Confidence Intervals Notes

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Objectives

- 1) Using asymptotic method assuming a normal, obtain and interpret a confidence interval for an unknown parameter, based on a random sample.
- 2) Describe the relationships between confidence intervals, confidence level, and sample size.

Confidence interval

A point estimate provides a single plausible value for a parameter. However, a point estimate is rarely perfect; usually there is some error in the estimate. In addition to supplying a point estimate of a parameter, a next logical step would be to provide a plausible **range of values** for the parameter.

Capturing the population parameter

A plausible range of values for the population parameter is called a **confidence interval**. Using only a point estimate is like fishing in a murky lake with a spear, and using a confidence interval is like fishing with a net. We can throw a spear where we saw a fish, but we will probably miss. On the other hand, if we toss a net in that area, we have a good chance of catching the fish.

If we report a point estimate, we probably will not hit the exact population parameter. On the other hand, if we report a range of plausible values – a confidence interval – we have a good shot at capturing the parameter.

Exercise: If we want to be very certain we capture the population parameter, should we use a wider interval or a smaller interval?¹

Constructing a confidence interval

A point estimate is our best guess for the value of the parameter, so it makes sense to build the confidence interval around that value. The standard error, which is a measure of the uncertainty associated with the point estimate, provides a guide for how large we should make the confidence interval.

Generally, what you should know about building confidence intervals is laid out in the following steps:

- 1. Identify the parameter you would like to estimate (for example, μ).
- 2. Identify a good estimate for that parameter (sample mean, \bar{X}).

¹If we want to be more certain we will capture the fish, we might use a wider net. Likewise, we use a wider confidence interval if we want to be more certain that we capture the parameter.

- 3. Determine the distribution of your estimate or a function of your estimate. This tells us where our estimate should be if we knew the value of our parameter. (According to the central limit theorem, $\frac{\bar{X}-\mu}{\sigma/\sqrt{n}} \sim \mathsf{Norm}(0,1)$ and $\frac{\bar{X}-\mu}{S/\sqrt{n}} \sim \mathsf{t}(n-1)$).
- 4. Use this distribution to obtain a range of feasible values (confidence interval) for the parameter. (For μ , we can solve for μ to find a reasonable range).

Constructing a 95% confidence interval for the mean

When the sampling distribution of a point estimate can reasonably be modeled as normal, the point estimate we observe will be within 1.96 standard errors of the true value of interest about 95% of the time. Thus, a 95% confidence interval for such a point estimate can be constructed:

$$\hat{\theta} \pm 1.96 \times SE_{\hat{\theta}}$$

We can be 95% confident this interval captures the true value.

Exercise:

Compute the area between -1.96 and 1.96 for a normal distribution with mean 0 and standard deviation 1.

$$pnorm(1.96) - pnorm(-1.96)$$

[1] 0.9500042

In mathematical terms, the derivation of this confidence is as follows:

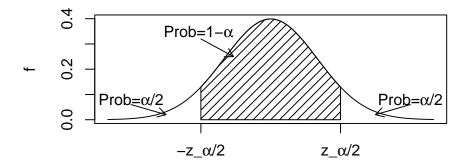
Let $X_1, X_2, ..., X_n$ be an iid sequence of random variables, each with mean μ and standard deviation σ . The central limit theorem tells us that

$$\frac{\bar{X} - \mu}{\sigma / \sqrt{n}} \stackrel{approx}{\sim} \mathsf{Norm}(0, 1)$$

Let $0 \le \alpha \le 1$ with the confidence level being $1 - \alpha$, yes α is the same as the significance level in hypothesis testing. Then,

$$P\left(-z_{\alpha/2} \le \frac{\bar{X} - \mu}{\sigma/\sqrt{n}} \le z_{\alpha/2}\right) = 1 - \alpha$$

where $z_{\alpha/2}$ is such that $P(Z \ge z_{\alpha/2}) = \alpha/2$, where $Z \sim \mathsf{Norm}(0,1)$. A picture would help:



So, I know that $(1-\alpha)*100\%$ of the time, $\frac{\bar{X}-\mu}{\sigma/\sqrt{n}}$ will be between $-z_{\alpha/2}$ and $z_{\alpha/2}$.

By rearranging the expression above and solving for μ , we get:

$$P\left(\bar{X} - z_{\alpha/2} \frac{\sigma}{\sqrt{n}} \le \mu \le \bar{X} + z_{\alpha/2} \frac{\sigma}{\sqrt{n}}\right) = 1 - \alpha$$

Be careful with the interpretation of this expression. As a reminder \bar{X} is the random variable here. The population mean, μ , is NOT a variable. It is an unknown parameter. Thus, the above expression is NOT a probabilistic statement about μ , but rather about \bar{X} .

Nonetheless, the above expression gives us a nice interval for "reasonable" values of μ given a particular sample.

A $(1 - \alpha) * 100\%$ confidence interval for the mean is given by:

$$\mu \in \left(\bar{X} \pm z_{\alpha/2} \frac{\sigma}{\sqrt{n}}\right)$$

In most applications, the most common value of α is 0.05. In that case, to construct a 95% confidence interval, we would need to find $z_{0.025}$ which can be found quickly with qnorm():

qnorm(1-0.05/2)

[1] 1.959964

qnorm(.975)

[1] 1.959964

Unknown Variance When inferring about the population mean, we usually will have to estimate the underlying standard deviation as well. This introduces an extra level of uncertainty. We found that while $\frac{\bar{X}-\mu}{\sigma/\sqrt{n}}$ has an approximate normal distribution, $\frac{\bar{X}-\mu}{S/\sqrt{n}}$ follows the t-distribution with n-1 degrees of freedom. This adds the additional assumption that the parent population, the distribution of X, must be normal.

Thus, when σ is unknown, a $(1-\alpha)*100\%$ confidence interval for the mean is given by:

$$\mu \in \left(\bar{X} \pm t_{\alpha/2, n-1} \frac{s}{\sqrt{n}}\right)$$

Similar to the case above, $t_{\alpha/2,n-1}$ can be found using the qt() function in R.

In practice, if X is close to symmetrical and unimodal, we can relax the assumption of normality. Always look at your sample data. Outliers or skewness can be causes of concern. You can always run other methods that don't require the assumption of normality, and compare results.

For large sample sizes, the choice of using the normal distribution or the t distribution is irrelevant since they are close to each other. The t distribution requires you to use the degrees of freedom so be careful.

Body Temperature Example

Example:

Find a 95% confidence interval for the body temperature data from last lesson.

We need the mean, standard deviation, and sample size from this data. The following R calculates the confidence interval, make sure you can follow the code.

```
## lower_bound upper_bound
## 1 98.122 98.37646
```

The 95% confidence interval for μ is (98.12, 98.38). I am 95% confident that μ , the average human body temperature, is in this interval. Also, we could say that 95% of similarly constructed intervals will contain the true mean, μ .

Remember that when we used this data in a the hypothesis test, the null hypothesis was H_0 :] The average body temperature is 98.6 μ = 98.6. This null hypothesized value is not in the interval, so we could reject the null hypothesis with this confidence interval. There is a link between hypothesis testing and confidence intervals.

We could also use R' to find the confidence interval.

```
t_test(~temperature,data=temperature)
```

```
##
## One Sample t-test
##
## data: temperature
## t = 1527.9, df = 129, p-value < 2.2e-16
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## 98.12200 98.37646
## sample estimates:
## mean of x
## 98.24923</pre>
```

Or if you just want the interval:

```
confint(t_test(~temperature,data=temperature))
```

```
## mean of x lower upper level
## 1 98.24923 98.122 98.37646 0.95
```

In reviewing the hypothesis test for a single mean, you can see how this confidence interval was formed by *inverting* the test statistic.

One-sided Intervals

If you remember the hypothesis test for temperature in the central limit theorem lesson, you may be crying foul. That was a one-sided hypothesis test and we just conducted a two-sided test. So far, we have discussed

only "two-sided" intervals. These intervals have an upper and lower bound. Typically, α is apportioned equally between the two tails. (Thus, we look for $Z_{\alpha/2}$.)

In "one-sided" intervals, we only bound the interval on one side. We construct one-sided intervals when we are concerned with whether a parameter exceeds or stays below some threshold. Building a one-sided interval is similar to building two-sided intervals, except rather than dividing α into two, you simply apportion all of α to the relevant side. The difficult part is to determine if I need an upper bound or lower bound.

For the body temperature study, the alternative hypothesis was that the mean was less than 98.6. In our confidence interval, we want to find the largest value the mean could be and thus we want the upper bound. Repeating the analysis with this in mind.

```
temperature %>%
  favstats(~temperature,data=.) %>%
  select(mean,sd,n) %>%
  summarise(upper_bound=mean+qt(0.95,129)*sd/sqrt(n))

## upper_bound
## 1 98.35577

confint(t_test(~temperature,data=temperature,alternative="less"))

## mean of x lower upper level
## 1 98.24923 -Inf 98.35577 0.95
```

Notice that upper bound is smaller since all 0.05 is going into the right tail.

Confidence intervals for two proportions

191

405

33

46

224

451

##

##

trmt Total

In hypothesis testing we had several examples of two proportions. We tested this with a permutation test. In our notes or homework, we have not presented the hypothesis test for two proportions using the asymptoic distribution, the central limit theorem. So in this section we will present three methods to answering our research question, a permutation test, a hypothesis test using the normal distribution, and a confidence interval.

Earlier this semester, in fact in the first lesson, we encountered an experiment that examined whether implanting a stent in the brain of a patient at risk for a stroke helps reduce the risk of a stroke. The results from the first 30 days of this study, which included 451 patients, are summarized in the R code below. These results are surprising! The point estimate suggests that patients who received stents may have a **higher** risk of stroke: $p_{trmt} - p_{control} = 0.090$.

```
stent <- read_csv("data/stent_study.csv")

tally(~group+outcome30,data=stent,margins = TRUE)

## outcome30
## group no_event stroke Total
## control 214 13 227</pre>
```

```
tally(outcome30~group,data=stent,margins = TRUE,format="proportion")
```

```
## group
## outcome30 control trmt
## no_event 0.94273128 0.85267857
## stroke 0.05726872 0.14732143
## Total 1.00000000 1.00000000

obs<-diffprop(outcome30~group,data=stent)
obs</pre>
```

```
## diffprop
## -0.09005271
```

Notice that because uses the variables by names in alphabetic order we have $p_{control} - p_{trmt} = -0.090$. This is not a problem.

Permutation test for two proportions

We start with the null hypothesis which is two-sided since we don't know if the treatment is harmful or beneficial.

```
H_0: The treatment and outcome are independent. p_{control} - p_{trmt} = 0 or p_{control} = p_{trmt}.
```

 H_A : The treatment and outcome are dependent $p_{control} \neq p_{trmt}$.

We will use $\alpha = 0.05$.

The test statistic is the difference in proportions of patients with stroke in the control and treatment groups.

```
obs<-diffprop(outcome30~group,data=stent)
obs</pre>
```

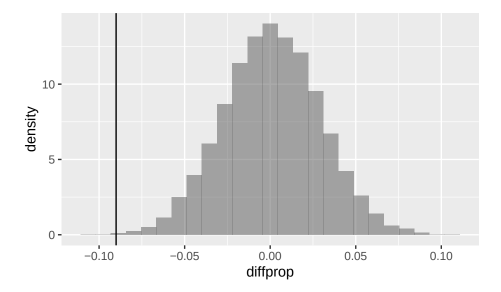
```
## diffprop
## -0.09005271
```

To calculate the p-value, we will shuffle the treatment and control labels because under the null hypothesis, there is no difference.

```
set.seed(2027)
results <- do(10000)*diffprop(outcome30~shuffle(group),data=stent)</pre>
```

We now create a visual summary of the test statistic under the null hypothesis, the sampling distribution.

```
results %>%
  gf_dhistogram(~diffprop) %>%
  gf_vline(xintercept =obs )
```



2*prop1(~(diffprop<=obs),data=results)</pre>

```
## prop_TRUE
## 0.00259974
```

Based on the data, if there were no difference between the treatment and control groups, the probability of the observed differences in proportion of strokes being - 0.09 or more extreme is 0.0026. This is too unlikely, so we reject that there is no difference between control and stroke groups.

Hypothesis test for two proportions using normal model

We must check two conditions before applying the normal model to a generic test of $\hat{p}_1 - \hat{p}_2$. First, the sampling distribution for each sample proportion must be nearly normal, and secondly, the samples must be independent. Under these two conditions, the sampling distribution of $\hat{p}_1 - \hat{p}_2$ may be well approximated using the normal model.

The hypotheses are the same as above.

Conditions for the sampling distribution of $\hat{p}_1 - \hat{p}_2$ to be normal The difference $\hat{p}_1 - \hat{p}_2$ tends to follow a normal model when

- each proportion separately follows a normal model, and
- the two samples are independent of each other.

Standard error The standard error of the difference in sample proportions is

$$SE_{\hat{p}_1-\hat{p}_2} = \sqrt{SE_{\hat{p}_1}^2 + SE_{\hat{p}_2}^2}$$

$$= \sqrt{\frac{p_1(1-p_1)}{n_1} + \frac{p_2(1-p_2)}{n_2}}$$

where p_1 and p_2 represent the population proportions, and n_1 and n_2 represent the sample sizes.

For our research question the conditions must be verified. Because each group is a simple random sample from less than 10% of the population, the observations are independent, both within the samples and between the samples. The success-failure condition also holds for each sample, at least 5 in each cell is the easiest way to think about it. Because all conditions are met, the normal model can be used for the point estimate of the difference in proportion of strokes

$$p_{control} - p_{trmt} = 0.05726872 - 0.14732143 = -0.090$$

The calculation of the standard error must be done carefully. Remember in hypothesis testing, we assume the null hypothesis is true; this means the proportions of strokes must be the same.

$$SE = \sqrt{\frac{p(1-p)}{n_{control}} + \frac{p(1-p)}{n_{trmt}}}$$

We don't know the exposure rate, p, but we can obtain a good estimate of it by **pooling** the results of both samples:

$$\hat{p} = \frac{\text{# of "successes"}}{\text{# of cases}} = \frac{13 + 33}{451} = 0.102$$

This is called the *pooled estimate* of the sample proportion, and we use it to compute the standard error when the null hypothesis is that $p_{control} = p_{trmt}$.

$$SE \approx \sqrt{\frac{\hat{p}(1-\hat{p})}{n_{control}} + \frac{\hat{p}(1-\hat{p})}{n_{trmt}}}$$

 \mathbf{m}

$$SE \approx \sqrt{\frac{0.102(1 - 0.102)}{227} + \frac{0.102(1 - 0.102)}{224}} = 0.0285$$

The test statistic is

$$Z = \frac{\text{point estimate} - \text{null value}}{SE} = \frac{-.09 - 0}{0.0285} = -3.16$$

The p-value is

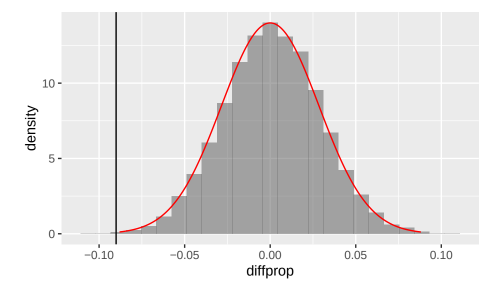
```
2*pnorm(-3.16)
```

[1] 0.001577691

Which is close to what we got with permutation test. This should not surprise us as the sampling distribution under the permutation test looked normal. We plot the empirical sampling distribution from the permutation test again with a normal density curve overlayed.

```
results %>%
  gf_dhistogram(~diffprop) %>%
  gf_vline(xintercept =obs ) %>%
  gf_dist("norm",sd=0.0285,color="red")
```

Warning: geom_vline(): Ignoring `mapping` because `xintercept` was provided.



Confidence interval for two proportions using normal model

The conditions for applying the normal model have already been verified, so we can proceed to the construction of the confidence interval. Remember the form of the confidence interval is

point estimate
$$\pm z^*SE$$

Our point estimate is -0.09. The standard error is different since we can't assume the proportion of strokes are equal. We will estimate the standard error from

$$SE = \sqrt{\frac{p_{control}(1 - p_{control})}{n_{control}} + \frac{p_{trmt}(1 - p_{trmt})}{n_{trmt}}}$$

$$SE \approx \sqrt{\frac{0.057(1 - 0.057)}{227} + \frac{0.15(1 - 0.15)}{224}} = 0.0284$$

It is close to the pooled value because of the nearly equal sample sizes.

The critical value is found from the normal quantile.

qnorm(.975)

[1] 1.959964

The 95% confidence interval is

$$-0.09 \pm 1.96 \times 0.0284 \rightarrow (-0.146, -0.034)$$

We are 95% confident that the difference in proportions of strokes in the control and treatment groups is between -0.146 and -0.034. Since this does not include zero, we are confident they are different. This supports the hypothesis tests.

Of course, R has a built in function to calculate the hypothesis test and confidence interval for two proportions.

prop_test(outcome30~group,data=stent)

```
##
## 2-sample test for equality of proportions with continuity correction
##
## data: tally(outcome30 ~ group)
## X-squared = 9.0233, df = 1, p-value = 0.002666
## alternative hypothesis: two.sided
## 95 percent confidence interval:
## 0.03022922 0.14987619
## sample estimates:
## prop 1 prop 2
## 0.9427313 0.8526786
```

The p-value is a little different from the one we calculated and closer to the permutation test, which is an exact test, because a correction factor was applied. Read online about this correction to learn more. We run the code below with the correction factor off and get the same p-value. The confidence interval is a little different because the function used no stroke as its event, but since zero is not in the interval, we get the same conclusion.

```
prop_test(outcome30~group,data=stent,correct=FALSE)
```

```
##
   2-sample test for equality of proportions without continuity
##
##
   correction
##
## data: tally(outcome30 ~ group)
## X-squared = 9.9823, df = 1, p-value = 0.001581
## alternative hypothesis: two.sided
## 95 percent confidence interval:
## 0.03466401 0.14544140
## sample estimates:
##
      prop 1
                prop 2
## 0.9427313 0.8526786
```

Essentially, confidence intervals and hypothesis tests serve similar purposes, but answer slightly different questions. A confidence interval gives you a range of feasible values of a parameter given a particular sample. A hypothesis test tells you whether a specific value is feasible given a sample. Sometimes you can informally conduct a hypothesis test simply by building an interval and observing whether the hypothesized value is contained in the interval. The disadvantage to this approach is that it does not yield a specific p-value. The disadvantage of the hypothesis test is that it does not give a range of values for the test statistic.

As with hypothesis tests, confidence intervals are imperfect. About 1-in-20 properly constructed 95% confidence intervals will fail to capture the parameter of interest. This is similar idea to our Type 1 error.

Changing the confidence level

Suppose we want to consider confidence intervals where the confidence level is somewhat higher than 95%: perhaps we would like a confidence level of 99%. Think back to the analogy about trying to catch a fish: if we want to be more sure that we will catch the fish, we should use a wider net. To create a 99% confidence

level, we must also widen our 95% interval. On the other hand, if we want an interval with lower confidence, such as 90%, we could make our original 95% interval slightly slimmer.

The 95% confidence interval structure provides guidance in how to make intervals with new confidence levels. Below is a general 95% confidence interval for a point estimate that comes from a nearly normal distribution:

```
point estimate \pm 1.96 \times SE
```

There are three components to this interval: the point estimate, "1.96", and the standard error. The choice of $1.96 \times SE$, which is also called **margin of error**, was based on capturing 95% of the data since the estimate is within 1.96 standard errors of the true value about 95% of the time. The choice of 1.96 corresponds to a 95% confidence level.

Exercise: If X is a normally distributed random variable, how often will X be within 2.58 standard deviations of the mean?²

To create a 99% confidence interval, change 1.96 in the 95% confidence interval formula to be 2.58.

The normal approximation is crucial to the precision of these confidence intervals. We will learn a method called the **bootstrap** that will allow us to find confidence intervals without the assumption of normality.

Interpreting confidence intervals

A careful eye might have observed the somewhat awkward language used to describe confidence intervals.

Correct interpretation:

We are XX% confident that the population parameter is between...

Incorrect language might try to describe the confidence interval as capturing the population parameter with a certain probability. This is one of the most common errors: while it might be useful to think of it as a probability, the confidence level only quantifies how plausible it is that the parameter is in the interval.

Another especially important consideration of confidence intervals is that they **only try to capture the population parameter**. Our intervals say nothing about the confidence of capturing individual observations, a proportion of the observations, or about capturing point estimates. Confidence intervals only attempt to capture population parameters.

File Creation Information

• File creation date: 2020-07-21

• Windows version: Windows 10 x64 (build 17763)

R version 3.6.3 (2020-02-29)mosaic package version: 1.6.0

• tidyverse package version: 1.3.0

²This is equivalent to asking how often a standard normal variable will be larger than -2.58 but less than 2.58. To determine this probability, look up -2.58 and 2.58 in R using pnorm() (0.0049 and 0.9951). Thus, there is a 0.9951 - 0.0049 \approx 0.99 probability that the unobserved random variable X will be within 2.58 standard deviations of the mean.