# Maximum Likelihood Estimation

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```
# Load libraries
library(tidyverse)
library(broom)
```

# Joint Probability Density

Joint probability density, or simply joint density, is simply the density of  $x_1, x_2, ...$  AND  $x_k$ . If we can make an assumption about INDEPENDENCE, then the joint density would be the product of the individual densities,

$$p(x_1, x_2, x_3, \dots, x_K) = p(x_1) \times p(x_2) \times p(x_3) \times \dots \times p(x_k)$$

Say we had three independent observations from our  $\sim \mathcal{N}(50, 10)$  distribution, namely  $x = \{60, 65, 67\}$ . Then the joint density would be,

```
dnorm(x = 60, mean = 50, sd = 10) * dnorm(x = 65, mean = 50, sd = 10) * dnorm(x = 67, mean = 50, sd = 10)
```

## [1] 0.000002947448

We could also shortcut this computation,

```
prod(dnorm(x = c(60, 65, 67), mean = 50, sd = 10))
```

## [1] 0.000002947448

This value is the joint probability density.

In the previous example, the joint probability density indicates the probability of observing the data ( $x = \{60, 65, 67\}$ ) GIVEN (1) they are drawn from a normal distribution and (2) the normal distribution has a mean of 50 and a standard deviation of 10. In other words, it is the probability of the data given the distribution and parameters. Symbolically,

Joint Density =  $P(Data \mid Distribution and Parameters)$ 

#### Likelihood

Likelihood is the probability of a particular set of parameters GIVEN (1) the data, and (2) the data are from a normal distribution. Symbolically,

```
Likelihood = P(Parameters | Distribution and Data)
```

Likelihood takes the data as given and computes the probability of a set of parameters. Symbolically we denote likelihood with a caligraphic letter "L" ( $\mathcal{L}$ ). For example, we might ask the question, given the observed data  $x = \{30, 20, 24, 27\}$  come from a normal distribution, what is the likelihood (probability) that the mean is 20 and the standard deviation is 4? We might denote this as,

$$\mathcal{L}(\mu = 20, \sigma = 4 \mid x)$$

Note that although we need to specify the distribution (e.g., normal), this is typically not included in the symbolic notation; instead it is typically included in the assumptions.

To compute the likelihood we compute the joint density of the data under that particular set of parameters.

```
prod(dnorm(x = c(30, 20, 24, 27), mean = 20, sd = 4))
```

```
## [1] 0.0000005702554
```

What is the likelihood (probability) that the mean is 25 and the standard deviation is 4?

```
prod(dnorm(x = c(30, 20, 24, 27), mean = 25, sd = 4))
```

```
## 「17 0.00001774012
```

Which set of parameters is *more likely* given the data and that they come from a normal distribution? The second set of parameters is more likely since it has a higher likelihood.

It is important to note that although we use the joint probability under a set of parameters to compute the likelihood of those parameters, theoretically joint density and likelihood are very different. Likelihood refers to the probability of the parameters and joint density refers to the probability of the data.

#### Maximum Likelihood

So now we come to the crux of Maximum Likelihood Estimation (MLE). The goal of MLE is to find a set of parameters that MAXIMIZES the likelihood given the data and a distribution. For example, given the observed data  $x = \{30, 20, 24, 27\}$  come from a normal distribution, what are the values for the parameters (mean and standard deviation) that give the highest liklihood?

#### Solution 1: Grid Search

One metho for finding the parameters (mean and standard deviation) that produce the MAXIMUM likelihood, is to substitute several parameter values in the dnorm() function, compute the likelihood for each set of parameters, and determine which set produces the highest (maximum) likelihood.

In computer science, this method for finding the MLE is referred to as a *grid search*. Below is some syntax to carry out a grid search. The syntax creates several sets of parameter values (called the search space), computes the likelihood for each combination of parameter values, and then arranges the likelihoods in descending order.

```
expand.grid(
  mu = seq(from = 10, to =30, by = 0.1),
  sigma = seq(from = 0, to = 10, by = 0.1)
  ) %>%
  rowwise() %>%
  mutate( lik = prod(dnorm(c(30, 20, 24, 27), mean = mu, sd = sigma)) ) %>%
  arrange(desc(lik))
```

```
## # A tibble: 20,301 x 3
##
                       lik
        mu sigma
     <dbl> <dbl>
##
      25.2 3.70 0.0000183
##
##
   2
      25.3 3.70 0.0000183
##
      25.3 3.80 0.0000182
      25.2 3.80 0.0000182
      25.4 3.70 0.0000182
##
##
   6
      25.1
            3.70 0.0000182
##
   7
      25.2 3.60 0.0000182
   8 25.3 3.60 0.0000182
##
## 9
      25.1
            3.80 0.0000182
## 10 25.4 3.80 0.0000182
```

```
## # ... with 20,291 more rows
```

The parameters that maximize the likelihood (in our search space) are a mean of 25.2 and a standard deviation of 3.7.

#### Log-Likelihood

The likelihood values are quite small since we are multiplying several probabilities together. We could take the natural logarithm of the likelihood to alleviate this issue. So in our example,  $\mathcal{L}=.00001829129$  and the log-likelihood would be

```
log(.00001829129)
```

```
## [1] -10.90909
```

We typically denote log-likelihood using a caligraphic lower-case "1" (1).

Going back to how we compute the likelihood, we assumed a set of parameters and then found the joint density, which assuming normality and independence is the product of the individual densities.

$$\mathcal{L}(\text{parameters}|\text{data}) = p(x_1) \times p(x_2) \times \ldots \times p(x_n)$$

If we compute the log of the likelihood instead:

$$\updownarrow (parameters|data) = \ln \Big( \mathcal{L}(parameters|data) \Big) = \ln \Big( p(x_1) \times p(x_2) \times \ldots \times p(x_n) \Big)$$

Using the rules of logarithms, the right-hand side of the equation can be manipulated to

$$= \ln(p(x_1)) + \ln(p(x_2)) + \ldots + \ln(p(x_n))$$

The log-likelihood is the *sum of the log-transformed densities*. This means we could re-write our grid search syntax to compute the log-likelihood. Since finding the log of the densities is so useful, there is even an argument in dnorm() of log=TRUE that does this for us. Our revised grid search syntax is:

```
expand.grid(
  mu = seq(from = 10, to =30, by = 0.1),
  sigma = seq(from = 0, to = 10, by = 0.1)
  ) %>%
  rowwise() %>%
  mutate( log_lik = sum(dnorm(c(30, 20, 24, 27), mean = mu, sd = sigma, log = TRUE)) ) %>%
  arrange(desc(log_lik))
```

```
## # A tibble: 20,301 x 3
##
         mu sigma log_lik
##
      <dbl> <dbl>
                    <dbl>
       25.2
             3.70
##
                    -10.9
##
       25.3
             3.70
                    -10.9
##
       25.3
             3.80
                    -10.9
##
       25.2
             3.80
                    -10.9
##
       25.1
             3.70
                    -10.9
             3.70
                    -10.9
       25.4
##
   7
       25.3
             3.60
                    -10.9
##
       25.2
             3.60
                    -10.9
##
  9
       25.1 3.80
                    -10.9
## 10 25.4 3.80
                    -10.9
## # ... with 20,291 more rows
```

Maximizing the log-likelihood gives the same parameter values as maximizing the likelihood. Remember that the log computation keeps the same ordination of values as the original data, so maximizing the log-likelihood is the same as maximizing the likelihood.

## Maximum Likelihood Estimation for Regression

In model fitting, the components we care about are the residuals. Those are the things we put distributional assumptions on (e.g., normality, homogeneity of variance, independent). Our goal in regression is to estimate a set of parameters ( $\beta_0$ ,  $\beta_1$ ) that maximize the likelihood for a given set of residuals that come from a normal distribution.

To understand this, let's use a toy example of n = 10 X and Y values.

```
# Enter data into vectors

x = c(4, 0, 3, 4, 7, 0, 0, 3, 0, 2)

y = c(53, 56, 37, 55, 50, 36, 22, 75, 37, 42)
```

We will write a function to compute the likelihood (or log-likelihood) of the residuals given a particular b0 and b1 estimate that will be inputted to the function. One issue is that in using the dnorm() function we need to specify the mean and standard deviation. The regression assumptions help with this.

The conditional mean residual value is 0. So we will set the mean value to 0. The asumption about the standard deviation is that the conditional distributions all have the same SD, but it doesn't specify what that is. However, the SD of the errors seems like a reasonable value, so let's use that.

Below, we will write a function called log\_likelihood() that takes two arguments as input, b0= and b1=, and outputs the log-likelihood.

```
log_likelihood = function(b0, b1){
    # Use the following x and y values
    x = c(4, 0, 3, 4, 7, 0, 0, 3, 0, 2)
    y = c(53, 56, 37, 55, 50, 36, 22, 75, 37, 42)

# Compute the yhat and residuals based on the two input values
    yhats = b0 + b1*x
    errors = y - yhats

# Compute the sd of the residuals
    sigma = sd(errors)

# Compute the log-likelihood
    log_lik = sum(dnorm(errors, mean = 0, sd = sigma, log = TRUE))

# Output the log-likelihood
    return(log_lik)
}
```

Now we read in our function by highlighting the whole thing and running it. Once it has been read in, we can use it just like any other function. For example to find the log-likelihood for the parameters  $\beta_0 = 10$  and  $\beta_1 = 3$  we use:

```
log_likelihood(b0 = 10, b1 = 3)
```

```
## [1] -64.29224
```

We can also use our function in a grid search.

```
expand.grid(
  b0 = seq(from = 30, to = 50, by = 0.1),
  b1 = seq(from = -5, to = 5, by = 0.1)
) %>%
  rowwise() %>%
  mutate( log_lik = log_likelihood(b0 = b0, b1 = b1) ) %>%
  arrange(desc(log_lik))
```

```
## # A tibble: 20,301 x 3
##
               b1 log_lik
        b0
      <dbl> <dbl>
                    <dbl>
##
##
   1
      40.1
            2.70
                   -39.5
##
   2 40.0
            2.70
                    -39.5
            2.70
   3
      40.2
                    -39.5
##
      39.9
            2.80
                    -39.5
##
   4
##
   5 39.8 2.80
                    -39.5
##
   6 40.0
            2.80
                    -39.5
##
   7
      39.9
            2.70
                    -39.5
##
   8
      39.7
            2.80
                    -39.5
##
  9 40.3
            2.70
                    -39.5
## 10 40.1
            2.80
                    -39.5
## # ... with 20,291 more rows
```

Here the parameter values that maximize the likelihood are  $\beta_0 = 40.1$  and  $\beta_1 = 2.7$ . We can also compute what the standard deviation value was using the estimated parameter values.

```
errors = y - 40.1 - 2.7*x sd(errors)
```

```
## [1] 13.18665
```

This value is an estimate of the RMSE.

In practice, there are a couple subtle differences, namely that the estimate for the SD value we use in dnorm() is slightly different. This generally does not have an effect on the coefficient estimates, but does impact the estimate of the RMSE. We will talk more about this when we talk about *Restricted Maximum Likelihood Estimation* (REML).

### Large Search Spaces

So far, we have been using a very finite search space that has been defined for us. For example, we limited the search space to 20,301 combinatins of  $\beta_0$  and  $\beta_1$ .

```
nrow(
  expand.grid(
    b0 = seq(from = 30, to = 50, by = 0.1),
    b1 = seq(from = -5, to = 5, by = 0.1)
)
)
```

```
## [1] 20301
```

This allowed us to find the coefficient estimates to the nearest tenth. If we instead needed to find the estimates to the nearest hundredth, we would need to expand the number of combinations:

```
nrow(
  expand.grid(
```

```
b0 = seq(from = 30, to = 50, by = 0.01),
b1 = seq(from = -5, to = 5, by = 0.01)
)
```

#### ## [1] 2003001

This leads to a search space of 2,003,001 parameter combinations. If we need them to the nearest thousandth, the search space is 200,030,001 combinations. Furthermore, in practice you would not have any idea which values of  $\beta_0$  and  $\beta_1$  to limit the search space to.

Essentially you would need to search an infinite number of values unless you could limit the search space in some way. For many common methods (e.g., linear regression) finding the ML estimates is mathematically pretty easy (if we know calculus; see *Way, Way, Way too Much Mathematics* below). For more complex methods (e.g., mixed-effect models) there is not a mathematical solution. Instead, mathematics is used to limit the search space and then a grid search is used to hone in on the estimates.

## ML Estimation in Regression Using R

Recall that the lm() function uses Ordinary Least Squares (OLS) estimation—it finds the coefficient estimates and RMSE that minimize the sum of squared residuals.

```
lm.1 = lm(y \sim 1 + x)
# Get coefficient estimates
tidy(lm.1)
            term estimate std.error statistic
                                                    p.value
## 1 (Intercept) 40.005988 6.341633 6.308468 0.0002306738
              x 2.736527 1.975980 1.384896 0.2034785827
# Get estimate for RMSE
glance(lm.1)
    r.squared adj.r.squared
                                sigma statistic p.value df
                                                                logLik
                  0.09255326 13.98625 1.917937 0.2034786 2 -39.45442
## 1 0.1933807
##
         AIC
                  BIC deviance df.residual
## 1 84.90883 85.81659 1564.922
Under OLS estimation:
```

- $\hat{\beta}_0 = 40.01$
- $\hat{\beta}_1 = 2.74$
- $\hat{\sigma}_{\epsilon} = 13.99$

To compute ML estimates of the coefficients we will use the mle2() function from the **bbmle** package. To use the mle2() function, we need to provide a user-written function that returns the *negative log-likelihood* given a set of parameter inputs.

For simple regression, recall that we need to estimate three parameters:  $\beta_0$ ,  $\beta_1$ , and  $\sigma_{\epsilon}$  (RMSE). Below we have a function that inputs the three parameters, uses the and returns the negative of the log-likelihood for a simple regression.

```
regress.ll = function(b0, b1, rmse) {

# Use the following x and y values

x = c(4, 0, 3, 4, 7, 0, 0, 3, 0, 2)

y = c(53, 56, 37, 55, 50, 36, 22, 75, 37, 42)
```

```
# Compute yhats and residuals
  yhats = b0 + b1 * x
  errors = y - yhats
  # Compute the negative log-likelihood
  neg_log_lik = -sum(dnorm(errors, mean = 0, sd = rmse, log = TRUE))
  return(neg_log_lik)
}
```

Now we can implement the mle2() function. This function requires the argument, minuslogl=, which gives the user written function returning the negative log-likelihood. It also requires a list of starting values for the input parameters in the user-written function. (Here we give values close to the OLS estimates as starting values.)

```
# Fit model using ML
library(bbmle)
mle.results = mle2(minuslog1 = regress.11, start = list(b0 = 40.0, b1 = 2.7, rmse = 13.98))
# View results
summary(mle.results)
## Maximum likelihood estimation
##
## Call:
## mle2(minuslogl = regress.ll, start = list(b0 = 40, b1 = 2.7,
       rmse = 13.98))
##
##
## Coefficients:
        Estimate Std. Error z value
##
                                                 Pr(z)
## b0
         40.0075
                     5.6721 7.0533 0.000000000001747 ***
          2.7361
## b1
                     1.7674 1.5481
                                                0.1216
## rmse 12.5097
                     2.7973 4.4721 0.000007744309040 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## -2 log L: 78.90883
Under ML estimation:
```

- $\hat{\beta}_0 = 40.01$   $\hat{\beta}_1 = 2.74$
- $\hat{\sigma}_{\epsilon} = 12.51$

Comparing the coefficient estimates ( $\hat{\beta}_0$  and  $\hat{\beta}_1$ ) between the two methods of estimation, we find they are quite similar. The estimate of  $\sigma_{\epsilon}$  is different between the two estimation methods (although they are somewhat close in value). This is because in OLS estimation, the estimate of  $\hat{\sigma}_{\epsilon}$  turns out to be,

$$\hat{\sigma}_{\epsilon} = \frac{\left(Y_i - \hat{Y}_i\right)^2}{n - 2},$$

and for ML estimation, the estimate for  $\hat{\sigma}_{\epsilon}$  is,

$$\hat{\sigma}_{\epsilon} = \frac{\left(Y_i - \hat{Y}_i\right)^2}{n},$$

The smaller denominator in OLS results in a higher estimate of the variation. This, in turn, affects the size of the SE estimates for the coefficients (and thus the t- and p-values). When n is large, the differences in the estimates of  $\hat{\sigma}_{\epsilon}$  are minimal and can safely be ignored.

Lastly, we note that the value of  $-2(\log-\text{likelihood})$  is the same for both the ML and OLS estimated models. This is a useful result. It allows us to use 1m() to estimate the coefficients from a model and then use its  $\log-\text{likelihood}$  as if we had fitted the model using ML.

### Using R to Directly Compute the Likelihood and Log-Likelihood

We can use R to directly compute the log-likelihood after we fit a model using the lm() function. To do this, we use the logLik() function.

```
lm.1 = lm(y \sim 1 + x)
logLik(lm.1)
```

```
## 'log Lik.' -39.45442 (df=3)
```

To compute the likelihood, we can use the exp() function to back-transform the log-likelihood to the likelihood (although generally we will work with the log-likelihood).

```
exp(-39.45442)
```

## [1] 0.000000000000000007330998

## Way, Way, Way too Much Mathematics

Let's also expound on this more mathematically. Going back to the mathematical notation, we can specify the liklihood as,

$$\mathcal{L}(\beta_0, \beta_1 | \text{data}) = p(\epsilon_1) \times p(\epsilon_2) \times \ldots \times p(\epsilon_n)$$

where

$$p(\epsilon_i) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left[-\frac{(\epsilon_i - \mu)^2}{2\sigma^2}\right]$$

The regression assumptions specify that each conditional error distribution is normal distributed with a mean of 0 and some variance (that is the same for all conditional error distributions), we can call it  $\sigma^2$ . Substituting these things in to the density function, we get,

$$p(\epsilon_i) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left[-\frac{(\epsilon_i - 0)^2}{2\sigma^2}\right]$$
$$= \frac{1}{\sigma\sqrt{2\pi}} \exp\left[-\frac{(\epsilon_i)^2}{2\sigma^2}\right]$$

Now we use this expression for each of the  $p(\epsilon_i)$  values in the liklihood computation.

$$\mathcal{L}(\beta_0, \beta_1 | \text{data}) = p(\epsilon_1) \times p(\epsilon_2) \times \ldots \times p(\epsilon_n)$$

$$= \frac{1}{\sigma \sqrt{2\pi}} \exp\left[-\frac{\epsilon_1^2}{2\sigma^2}\right] \times \frac{1}{\sigma \sqrt{2\pi}} \exp\left[-\frac{\epsilon_2^2}{2\sigma^2}\right] \times \ldots \times \frac{1}{\sigma \sqrt{2\pi}} \exp\left[-\frac{\epsilon_n^2}{2\sigma^2}\right]$$

We can simplify this

$$\mathcal{L}(\beta_0, \beta_1 | \text{data}) = \left[ \frac{1}{\sigma \sqrt{2\pi}} \right]^n \times \exp\left[ -\frac{\epsilon_1^2}{2\sigma^2} \right] \times \exp\left[ -\frac{\epsilon_2^2}{2\sigma^2} \right] \times \dots \times \exp\left[ -\frac{\epsilon_n^2}{2\sigma^2} \right]$$

Now we will take the natural logarithm of both sides of the expression

$$\ln\left(\mathcal{L}(\beta_0, \beta_1 | \text{data})\right) = \ln\left(\left[\frac{1}{\sigma\sqrt{2\pi}}\right]^n \times \exp\left[-\frac{\epsilon_1^2}{2\sigma^2}\right] \times \exp\left[-\frac{\epsilon_2^2}{2\sigma^2}\right] \times \dots \times \exp\left[-\frac{\epsilon_n^2}{2\sigma^2}\right]\right)$$

Using our rules for logarithms and re-arranging gives,

$$\updownarrow(\beta_0, \beta_1|\text{data}) = -\frac{n}{2} \times \ln(2\pi\sigma^2) - \frac{1}{2\sigma^2} \times \sum_{i=1}^{n} \epsilon_i^2$$

The log-liklihood is a function of n,  $\sigma^2$  and the SSE. The data define n (the sample size) and the other two come from the error terms which are a function of the parameters and the data.

Once we have this function, calculus can be used to find the analytic maximum. Typically before we do this, we replace  $\epsilon_i$  with  $Y_i - \beta_0 - \beta_1(X_i)$ . Then the partial derivatives w.r.t.  $\beta_0$  and  $\beta_1$  are both compued, set equal to zero, and solved for the respective parameter.