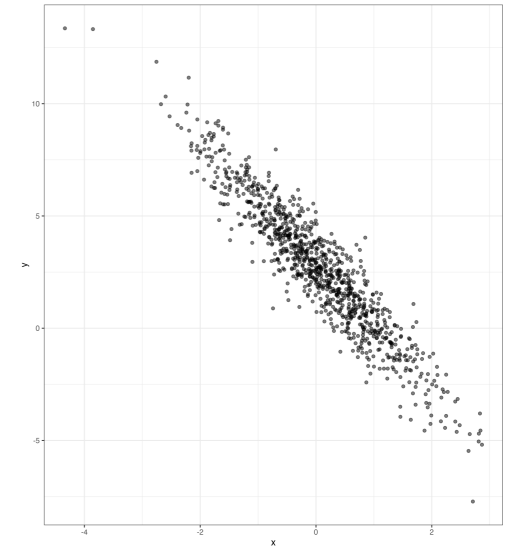
Gradient Descent with OLS

- Next we pick a step size, λ
 - Smaller values of λ result in a more precise solution, but slower convergence time
 - Larger values of λ are less precise but converge faster
 - If you make λ too large, the optimization may become unstable and never converge!
- Initialize starting values for your parameters
 - Lots of choices! Start at zero? Pick randomly?
- Update your parameters:
 - Recall $\theta_{n+1} \leftarrow \theta_n \lambda \nabla f$
 - ullet $\hat{eta}_{0,n+1} \leftarrow \hat{eta}_{0,n} + 2\lambda \sum_{x_i,y_i \in \mathbf{X}} \left(y_i (\hat{eta}_{0,n} + \hat{eta}_{1,n}x_i)
 ight)$
 - $oldsymbol{\hat{eta}}_{1,n+1} \leftarrow \hat{eta}_{1,n} + 2\lambda \sum_{x_i,y_i \in \mathbf{X}} x_i ig(y_i (\hat{eta}_{0,n} + \hat{eta}_{1,n} x_i)ig)$

Simulating some OLS data

Simulating some OLS data

```
1     ggplot(d, aes(x = x, y = y)) +
2     geom_point(alpha = 0.5) +
3     theme_bw()
```



Implementing the Gradient Descent Update

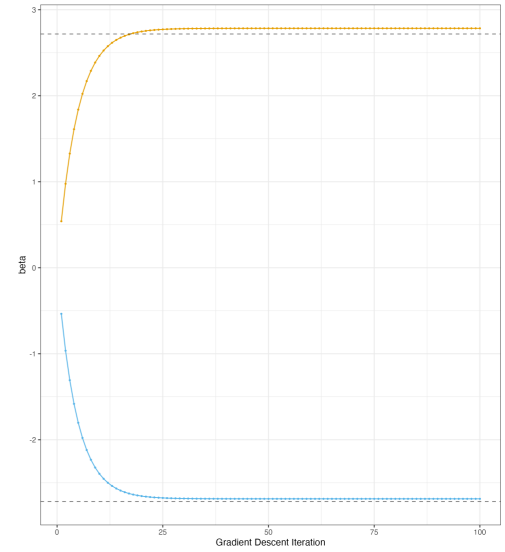
```
1 lr <- 1e-4
      2 beta <-c(0, 0)
     4 beta\lceil 1 \rceil <- beta\lceil 1 \rceil + 2 * lr * sum(d\( \frac{1}{2} \) - (beta\lceil 1 \rceil + beta\lceil 2 \rceil * d\( \frac{1}{2} \) \( \frac{1}{2} \) * d\( \frac{1}{2} \) \( \frac{1}{2} \) * d\( \frac{1}{2} \) \( \frac{1}{2} \) * d\( \fr
      5 beta[2] <- beta[2] + 2 * lr * sum(d$x * (d$y - (beta[1] + beta[2] * d$x)))
                        beta
     9 # [1] 0.5407162 -0.5354659
                        beta[1] \leftarrow beta[1] + 2 * lr * sum(d$y - (beta[1] + beta[2] * d$x))
                     beta[2] \leftarrow beta[2] + 2 * lr * sum(d$x * (d$y - (beta[1] + beta[2] * d$x)))
13
                      beta
16 # [1] 0.9764911 -0.9641414
```

How long until convergence? $\lambda=10^{-4}$

```
1 M <- 1e2
 2 lr <- 1e-4
     beta \leftarrow c(0, 0)
     betas <- data.frame(i = 1:M, beta 0 = numeric(M), beta 1 = numeric(M))
     for (i in 1:M) {
      beta[1] \leftarrow beta[1] + 2 * lr * sum(d$y - (beta[1] + beta[2] * d$x))
       beta[2] \leftarrow beta[2] + 2 * lr * sum(d$x * (d$y - (beta[1] + beta[2] * d$x)))
 9
10
       betas$beta 0[i] <- beta[1]
       betas$beta 1[i] <- beta[2]
11
12
13
14
     beta
15
16 # [1] 2.783908 -2.686621
```

$\lambda = 10^{-4}$

```
ggplot(betas, aes(x = i)) +
       geom line(aes(y = beta 0),
                 color = okabeito colors(1)) +
       geom_line(aes(y = beta_1),
                 color = okabeito_colors(2)) +
       geom_point(aes(y = beta_0),
                  size = 0.5,
                  color = okabeito colors(1)) +
       geom point(aes(y = beta 1),
                  size = 0.5,
10
                  color = okabeito colors(2)) +
11
12
       geom_hline(aes(yintercept = true_beta[1]),
13
                  lty = 2,
                  alpha = 0.5) +
14
15
       geom_hline(aes(yintercept = true_beta[2]),
                      lty = 2,
16
17
                      alpha = 0.5) +
       labs(x='Gradient Descent Iteration',
18
            y = 'beta') +
19
20
       theme_bw()
```

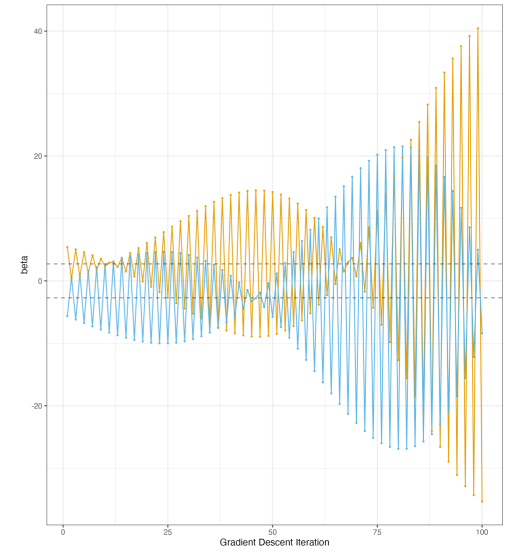


What about a faster learning rate? $\lambda=10^{-3}$

```
1 M <- 1e2
 2 lr <- 1e-3
     beta \leftarrow c(0, 0)
     betas <- data.frame(i = 1:M, beta 0 = numeric(M), beta 1 = numeric(M))
     for (i in 1:M) {
      beta[1] \leftarrow beta[1] + 2 * lr * sum(d$y - (beta[1] + beta[2] * d$x))
       beta[2] \leftarrow beta[2] + 2 * lr * sum(d$x * (d$y - (beta[1] + beta[2] * d$x)))
 9
10
       betas$beta 0[i] <- beta[1]
       betas$beta 1[i] <- beta[2]
11
12
13
14
     beta
15
16 # [1] -35.335523 -8.404527
```

$\lambda = 10^{-3}$

```
ggplot(betas, aes(x = i)) +
       geom line(aes(y = beta_0),
                 color = okabeito colors(1)) +
       geom_line(aes(y = beta_1),
                 color = okabeito_colors(2)) +
       geom_point(aes(y = beta_0),
                  size = 0.5,
 8
                  color = okabeito colors(1)) +
       geom point(aes(y = beta_1),
 9
                  size = 0.5,
10
                  color = okabeito colors(2)) +
11
12
       geom_hline(aes(yintercept = true_beta[1]),
13
                  lty = 2,
                  alpha = 0.5) +
14
15
       geom_hline(aes(yintercept = true_beta[2]),
                      lty = 2,
16
17
                      alpha = 0.5) +
       labs(x='Gradient Descent Iteration',
18
            y = 'beta') +
19
20
       theme_bw()
```

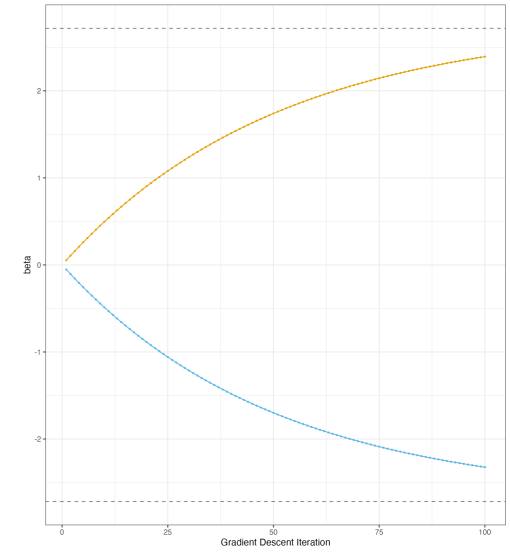


What about a slower learning rate? $\lambda=10^{-5}$

```
1 M <- 1e2
 2 lr <- 1e-5
     beta \leftarrow c(0, 0)
     betas <- data.frame(i = 1:M, beta 0 = numeric(M), beta 1 = numeric(M))
     for (i in 1:M) {
      beta[1] \leftarrow beta[1] + 2 * lr * sum(d$y - (beta[1] + beta[2] * d$x))
       beta[2] \leftarrow beta[2] + 2 * lr * sum(d$x * (d$y - (beta[1] + beta[2] * d$x)))
 9
10
       betas$beta 0[i] <- beta[1]
       betas$beta_1[i] <- beta[2]</pre>
11
12
13
14
     beta
15
16 # [1] 2.392762 -2.323370
```

$\lambda = 10^{-5}$

```
ggplot(betas, aes(x = i)) +
       geom line(aes(y = beta 0),
                 color = okabeito colors(1)) +
       geom_line(aes(y = beta_1),
                 color = okabeito_colors(2)) +
       geom_point(aes(y = beta_0),
                  size = 0.5,
                  color = okabeito colors(1)) +
 9
       geom point(aes(y = beta 1),
                  size = 0.5,
10
                  color = okabeito colors(2)) +
11
12
       geom_hline(aes(yintercept = true_beta[1]),
13
                  lty = 2,
                  alpha = 0.5) +
14
15
       geom_hline(aes(yintercept = true_beta[2]),
                      lty = 2,
16
17
                      alpha = 0.5) +
       labs(x='Gradient Descent Iteration',
18
            y = 'beta') +
19
20
       theme bw()
```



When do you stop?

- The easiest way to do it is just pick a number of iterations and then stop
 - You may not reach the solution
 - You may take too many iterations that you didn't need to
- Typically we stop when the solution converges
 - Often this looks like picking a threshold value, ϵ
 - After every iteration, check to see how much the estimate has changed by
 - Stop when the change in parameter estimates is smaller than the threshold
 - Lets you set the level of precision you want in your answer
 - If your threshold is too small relative to the step size, the optimization routine may never converge

More Tools

Maximum Likelihood Estimation

Maximum Likelihood Estimation

• A linear regression assumes normally distributed error with constant error variance, or:

$$y_i \sim \mathcal{N}ig(eta_0 + eta_1 x_i, \sigma^2ig)$$

• We can write down the *likelihood* (think Bayes' theorem) of observing a set of parameters conditioned on our observed data by multiplying together the normal density functions for each observation:

$$L(eta|\mathbf{X}) = \prod_{y_i,x_i \in \mathbf{X}} rac{1}{\sigma \sqrt{2\pi}} e^{-rac{1}{2}\left(rac{y_i - (eta_0 + eta_1 x_i)}{\sigma}
ight)^2}$$

Log Transformin'

- We can make our lives way easier with a log transform! Why?
- Multiplying lots of probabilities results in numerical instability by ending up with tiny numbers!
- The log transform can turn these products into a sum!

$$egin{aligned} \log \prod_{y_i, x_i \in \mathbf{X}} rac{1}{\sigma \sqrt{2\pi}} e^{-rac{1}{2} \left(rac{y_i - (eta_0 + eta_1 x_i)}{\sigma}
ight)^2} &= \sum_{y_i, x_i \in \mathbf{X}} \log rac{1}{\sigma \sqrt{2\pi}} e^{-rac{1}{2} \left(rac{y_i - (eta_0 + eta_1 x_i)}{\sigma}
ight)^2} \ &= \sum_{y_i, x_i \in \mathbf{X}} \log rac{1}{\sigma \sqrt{2\pi}} + \log e^{-rac{1}{2} \left(rac{y_i - (eta_0 + eta_1 x_i)}{\sigma}
ight)^2} \ &= \sum_{y_i, x_i \in \mathbf{X}} \log rac{1}{\sigma \sqrt{2\pi}} - rac{1}{2} \left(rac{y_i - (eta_0 + eta_1 x_i)}{\sigma}
ight)^2 \end{aligned}$$

Maximum (Log) Likelihood Estimation

- Maximum Likelihood Estimation (MLE) is just finding the values of your parameters that maximize the likelihood
- Because the log transform is monotonic, we can just maximize the log likelihood instead

$$\ell(heta|\mathbf{X}) = \log L(heta|\mathbf{X})$$
 $\hat{ heta} = rgmax_{ heta} L(heta|\mathbf{X}) = rgmax_{ heta} \ell(heta|\mathbf{X})$

- We have a few options on how this works:
- 1. Take the gradient of the log likelihood and set it equal to zero to find the estimates $\hat{ heta}$ analytically
- 2. Use numerical optimization
- 3. Some secret third thing that we haven't really discussed yet

MLE for OLS

$$\hat{eta} = rgmax_{eta} \left[\sum_{y_i, x_i \in \mathbf{X}} \log \frac{1}{\sigma \sqrt{2\pi}} - \frac{1}{2} \left(\frac{y_i - (eta_0 + eta_1 x_i)}{\sigma} \right)^2 \right]$$
 $\hat{eta} = rgmax_{eta} \left[N \log \frac{1}{\sigma \sqrt{2\pi}} - \frac{1}{2\sigma^2} \sum_{y_i, x_i \in \mathbf{X}} \left(y_i - (eta_0 + eta_1 x_i) \right)^2 \right]$
 $\hat{eta} = rgmax_{eta} \left[N \log \frac{1}{\sigma \sqrt{2\pi}} - \underbrace{\frac{1}{2\sigma^2}}_{Constant} \sum_{y_i, x_i \in \mathbf{X}} \left(y_i - (eta_0 + eta_1 x_i) \right)^2 \right]$
 $\hat{eta} = rgmax_{eta} \left[- \sum_{y_i, x_i \in \mathbf{X}} \left(y_i - (eta_0 + eta_1 x_i) \right)^2 \right]$

Does this last line look familiar?

MLE for OLS

$$\hat{eta} = \operatorname*{argmin}_{eta} \sum_{y_i, x_i \in \mathbf{X}} ig(y_i - (eta_0 + eta_1 x_i) ig)^2$$

- The MLE for OLS is identical to minimizing the sum of squared residuals
- This is not always the case for other estimators, though!
- Now we have to actually minimize it—for this, we'll use optim()

optim()

All hail the greatest built-in function in base R, optim()

- optim() is a general purpose optimization function in R
- It has a bunch of different optimization methods and is used by just about every single modeling function under the hood
- optim() specifically does argmin
 - You write a function that takes parameters as inputs
 - You give optim() starting values for those parameters (as a vector) and the function to be argmin'd
 - optim() returns the values of the parameters that minimize the function
- If you can write something as a maximization or minimization problem, optim() can solve it for you
 - Caveats about convexity and identifiability and some other stuff

MLE with optim() for OLS

```
SumSquaredResid <- function(beta, d){</pre>
     # TODO: Compute sum of squared residual
     resid \leftarrow d$y - (beta[1] + beta[2]*d$x)
     ssr <- sum(resid^2)</pre>
     return(ssr)
   SumSquaredResid(c(0,0), d)
   # [1] 15666.95
   out <- optim(c(0,0), # starting vals for parameters
                 SumSquaredResid, # fn to minimize
14
                 d=d)
                      # args to pass to fn
   # pass control=list(scale=-1) to make optim argmax
18
   mle beta <- out$par</pre>
   mle beta
23 # [1] 2.784030 -2.686716
```

Checking results

```
1 true_beta
3 # [1] 2.718 -2.718
5 ols_beta
7 # (Intercept) x
8 # 2.783908 -2.686621
10 gd_beta
12 # [1] 2.783908 -2.686621
13
14 mle_beta
16 # [1] 2.784030 -2.686716
```

Some notes on optim()

- The function you minimize needs to output only a single scalar value
- The first argument of the function you minimize needs to be all the parameters you want to optimize,
 passed as a vector
- There are a whole bunch of options you can fiddle with in order to change convergence conditions, speed, or optimization algorithm
- If you make a function that also returns the gradient, you can gain a lot of speed and precision!
- Works best with convex problems
 - Otherwise you can get stuck in local minima!
 - If you can't be convex, just start from a bunch of random spots and take the best solution
 - If your problem involves finding where a function is equal to zero, squaring it and finding the minimum will do the job!

Wrap Up

Recap

- Numerical optimization is the most useful skill we learn in this course
- If you can frame something as an optimization problem, optim() can generally solve it
- Working in log space has tons of advantages, especially when you're dealing with probabilities

Final Thoughts

PollEv.com/klintkanopka