Bootstrap Confidence Intervals

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1 Introduction

In previous weeks we have seen many examples of calculating sample statistics such as means, percentiles, standard deviations and regression coefficients. These sample statistics are used as point estimates of population parameters which describe the population from which the sample of data was taken. That last sentence assumes you're familiar with concepts and terminology about sampling (e.g. from the Statistical Inference course in 1st Semester) so here is a summary of some key terms:

- 1. **Population**: The population is a set of N observations of interest.
- 2. **Population parameter**: A population parameter is a numerical summary value about the population. In most settings, this is a value that's unknown and you wish you knew it.
- 3. **Census**: An exhaustive enumeration/counting of all observations in the population in order to compute the population parameter's numerical value exactly. When N is small, a census is feasible. However, when N is large, a census can get very expensive, either in terms of time, energy, or money.
- 4. Sampling: Collecting a sample of size n of observations from the population. Typically the sample size n is much smaller than the population size N, thereby making sampling a much cheaper procedure than a census. It is important to remember that the lowercase n corresponds to the sample size and uppercase N corresponds to the population size, thus n???N. Point estimates/sample statistics: A summary statistic based on the sample of size n that estimates the unknown population parameter.
- 5. **Representative sampling**: A sample is said be a representative sample if it "looks like the population". In other words, the sample's characteristics are a good representation of the population's characteristics.
- 6. **Generalizability**: We say a sample is generalizable if any results based on the sample can generalize to the population.
- 7. **Bias**: In a statistical sense, we say bias occurs if certain observations in a population have a higher chance of being sampled than others. We say a sampling procedure is unbiased if every observation in a population had an equal chance of being sampled.
- 8. **Random sampling**: We say a sampling procedure is random if we sample randomly from the population in an unbiased fashion.

1.1 Inference via Sampling

The logic of inference via sampling is:

- If the sampling of a sample of size n is done at **random**, then
- The sample is **unbiased** and **representative** of the population, thus
- Any result based on the sample can **generalize** to the population, thus
- The point estimate/sample statistic is an estimate of the unknown population parameter of interest

and thus we have **inferred** something about the population based on our sample.

1.2 Task 1

In 2013 National Public Radio in the USA reported a poll of President Obama's approval rating among young Americans aged 18-29 in an article Poll: Support For Obama Among Young Americans Eroding. Here is a quote from the article:

"After voting for him in large numbers in 2008 and 2012, young Americans are souring on President Obama.

According to a new Harvard University Institute of Politics poll, just 41 percent of millennials (adults ages 18-29) approve of Obama's job performance, his lowest-ever standing among the group and an 11-point drop from April."

Identify each of the following terms in this context. (NB. Do not enter any R code below, but you can access the solution by clicking "Hint".)

- Population: Millennials (Americans aged 18-29)
- Population parameter: The true population proportion p of young Americans who approve of Obama's job performance.
- Census: Young Americans and asking them if they approve of Obama's job performance.
- Sampling: One way is to get phone records from a dtabase and pick out n phone numbers.
- Point estimates/sample statistics: Th sample proportion \hat{p} of young Americans in the sample that approve of Obama's job performance.
- Representative sampling: Does the sampe of n = 2089 young Americans accurately represent the population of all young Americans age 18-29?
- Generalizability: is $\hat{p} = 0.41$ a good estimate of p?
- Bias: Are there any sources of bias in the study? Sampling bias, self-selection bias...
- Random sampling: Was the sampling randomly done?

2 Inference Using Sample Statistics

Scenario	Population parameter	Population Notation	Point estimate/sample statistic	Sample Notation
1	Population proportion	\overline{p}	Sample proportion	\hat{p}
2	Population mean	μ	Sample mean	$ar{x}$
3	Difference in population proportions	$p_1 - p_2$	Difference in sample proporti	ons $\hat{p}_1 - \hat{p}_2$
4	Difference in population means	$\mu_1 - \mu_2$	Difference in sample means	$\bar{x}_1 - \bar{x}_2$
5	Population regression intercept	eta_0	Sample regression intercept	$\hat{eta}_0 \ or \ b_0$
6	Population regression slope	eta_1	Sample regression slope	\hat{eta}_1 or b_1

3 Bootstrapping

The moderndive package contains a sample of 40 pennies collected and minted in the United States. Let's explore this sample data first: The pennies_sample data frame has rows corresponding to a single penny with two variables:

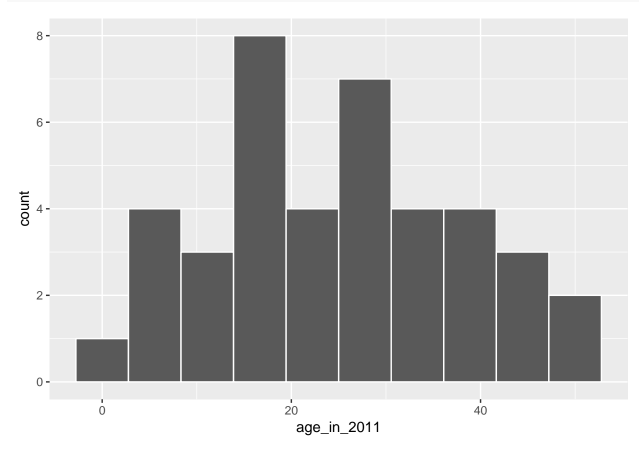
- year of minting as shown on the penny and
- age_in_2011 giving the years the penny had been in circulation in 2011 as an integer, e.g. 15, 2, etc.

Suppose we are interested in understanding some properties of the mean age of all US pennies from this data collected in 2011. How might we go about that? Let's begin by understanding some of the properties of pennies sample using data wrangling from Week 2 and data visualization from Week 1.

3.1 EDA

First, let's visualize the values in this sampl as a histogram:

```
ggplot(pennies_sample, aes(x = age_in_2011)) +
geom_histogram(bins = 10, color = "white")
```



We see a roughly symmetric distribution here that has quite a few values near 20 years in age with only a few larger than 40 years or smaller than 5 years. If pennies_sample is a representative sample from the population, we'd expect the age of all US pennies collected in 2011 to have a similar shape, a similar spread, and similar measures of central tendency like the mean.

So where does the mean value fall for this sample? This point will be known as our **point estimate** and provides us with a single number that could serve as the guess to what the true population mean age might be. Recall how to find this using the dplyr package:

```
x_bar <- pennies_sample %>%
summarize(stat = mean(age_in_2011))
```

We've denoted this sample mean as \hat{x} , which is the standard symbol for denoting the mean of a sample. Our point estimate is, thus, $\hat{x} = 25.1$. Note that this is just one sample providing just one sample mean to estimate the population mean. To construct a **confidence interval** (and to do any sort of *statistical inference* for that matter) we need to know about the **sampling distribution** of this sample mean, i.e. how would its values vary if many samples of the same size were drawn from the same population.

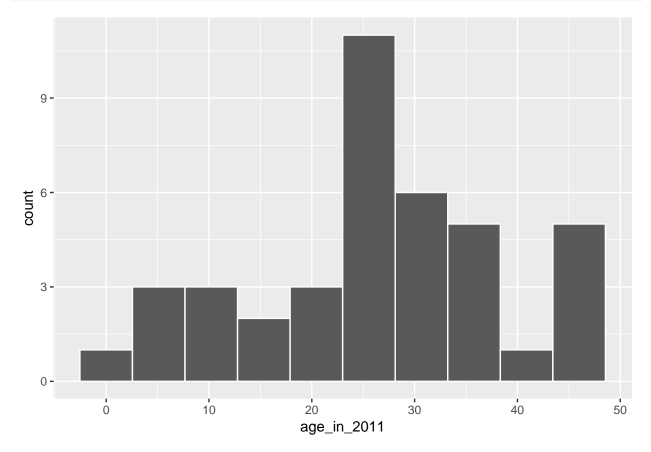
The process of **bootstrapping** allows us to use a single sample to generate many different samples that will act as our way of approximating a sampling distribution using a created **bootstrap distribution** instead. We will "pull ourselves up by our bootstraps" (as the saying goes in English, see here) using a single sample (pennies_sample) to get an idea of the sampling distribution of the sample mean.

3.2 The Bootstrapping Process

Bootstrapping uses a process of sampling with replacement from our original sample to create new bootstrap samples of the *same size* as our original sample. We can use the rep_sample_n() function in the infer package to explore what one such bootstrap sample would look like. Remember that we are randomly sampling from the original sample here with replacement and that we always use the same sample size for the bootstrap samples as the size of the original sample (pennies_sample).

```
bootstrap_sample1 <- pennies_sample %>%
    rep_sample_n(size = 40, replace = TRUE, reps = 1)

ggplot(bootstrap_sample1, aes(x = age_in_2011)) +
    geom_histogram(bins = 10, color = "white")
```



We now have another sample from what we could assume comes from the population of interest. We can similarly calculate the sample mean of this bootstrap sample, called a **bootstrap statistic**.

We'll come back to analyzing the variation in the values of different bootstrap samples' statistics shortly. But first, let's recap what was done to get to this single bootstrap sample using a tactile explanation:

1. First, pretend that each of the 40 values of age_in_2011 in pennies_sample were written on a small

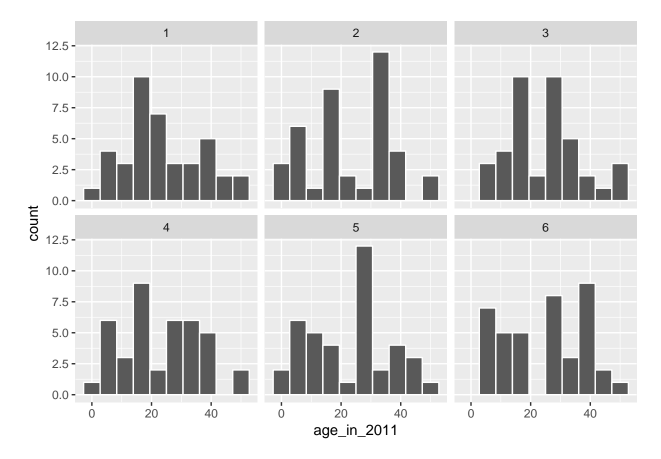
- piece of paper. Recall that these values were 6, 30, 34, 19, 6, etc.
- 2. Now, put the 40 small pieces of paper into a receptacle such as a baseball cap.
- 3. Shake up the pieces of paper.
- 4. Draw "at random" from the cap to select one piece of paper.
- 5. Write down the value on this piece of paper. Say that it is 28.
- 6. Now, place this piece of paper containing 28 back into the cap.
- 7. Draw "at random" again from the cap to select a piece of paper. Note that this is the sampling with replacement part since you may draw 28 again.
- 8. Repeat this process until you have drawn 40 pieces of paper and written down the values on these 40 pieces of paper. Completing this repetition produces ONE bootstrap sample.

If you look at the values in bootstrap_sample1, you can see how this process plays out. We originally drew 28, then we drew 11, then 7, and so on. Of course, we didn't actually use pieces of paper and a cap here. We just had the computer perform this process for us to produce bootstrap_sample1 using rep_sample_n() with replace = TRUE set.

The process of *sampling with replacement* is how we can use the original sample to take a guess as to what other values in the population may be. Sometimes in these bootstrap samples, we will select lots of larger values from the original sample, sometimes we will select lots of smaller values, and most frequently we will select values that are near the center of the sample. Let's explore what the distribution of values of age_in_2011 for six different bootstrap samples looks like to further understand this variability.

```
six_bootstrap_samples <- pennies_sample %>%
  rep_sample_n(size = 40, replace = TRUE, reps = 6)

ggplot(six_bootstrap_samples, aes(x = age_in_2011)) +
  geom_histogram(bins = 10, color = "white") +
  facet_wrap(~ replicate)
```



We can also look at the six different means using dplyr syntax:

```
six_bootstrap_samples %>%
group_by(replicate) %>%
summarize(stat = mean(age_in_2011))
```

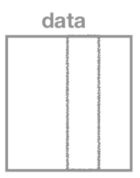
```
# A tibble: 6 x 2
  replicate stat
      <int> <dbl>
1
           1
              24.0
2
           2
              23.1
3
           3
              24.4
4
           4
              23.4
5
           5
              23.0
6
              24.6
```

Instead of doing this six times, we could do it 1000 times and then look at the distribution of stat across all 1000 of the replicates. This sets the stage for the infer R package (see documentation here or the "Cheat Sheet" on the DA Moodle page) that helps users perform statistical inference such as confidence intervals and hypothesis tests using verbs similar to what you've seen with dplyr. In the next section we'll walk through setting up each of the infer verbs for confidence intervals using this pennies_sample example, while also explaining the purpose of the verbs in a general framework.

4 Infer Package for Statistical Inference

The infer package makes great use of the tidyverse "pipe" %>% to create a pipeline for statistical inference. The goal of the package is to provide a way for its users to explain the computational process of confidence intervals and hypothesis tests using the code as a guide. The verbs build in order here, so you'll want to start with specify() and then continue through the others as needed.

4.1 Specify Variables



specify()

The specify() function is used primarily to choose which variables will be the focus of the statistical inference. In addition, a setting of which variable will act as the explanatory and which acts as the response variable is done here. For proportion problems (i.e. Scenarios 1 & 3 in Table 1) we also specify which of the different levels we are calculating the proportion of (e.g. "females", "approve of Obama's job performance", etc.). To begin to create a confidence interval for the population mean age of US pennies in 2011, we start by using specify() to choose which variable in our pennies_sample data we'd like to work with. This can be done in one of two ways:

1. Using the response argument:

pennies sample %>%

```
specify(response = age_in_2011)
Response: age_in_2011 (integer)
# A tibble: 40 \times 1
   age_in_2011
          <int>
 1
               6
 2
              30
 3
              34
 4
              19
 5
               6
 6
               5
 7
              11
 8
              19
 9
              23
10
              15
```

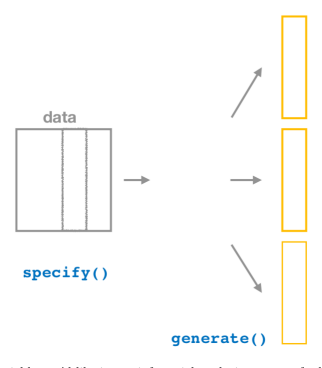
... with 30 more rows

2. Using formula notation:

```
pennies_sample %>%
  specify(formula = age_in_2011 ~ NULL)
Response: age_in_2011 (integer)
# A tibble: 40 \times 1
   age_in_2011
          <int>
 1
              6
 2
             30
             34
 3
 4
             19
 5
              6
 6
              5
 7
             11
 8
             19
9
             23
10
             15
# ... with 30 more rows
```

Note that the formula notation uses the common R methodology to include the response y variable on the left of the \sim and the explanatory x variable on the right of the "tilde." Recall that you used this notation frequently with the lm() function in Weeks 4 and 6 when fitting regression models. Either notation works just fine, but a preference is usually given here for the formula notation to further build on the ideas from earlier chapters.

4.2 Generate Replicates



After specify()ing the variables we'd like in our inferential analysis, we next feed that into the generate() verb. The generate() verb's main argument is reps, which is used to give how many different repetitions one would like to perform. Another argument here is type, which is automatically determined by the kinds

of variables passed into specify(). We can also be explicit and set this type to be type = "bootstrap". Make sure to check out ?generate to see the options here and use the ? operator to better understand other verbs as well. Let's generate() 1000 bootstrap samples:

```
thousand_bootstrap_samples <- pennies_sample %>%
  specify(response = age_in_2011) %>%
  generate(reps = 1000, type = "bootstrap")
```

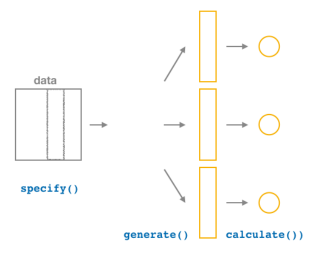
We can use the dplyr count() function to help us understand what the thousand_bootstrap_samples data frame looks like:

```
thousand_bootstrap_samples %>% count(replicate)
```

```
# A tibble: 1,000 x 2
            replicate [1,000]
# Groups:
   replicate
                  n
       <int> <int>
           1
 1
                 40
           2
 2
                 40
 3
           3
                 40
 4
            4
                 40
 5
           5
                 40
 6
           6
                 40
 7
           7
                 40
 8
           8
                 40
9
           9
                 40
10
          10
                 40
# ... with 990 more rows
# Equivalent to:
# thousand_bootstrap_samples %>%
    group_by(replicate) %>%
    summarise(n=n())
```

Notice that each replicate has 40 entries here. Now that we have 1000 different bootstrap samples, our next step is to calculate the bootstrap statistics for each sample.

4.3 Calculate Summary Statistics



After generate()ing many different samples, we next want to condense those samples down into a single statistic for each replicated sample. As seen in the diagram, the calculate() function is helpful here.

As we did at the beginning of this chapter, we now want to calculate the mean age_in_2011 for each bootstrap sample. To do so, we use the stat argument and set it to "mean" below. The stat argument has a variety of different options here and we will see further examples of this throughout the remaining chapters.

```
bootstrap_distribution <- pennies_sample %>%
  specify(response = age_in_2011) %>%
  generate(reps = 1000) %>%
  calculate(stat = "mean")
bootstrap_distribution
```

```
# A tibble: 1,000 x 2
   replicate stat
       <int> <dbl>
           1 22.8
 1
 2
           2 27.6
           3
              27.4
 3
 4
           4
             21.0
 5
           5
             29.1
 6
           6
             26.2
 7
           7
              25.3
 8
           8 22.9
 9
           9 22.7
10
          10 26.4
# ... with 990 more rows
```

Observed statistic / point estimate calculations Just as group_by() %>% summarize() produces a useful workflow in dplyr, we can also use specify() %>% calculate() to compute summary measures on our original sample data. It's often helpful both in confidence interval calculations and in hypothesis testing to identify what the corresponding statistic is in the original data. For our example on penny age, we computed above a value of x_bar using the summarize() verb in dplyr:

```
pennies_sample %>%
   summarize(stat = mean(age_in_2011))

# A tibble: 1 x 1
   stat
   <dbl>
1 25.1
```

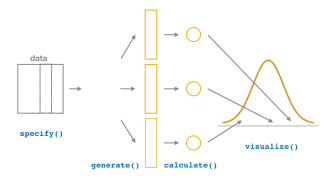
This can also be done by skipping the generate() step in the pipeline feeding specify() directly into calculate():

```
pennies_sample %>%
    specify(response = age_in_2011) %>%
    calculate(stat = "mean")

# A tibble: 1 x 1
    stat
    <dbl>
1 25.1
```

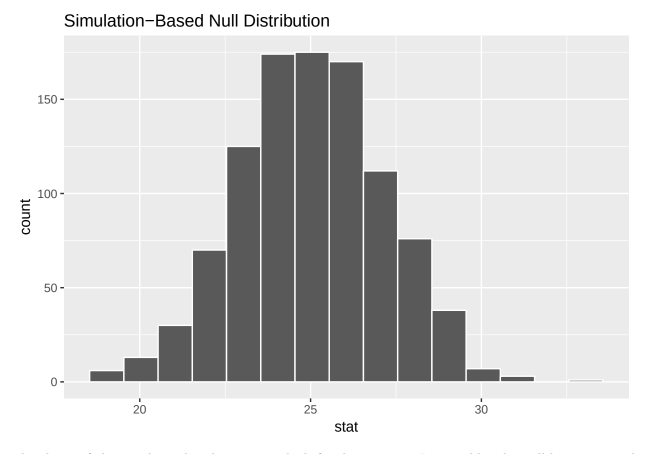
This shortcut will be particularly useful when the calculation of the observed statistic is tricky to do using dplyr alone. This is particularly the case when working with more than one variable.

4.4 Visualize the Results



The visualize() verb provides a simple way to view the bootstrap distribution as a histogram of the stat variable values. It has many other arguments that one can use as well including the shading of the histogram values corresponding to the confidence interval values.

bootstrap_distribution %>%
 visualize()



The shape of this resulting distribution may look familiar to you. It resembles the well-known normal (bell-shaped) curve. It is, in fact, an estimate of the **sampling variability** of the sample statistic. If you think back to *Statistical Inference* in Semester 1 you will remember that the *Central Limit Theorem* predicted that the sampling distribution would be a **normal distribution**, as seen in the bell-shaped distribution here.

The following diagram recaps the infer pipeline for creating a bootstrap distribution.

Confidence Interval in infer



5 Constructing Confidence Intervals

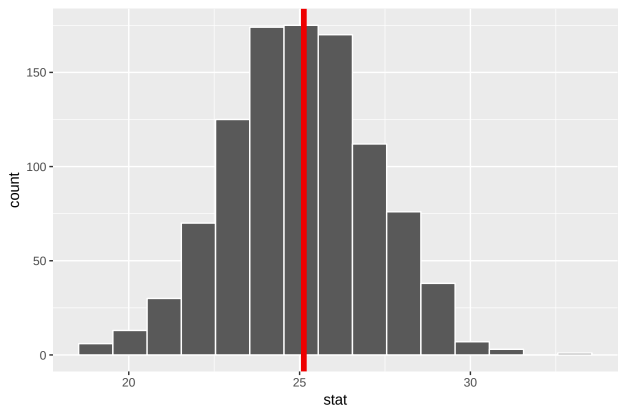
confidence interval: a range of plausible values for a population parameter. It depends on a specified confidence level with higher *confidence levels* corresponding to wider confidence intervals and lower confidence levels corresponding to narrower confidence intervals. Common confidence levels include 90%, 95%, and 99%.

Confidence intervals play an important role in the sciences and any field that uses data. You can think of a confidence interval as playing the role of a net when fishing. Instead of just trying to catch a fish with a single spear (estimating an unknown parameter by using a single point estimate/sample statistic), we can use a net to try to provide a range of possible locations for the fish (use a range of possible values based around our sample statistic to make a plausible guess as to the location of the parameter).

The bootstrapping process provides bootstrap statistics that have a bootstrap distribution with center at (or extremely close to) the mean of the original sample. This can be seen by giving the observed statistic obs_stat argument the value of the point estimate x_bar.

```
bootstrap_distribution %>%
  visualize(obs_stat = x_bar)
```





We can also compute the mean of the bootstrap distribution of means to see how it compares to x_bar:

5.1 The Percentile Method

One way to calculate a range of plausible values for the unknown mean age of coins in 2011 is to use the middle 95% of the bootstrap_distribution to determine our endpoints. Our endpoints are thus at the 2.5th and 97.5th percentiles. This can be done with infer using the get_ci() function. (You can also use the conf_int() or get_confidence_interval() functions here as they are aliases that work the exact same way.)

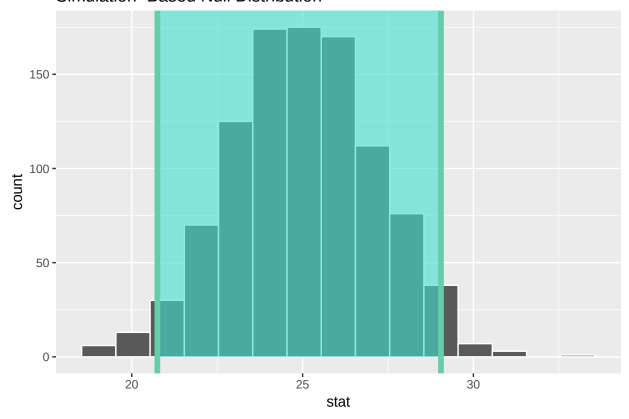
These options are the default values for level and type so we can also just do:

```
percentile_ci <- bootstrap_distribution %>%
  get_ci()
percentile_ci
# A tibble: 1 x 2
```

Using the percentile method, our range of plausible values for the mean age of US pennies in circulation in 2011 is 20.97 years to 29.25 years. We can use the visualize() function to view this using the endpoints and direction arguments, setting direction to "between" (between the values) and endpoints to be those stored with name percentile_ci.

```
bootstrap_distribution %>%
   visualize(endpoints = percentile_ci, direction = "between")
```





You can see that 95% of the data stored in the stat variable in bootstrap_distribution falls between the two endpoints with 2.5% to the left outside of the shading and 2.5% to the right outside of the shading. The cut-off points that provide our range are shown with the darker lines.

5.2 The Standard Error Method

If the bootstrap distribution is close to symmetric and bell-shaped, we can also use a shortcut formula for determining the lower and upper endpoints of the confidence interval. This is done by using the formula $\bar{x}\pm(multiplier???SE)$, where \bar{x} is our original sample mean and SE stands for **standard error** and

corresponds to the standard deviation of the bootstrap distribution.

Definition: The *stnadard error* is the standard deviation of the sampling distribution.

The variability of the sampling distribution may be approximated by the variability of the bootstrap distribution. Traditional theory-based methodologies for inference also have formulas for standard errors, assuming some conditions are met (you will have (seen some of these)[https://moodle.gla.ac.uk/pluginfile.php/1813455/mod_resource/content/1/Interval%20Estimates%20Summary.pdf] in Statistical Inference in Semester 1).

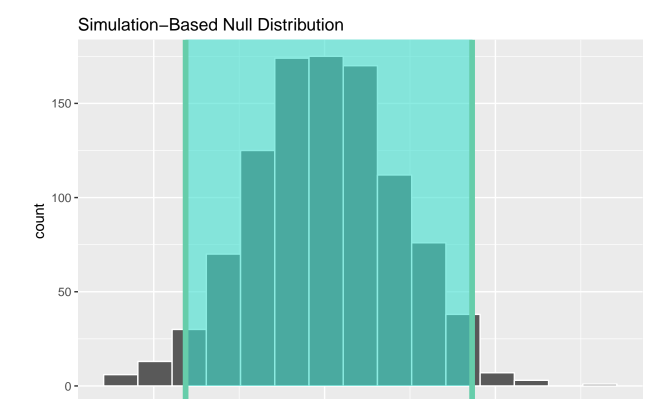
The value of *multiplier* here is the appropriate percentile of the standard normal distribution. This is automatically calculated when level is provided with level = 0.95 being the default. (95% of the values in a standard normal distribution fall within 1.96 standard deviations of the mean, so multiplier = 1.96 corresponds to level = 0.95, for example.) As mentioned, this formula assumes that the bootstrap distribution is symmetric and bell-shaped. This is often the case with bootstrap distributions, especially those in which the original distribution of the sample is not highly skewed.

This \bar{x} s(multiplier???SE) formula is implemented in the get_ci() function as shown with our pennies problem using the bootstrap distribution's variability as an approximation for the sampling distribution's variability. We'll see more on this approximation shortly.

Note that the center of the confidence interval (the point_estimate) must be provided for the standard error confidence interval.

```
standard_error_ci <- bootstrap_distribution %>%
  get_ci(type = "se", point_estimate = x_bar)
standard_error_ci

# A tibble: 1 x 2
  lower upper
  <dbl> <dbl>
1 20.9 29.3
bootstrap_distribution %>%
  visualize(endpoints = standard_error_ci, direction = "between")
```



We see that both methods produce nearly identical confidence intervals with the percentile method being [20.97, 29.25] and the standard error method being [20.97, 29.28].

stat

30

25

6 Interpreting the Confidence Interval

20

Recall that the confidence intervals we've produced are based on bootstrapping using the single sample pennies_sample. We have been claiming that this is a sample from all the pennies in circulation in 2011, but we can now reveal that it is actually a sample from a larger number of pennies stored as pennies in the moderndive package. The pennies data frame contains 800 rows of data and two columns pertaining to the same variables as pennies_sample. Its important to stress that this is very artificial, i.e. we would usually never have access to all the information about the larger group from which our sample is taken, but we have set up the data this way here to illustrate the properties of confidence intervals for the purpose of interpreting confidence intervals.

So let's assume that pennies is our population of interest (i.e. a population with N=800 units). We can therefor calculate the population mean age of pennies in 2011, denoted by the Greek letter μ , by calculating the mean of age_in_2011 for the pennies data frame.

```
pennies_mu <- pennies %>%
   summarize(overall_mean = mean(age_in_2011)) %>%
   pull() #Use this function to extract the single value from the data frame
pennies_mu
```

[1] 21.1525

As we saw at the end of the previous section, one range of plausible values for the population mean age of

pennies in 2011 (μ), is [20.97, 29.25]. Note that the value $\mu = 21.15$ (i.e.the mean of pennies calculated above) does fall in this confidence interval. So in this instance, the confidence interval based on pennies_sample was a good estimate of μ .

If we had a different sample of size 40 and constructed a confidence interval using the same method, would we be guaranteed that it contained the population parameter value μ as well? Let's try it out:

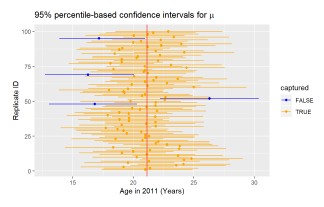
```
pennies_sample2 <- pennies %>%
sample_n(size = 40)
```

Note the use of the sample_n() function in the dplyr package here. This does the same thing as rep_sample_n(reps = 1) but omits the extra replicate column.

We next create an infer pipeline to generate a percentile-based 95% confidence interval for μ :

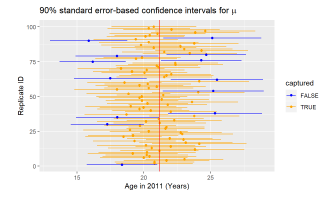
```
percentile_ci2 <- pennies_sample2 %>%
  specify(formula = age_in_2011 ~ NULL) %>%
  generate(reps = 1000) %>%
  calculate(stat = "mean") %>%
  get_ci()
percentile_ci2
```

This new confidence interval also contains the value of μ . Let's further investigate by repeating this process 100 times to get 100 different confidence intervals derived from 100 different samples of pennies. Each sample will have size of 40 just as the original sample. We will plot each of these confidence intervals as horizontal lines. We will also show a line corresponding to the known population value of 21.1525 years.



Of the 100 confidence intervals based on samples of size n=40, 96 of them captured the population mean $\mu=21.15$, whereas 4 of them did not include it. If we repeated this process of building confidence intervals more times with more samples, we'd expect 95% of them to contain the population mean. In other words, the procedure we have used to generate confidence intervals is "95% reliable" in that we can expect it to include the true population parameter 95% of the time if the process is repeated.

To further accentuate this point, let's perform a similar procedure using 90% confidence intervals instead. This time we will use the standard error method instead of the percentile method for computing the confidence intervals.



Repeating this process for more samples would result in us getting closer and closer to 90% of the confidence intervals including the true value. It is common to say while interpreting a confidence interval to be "95% confident" or "90% confident" that the specified confidence interval contains the true value. We will use this "confident" language throughout the rest of this chapter, but remember that it is a theoretical statement about what we would expect to happen if we to sample again and again from the same population (which we don't do in practice, of course).

7 Comparing Two Proportions

Let's start with an example. If you see someone else yawn, are you more likely to yawn? In an episode of the TV show Mythbusters, they tested the myth that yawning is contagious. Fifty adults who thought they were being considered for an appearance on the show were interviewed by a show recruiter ("confederate") who either yawned or did not. Participants then sat by themselves in a large van and were asked to wait. While in the van, the Mythbusters watched via hidden camera to see if the unaware participants yawned. The data frame containing the results is available at mythbusters_yawn in the moderndive package. Let's check it out.

mythbusters_yawn

```
# A tibble: 50 x 3
    subj group
                   yawn
   <int> <chr>
                   <chr>
       1 seed
 1
                   yes
 2
       2
          control
                  ves
 3
       3
          seed
                   no
 4
          seed
                   yes
 5
       5 seed
                   no
 6
       6
         control no
 7
          seed
 8
         control no
 9
         control no
10
      10 seed
                   no
      with 40 more rows
```

- The participant ID is stored in the subj variable with values of 1 to 50.
- The group variable is either "seed" for when a confederate was trying to influence the participant or "control" if a confederate did not interact with the participant.
- The yawn variable is either "yes" if the participant yawned or "no" if the participant did not yawn.

We can use the janitor package to get a glimpse into this data in a table format:

```
mythbusters_yawn %>%
  tabyl(group, yawn) %>%
  adorn_percentages() %>%
  adorn_pct_formatting() %>%
  adorn_ns() # To show original counts
```

```
group no yes
control 75.0% (12) 25.0% (4)
seed 70.6% (24) 29.4% (10)
```

We are interested in comparing the proportion of those that yawned after seeing a **seed** versus those that yawned with no seed interaction (i.e. **control**). We'd like to see if the difference between these two proportions is significantly larger than 0. If so, we'd have evidence to support the claim that yawning is contagious based on this study.

In looking over this problem, we can take note of some important details to include in our infer pipeline:

- We are calling a "success" having a yawn value of yes.
- Our response variable will always correspond to the variable used in the success so the response variable is yawn.
- The explanatory variable is the other variable of interest here: group.

7.1 Compute the Point Estimate

We are examining the relationship between yawning and whether or not the participant saw someone yawn (seed) or not (control).

```
mythbusters_yawn %>%
  specify(formula = yawn ~ group, success = "yes")
Response: yawn (factor)
Explanatory: group (factor)
# A tibble: 50 x 2
   yawn group
   <fct> <fct>
 1 yes
         seed
 2 yes
         control
 3 no
         seed
 4 yes
         seed
 5 no
         seed
 6 no
         control
7 yes
         seed
8 no
         control
9 no
         control
10 no
         seed
# ... with 40 more rows
```

We next want to calculate the statistic of interest for our sample. This corresponds to the difference in the proportion of successes.

```
obs_diff <- mythbusters_yawn %>%
   specify(formula = yawn ~ group, success = "yes") %>%
   calculate(stat = "diff in props", order = c("seed", "control")) # the order in which R should subtrac
obs_diff
```

A tibble: 1 x 1

```
stat
<dbl>
1 0.0441
```

This value represents the proportion of those that yawned after seeing a seed yawn (0.2941) minus the proportion of those that yawned with not seeing a seed (0.25).

7.2 Bootstrap Distribution

Our next step in building a confidence interval is to create a bootstrap distribution of statistics (differences in proportions of successes). We saw how it works with a single variable in computing bootstrap means in the pennies example but we haven't yet worked with bootstrapping involving multiple variables, i.e. comparing two groups. In the infer package, bootstrapping with multiple variables means that each **row** is potentially resampled. Let's investigate this by looking at the first few rows of mythbusters_yawn:

head(mythbusters_yawn)

```
# A tibble: 6 x 3
   subj group
                 yawn
  <int> <chr>
                 <chr>>
1
      1 seed
                 yes
2
      2 control yes
3
      3 seed
4
      4 seed
                 yes
5
      5 seed
                 nο
      6 control no
```

When we bootstrap this data, we are potentially pulling the subject's readings multiple times. Thus, we could see the entries of "seed" for group and "no" for yawn together in a new row in a bootstrap sample. This is further seen by exploring the sample_n() function in dplyr on this smaller 6 row data frame comprised of head(mythbusters_yawn). The sample_n() function can perform this bootstrapping procedure and is similar to the rep_sample_n() function in infer, except that it is not repeated but rather only performs one sample with or without replacement.

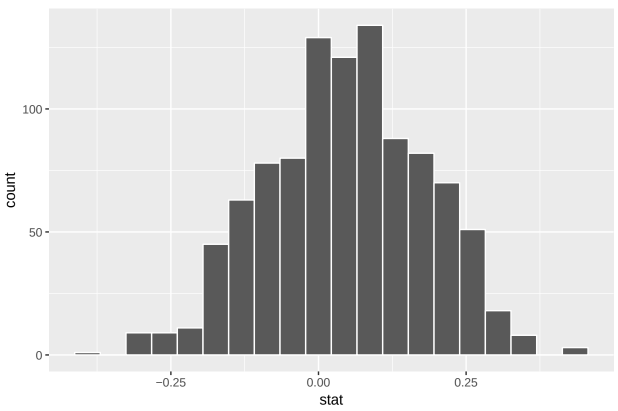
```
set.seed(2019)
head(mythbusters_yawn) %>%
  sample_n(size = 6, replace = TRUE)
```

```
# A tibble: 6 x 3
   subj group
                 yawn
  <int> <chr>
                 <chr>>
      5 seed
1
                 no
2
      5 seed
3
      2 control yes
4
      4 seed
                 yes
5
      1 seed
                 yes
      1 seed
```

We can see that in this bootstrap sample generated from the first six rows of mythbusters_yawn, we have some rows repeated. The same is true when we perform the generate() step in infer as done below.

```
bootstrap_distribution <- mythbusters_yawn %>%
  specify(formula = yawn ~ group, success = "yes") %>%
  generate(reps = 1000) %>%
  calculate(stat = "diff in props", order = c("seed", "control"))
bootstrap_distribution %>% visualize(bins = 20)
```





This distribution is roughly symmetric and bell-shaped but isn't quite there. Let's use the percentile-based method to compute a 95% confidence interval for the true difference in the proportion of those that yawn with and without a seed presented. The arguments are explicitly listed here but remember they are the defaults and simply get_ci() can be used.

```
bootstrap_distribution %>%
  get_ci(type = "percentile", level = 0.95)
# A tibble: 1 x 2
```

2.5% `97.5%` <dbl> <dbl>
1 -0.219 0.293

The confidence interval shown here is (-0.22, 0.29) and thus includes the value of 0. Therefore zero is a plausible value for the difference in the proportion of people that yawned with and without seeing someone yawn which is inconclusive evidence of a relationship between seeing someone yawn and yawning

Therefore, we are not sure which proportion is larger. Some of the bootstrap statistics showed the proportion without a seed to be higher and others showed the proportion with a seed to be higher. If the confidence interval was entirely above zero, we would be relatively sure (about "95% confident") that the seed group had a higher proportion of yawning than the control group.

Note that this all relates to the importance of denoting the order argument in the calculate() function. Since we specified "seed" and then "control" positive values for the statistic correspond to the "seed" proportion being higher, whereas negative values correspond to the "control" group being higher.

We, therefore, have evidence via this confidence interval suggesting that the conclusion from the Mythbusters show that "yawning is contagious" being "confirmed" is **not** statistically correct.

8 Further Tasks

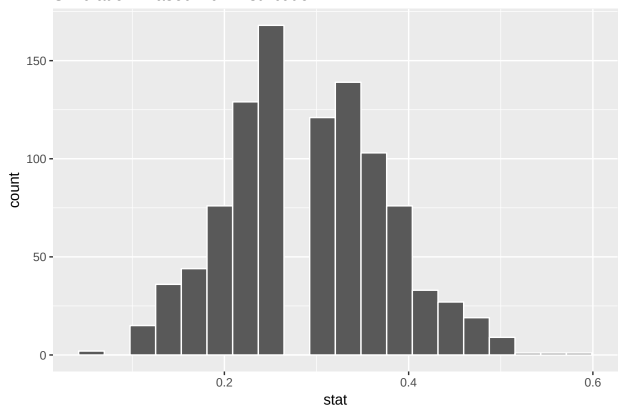
8.1 Confidence Intervals for Yawn Data

In the last section, we constructed a confidence interval for the difference in the proportion of people who yawned between the "seeded" group and the "control" group (Scenario 3).

By modifying the code in the last section in light of how we constructed a confidence interval for the age of pennies in the section on "Constructing confidence intervals" (Scenario 2), use mythbusters_yawn data to constuct a confidence interval for the proportion of people who yawn when they see someone else yawn (Scenario 1). Does this overlap with the confidence interval for the proportion of people who yawn when they did not see someone else yawn (Scenario 1 again!)? Are your findings here consistent with the findings in the last section?

```
bootstrap_distribution <- mythbusters_yawn %>%
  filter(group=="seed") %>%
  specify(formula = yawn ~ NULL, success = "yes") %>%
  generate(reps = 1000) %>%
  calculate(stat = "prop")
bootstrap_distribution %>% visualize(bins = 20)
```

Simulation—Based Null Distribution



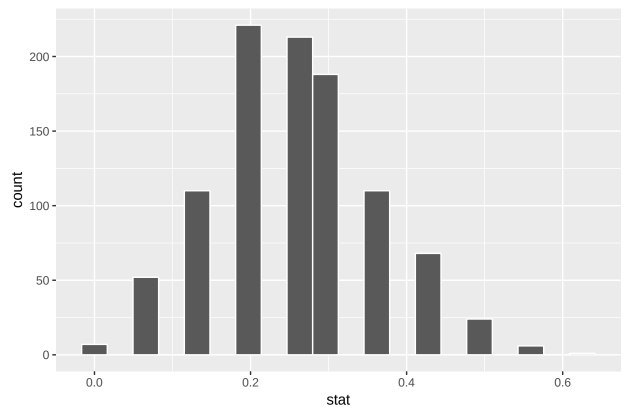
This distribution is roughly symmetric and bell-shaped but isn't quite there. Let's use the percentile-based method to compute a 95% confidence interval for the true proportion of those that yawn with a seed presented. The arguments are explicitly listed here but remember they are the defaults and simply get_ci() can be used.

```
bootstrap_distribution %>%
  get_ci(type = "percentile", level = 0.95)
```

The confidence interval shown here is (0.15, 0.44). The range of plausible values for the proportion of people that yawned with after seeing someone yawn is therefore between 0.15 and 0.44. We now repeat the exercise but for the group that didn't see someone yawn, i.e. group == "control"

```
bootstrap_distribution <- mythbusters_yawn %>%
  filter(group=="control") %>%
  specify(formula = yawn ~ NULL, success = "yes") %>%
  generate(reps = 1000) %>%
  calculate(stat = "prop")
bootstrap_distribution %>% visualize(bins = 20)
```

Simulation-Based Null Distribution



```
bootstrap_distribution %>%
  get_ci(type = "percentile", level = 0.95)
```

Setting type = "bootstrap" in generate(). The confidence interval shown here is (0.06, 0.5). The range of plausible values for the proportion of people that yawned without seeing someone yawn is therefore between 0.06 and 0.5.

Comparing the two CIs for the proportion of those that saw someone yawn (0.15, 0.44) and those that didn't

(0.06, 0.5), we note that these two CIs overlap, which is consistent with the findings from the CI for the difference in the two proptions (-0.23, 0.31) in the last section, i.e. since they **do overlap** its plausible that they could each take the *same value* and therefore its plausible that their **difference is zero**, which is exactly what the CI for the difference in proportions tells us.

8.2 Confidence Interval for Cats Data

Recall the data on 144 domestic male and female adult cats that we first saw in Week 4. Each cat had their heart weight in grams (Hwt) and body weight in kilograms (Bwt) measured, and interest lies in exploring difference between females and males. a. Construct a bootstrap confidence intervals for the avearge heart weight of female and male cats separately? Interpret your results. b. Construct a bootstrap confidence interval for the difference in the avearge heart weights of female and male cats. Interpret your result. c. Repeat a. and b. for the body weight of cats. Hint: You need to read in the cats data and remind yourself how it is organised, e.g.

```
cats <- read.csv("cats.csv")
glimpse(cats)</pre>
```

Solution 2.a.

a. Construct a bootstrap confidence intervals for the avearge heart weight of female and male cats separately? Interpret your results.

We start by focusing on female cats, i.e. Sex == "F":

```
cats <- read.csv("cats.csv")
#glimpse(cats)
bootstrap_distribution <- cats %>%
    filter(Sex == "F") %>%
    specify(response = Hwt) %>%
    generate(reps = 1000) %>%
    calculate(stat = "mean")

#bootstrap_distribution %>% visualize()

percentile_ci <- bootstrap_distribution %>%
    get_ci()
#percentile_ci
```

Using the percentile method, our range of plausible values for the mean heart weight of female adult cats is 8.82 grams to 9.58 grams.

Now repeat the analysis for male cats, i.e. Sex == "M":

```
bootstrap_distribution <- cats %>%
  filter(Sex == "M") %>%
  specify(response = Hwt) %>%
  generate(reps = 1000) %>%
  calculate(stat = "mean")
```

```
#bootstrap_distribution %>% visualize()
percentile_ci <- bootstrap_distribution %>%
   get_ci()
#percentile_ci
```

Using the percentile method, our range of plausible values for the mean heart weight of male adult cats is 10.83 grams to 11.79 grams.

Solution 2.b.

b. Construct a bootstrap confidence interval for the difference in the avearge heart weights of female and male cats. Interpret your result.

```
bootstrap_distribution <- cats %>%
  specify(Hwt~Sex) %>%
  generate(reps = 1000) %>%
  calculate(stat = "diff in means", order = c("F", "M"))
  percentile_ci <- bootstrap_distribution %>%
  get_ci()
```

Using the percentile method, our range of plausible values for the difference in the mean heart weight between female and male adult cats is -2.77 grams to -1.45 grams. That is to say that on average female adult cats' hearts weigh between 1.45 and 2.77 grams less than adult male cats' hearts. (Note: the fact that the individual CIs in part a. didn't overlap told us that zero wouldn't be in the interval for the difference in the population means)

Solution 2.c.

c. Repeat a. and b. for the body weight of cats.

RMarkdown makes it very easy to repeat the analysis in parts a. and b. on a different variable.

```
cats <- read.csv("cats.csv")
bootstrap_distribution <- cats %>%
  filter(Sex == "F") %>%
  specify(response = Bwt) %>%
  generate(reps = 1000) %>%
  calculate(stat = "mean")
percentile_ci <- bootstrap_distribution %>%
  get_ci()
```

Using the percentile method, our range of plausible values for the mean body weight of female adult cats is 2.28 kilograms to 2.43 kilograms.

Now repeat the analysis for male cats, i.e. Sex == "M":

```
bootstrap_distribution <- cats %>%
  filter(Sex == "M") %>%
  specify(response = Bwt) %>%
  generate(reps = 1000) %>%
  calculate(stat = "mean")
percentile_ci <- bootstrap_distribution %>%
  get_ci()
```

Using the percentile method, our range of plausible values for the mean body weight of male adult cats is 2.82 kilograms to 2.99 kilograms.

```
bootstrap_distribution <- cats %>%
   specify(Bwt~Sex) %>%
```

```
generate(reps = 1000) %>%
calculate(stat = "diff in means", order = c("F", "M"))
percentile_ci <- bootstrap_distribution %>%
get_ci()
```

Using the percentile method, our range of plausible values for the difference in the mean body weight between female and male adult cats is -0.67 kilograms to -0.41 kilograms. That is to say that on average female adult cats' bodies weigh between 0.41 and 0.67 kilograms less than adult male cats' bodies. (Note: the fact that the individual CIs in part a. didn't overlap told us that zero wouldn't be in the interval for the difference in the population means)