Foundations for statistical inference - Confidence intervals

Jimmy Ng

Sampling from Ames, Iowa

If you have access to data on an entire population, say the size of every house in Ames, Iowa, it's straight forward to answer questions like, "How big is the typical house in Ames?" and "How much variation is there in sizes of houses?". If you have access to only a sample of the population, as is often the case, the task becomes more complicated. What is your best guess for the typical size if you only know the sizes of several dozen houses? This sort of situation requires that you use your sample to make inference on what your population looks like.

The data

In the previous lab, "Sampling Distributions", we looked at the population data of houses from Ames, Iowa. Let's start by loading that data set.

```
load("more/ames.RData")
```

In this lab we'll start with a simple random sample of size 60 from the population. Specifically, this is a simple random sample of size 60. Note that the data set has information on many housing variables, but for the first portion of the lab we'll focus on the size of the house, represented by the variable Gr.Liv.Area.

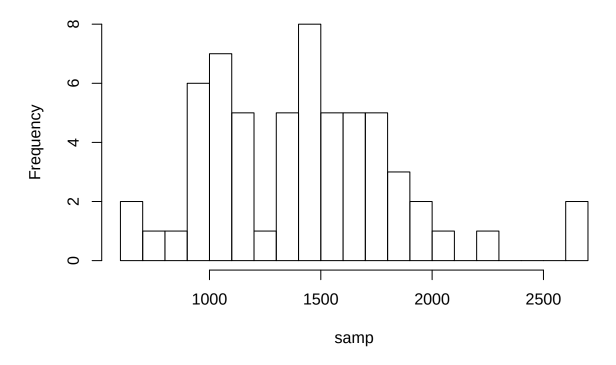
```
population <- ames$Gr.Liv.Area
samp <- sample(population, 60)</pre>
```

1. Describe the distribution of your sample. What would you say is the "typical" size within your sample? Also state precisely what you interpreted "typical" to mean.

```
shapiro.test(samp)
```

```
##
## Shapiro-Wilk normality test
##
## data: samp
## W = 0.95883, p-value = 0.04125
hist(samp, breaks = 15)
```

Histogram of samp



```
summary(samp)
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 630 1074 1436 1418 1667 2687
sd(samp)
```

[1] 426.0746

JN: The shapiro test is used to test whether the data is normally distributed and the result (p value > .05) suggested that it is normally distributed. The histogram also depicted a nearly normal distribution with slight skew on the right. The summary statistics as well as the SD also pointed to the same conclusion. With the mean and median in the center with equal spread on both sides, the distribution is normally distributed.

2. Would you expect another student's distribution to be identical to yours? Would you expect it to be similar? Why or why not?

JN: I expect that another student's distribution would be similar but not identical to "samp". There will always be variation when drawing random sample; however, the sampling distribution for each draw should be highly similar as long as the sampling is random and independent.

Confidence intervals

One of the most common ways to describe the typical or central value of a distribution is to use the mean. In this case we can calculate the mean of the sample using,

```
sample_mean <- mean(samp)</pre>
```

Return for a moment to the question that first motivated this lab: based on this sample, what can we infer about the population? Based only on this single sample, the best estimate of the average living area of houses sold in Ames would be the sample mean, usually denoted as \bar{x} (here we're calling it sample_mean). That serves as a good *point estimate* but it would be useful to also communicate how uncertain we are of that estimate. This can be captured by using a *confidence interval*.

We can calculate a 95% confidence interval for a sample mean by adding and subtracting 1.96 standard errors to the point estimate (See Section 4.2.3 if you are unfamiliar with this formula).

```
se <- sd(samp) / sqrt(60)
lower <- sample_mean - 1.96 * se
upper <- sample_mean + 1.96 * se
c(lower, upper)</pre>
```

```
## [1] 1310.622 1526.245
```

This is an important inference that we've just made: even though we don't know what the full population looks like, we're 95% confident that the true average size of houses in Ames lies between the values *lower* and *upper*. There are a few conditions that must be met for this interval to be valid.

3. For the confidence interval to be valid, the sample mean must be normally distributed and have standard error s/\sqrt{n} . What conditions must be met for this to be true?

JN: each sample must be randomly and independently selected in each sampling; less than 10% of the population is chosen in each sampling; and the size should be large enough (e.g. $n \ge 30$) to ignore any minor skew.

Confidence levels

4. What does "95% confidence" mean? If you're not sure, see Section 4.2.2.

JN: we are confident that 95% of the time the true population mean would fall between the lower and upper interval. In other words, we commit ourselves to make type 1 error about 5% of the time, i.e. the true mean is actually outside the intervales 5% of the time.

In this case we have the luxury of knowing the true population mean since we have data on the entire population. This value can be calculated using the following command:

```
mean(population)
## [1] 1499.69
```

5. Does your confidence interval capture the true average size of houses in Ames? If you are working on this lab in a classroom, does your neighbor's interval capture this value?

JN: Yes, it is captured in between the lower and upper intervals.

6. Each student in your class should have gotten a slightly different confidence interval. What proportion of those intervals would you expect to capture the true population mean? Why? If you are working in this lab in a classroom, collect data on the intervals created by other students in the class and calculate the proportion of intervals that capture the true population mean.

JN: 95% of us should capture the true mean in their intervals; in othe words, 5% of the time, we make type 1 error.

Using R, we're going to recreate many samples to learn more about how sample means and confidence intervals vary from one sample to another. *Loops* come in handy here (If you are unfamiliar with loops, review the Sampling Distribution Lab).

Here is the rough outline:

- Obtain a random sample.
- Calculate and store the sample's mean and standard deviation.
- Repeat steps (1) and (2) 50 times.
- Use these stored statistics to calculate many confidence intervals.

But before we do all of this, we need to first create empty vectors where we can save the means and standard deviations that will be calculated from each sample. And while we're at it, let's also store the desired sample size as n.

```
samp_mean <- rep(NA, 50)
samp_sd <- rep(NA, 50)
n <- 60</pre>
```

Now we're ready for the loop where we calculate the means and standard deviations of 50 random samples.

```
set.seed(123)
for(i in 1:50){
  samp <- sample(population, n) # obtain a sample of size n = 60 from the population
  samp_mean[i] <- mean(samp) # save sample mean in ith element of samp_mean
  samp_sd[i] <- sd(samp) # save sample sd in ith element of samp_sd
}</pre>
```

Lastly, we construct the confidence intervals.

```
lower_vector <- samp_mean - 1.96 * samp_sd / sqrt(n)
upper_vector <- samp_mean + 1.96 * samp_sd / sqrt(n)</pre>
```

Lower bounds of these 50 confidence intervals are stored in lower_vector, and the upper bounds are in upper_vector. Let's view the first interval.

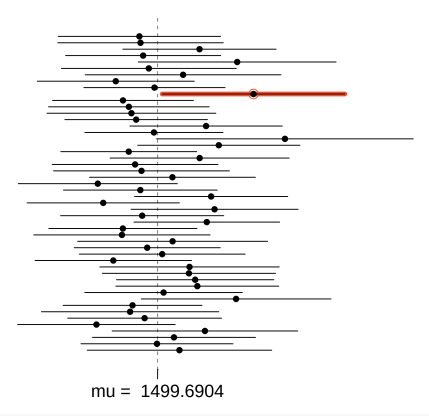
```
c(lower_vector[1], upper_vector[1])
## [1] 1390.046 1677.388
```

On your own

[1] 98

• Using the following function (which was downloaded with the data set), plot all intervals. What proportion of your confidence intervals include the true population mean? Is this proportion exactly equal to the confidence level? If not, explain why.

```
plot_ci(lower_vector, upper_vector, mean(population))
```



```
df <- data.frame(lower_vector, upper_vector, m = rep(mean(population), length(upper_vector)))
(sum(with(df, ifelse(lower_vector < m & upper_vector > m, 1, 0))) / 50) * 100
```

JN: [remember to set.seed(123)] About "(sum(with(df, ifelse(lower_vector < m & upper_vector > m, 1, 0))) / 50) * 100" percent of confidence intervals (CI) included the true population mean. That is, about 98%. That matches with the alpha level (p-value) and CI that we pick (95%) - in other words, we assume a 95% confidence interval for our sampling distribution and that means we expect to make a mistake (type 1 error) about 5 or less percent of the time. We control the error (type 1) about 5 percent or less, not exactly equal to 5%. In above example, 98% of our CI included the true population mean! We make 2% error (i.e. 1 out of 50).

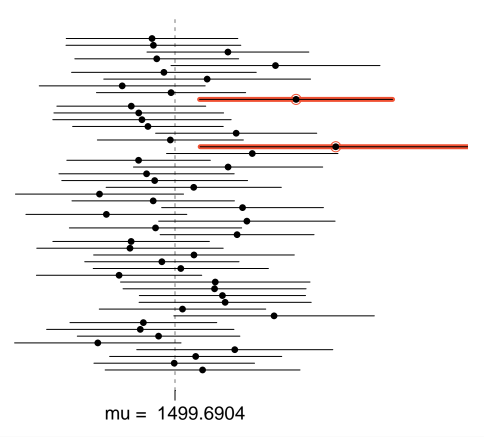
• Pick a confidence level of your choosing, provided it is not 95%. What is the appropriate critical value?

JN: let's pick a confidence leel of 90%. The p-value is 0.1 and the corresponding critical value is 1.64.

• Calculate 50 confidence intervals at the confidence level you chose in the previous question. You do not need to obtain new samples, simply calculate new intervals based on the sample means and standard deviations you have already collected. Using the plot_ci function, plot all intervals and calculate the proportion of intervals that include the true population mean. How does this percentage compare to the confidence level selected for the intervals?

```
lower_vector <- samp_mean - 1.64 * samp_sd / sqrt(n)
upper_vector <- samp_mean + 1.64 * samp_sd / sqrt(n)

plot_ci(lower_vector, upper_vector, mean(population))</pre>
```



```
df <- data.frame(lower_vector, upper_vector, m = rep(mean(population), length(upper_vector)))
(sum(with(df, ifelse(lower_vector < m & upper_vector > m, 1, 0))) / 50) * 100
## [1] 96
```

JN: [remember to set.seed(123)] About "(sum(with(df, ifelse(lower_vector < m & upper_vector > m, 1, 0))) / 50) * 100" percent of confidence intervals (CI) included the true population mean. That is, about 96%. Now, the accuracy declined after we loose up (or lower the threshold) the alpha level. We are allowed to make roughly 10% or less mistake and we did. Lowering the threshold (from 5% to 10%) would lead to higher chance of type 1 errors, but in exchange, we will get a set of smaller/tigher intervals.

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