# Assumptions in hypothesis testing

HYPOTHESIS TESTING IN R



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#### Randomness

#### **Assumption**

The samples are random subsets of larger populations.

#### Consequence

Sample is not representative of population.

#### How to check this

- Understand how your data was collected.
- Speak to the data collector/domain expert.



<sup>&</sup>lt;sup>1</sup> Sampling techniques are discussed in "Sampling in R".



#### Independence of observations

#### **Assumption**

Each observation (row) in the dataset is independent.

#### Consequence

Increased chance of false negative/positive error.

#### How to check this

Understand how your data was collected.

#### Large sample size

#### **Assumption**

The sample is big enough to mitigate uncertainty, and so that the Central Limit Theorem applies.

#### Consequence

- Really wide confidence intervals.
- Increased chance of false negative/positive error.

#### How to check this

It depends on the test.

# Large sample size: t-test

#### One sample

• At least 30<sup>1</sup> observations in the sample.

$$n \ge 30$$

n: sample size

#### Paired samples

• At least 30 pairs of observations across the samples.

Number of rows in your data  $\geq 30$ 

#### Two samples

• At least 30 observations in each sample.

$$n_1 \ge 30, n_2 \ge 30$$

 $n_i$ : sample size for group i

#### **ANOVA**

 At least pairs of 30 observations in each sample.

 $n_i \geq 30$  for all values of i

<sup>&</sup>lt;sup>1</sup> Sometimes you can get away with less than 30; the important thing is that the null distribution appears normal.

# Large sample size: proportion tests

#### One sample

• Number of successes in sample is greater than or equal to 10.

$$n imes \hat{p} \geq 10$$

• Number of failures in sample is greater than or equal to 10.

$$n imes (1-\hat{p}) \geq 10$$

n: sample size

 $\hat{p}$ : proportion of successes in sample

#### Two samples

 Number of successes in each sample is greater than or equal to 10.

$$n_1 imes \hat{p}_1 \geq 10$$

$$n_2 imes \hat{p}_2 \geq 10$$

• Number of failures in each sample is greater than or equal to 10.

$$n_1 imes (1-\hat{p}_1)\geq 10$$

$$n_2 imes (1-\hat{p}_2) \geq 10$$

# Large sample size: chi-square tests

• The number of successes in each group in greater than or equal to 5.

$$n_i imes \hat{p}_i \geq 5$$
 for all values of  $i$ 

• The number of failures in each group in greater than or equal to 5.

$$n_i imes (1-\hat{p}_i) \geq 5$$
 for all values of  $i$ 

 $n_i$ : sample size for group i

 $\hat{p}_i$ : proportion of successes in sample group i

### Sanity check

If the bootstrap distribution doesn't look normal, assumptions likely aren't valid.



# Let's practice!

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# The "There is only one test" framework

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#### Imbalanced data

```
stack_overflow_imbalanced %>%
  count(hobbyist, age_cat, .drop = FALSE)
```

A sample is *imbalanced* if some groups are much bigger than others.

#### Hypotheses

 $H_0$ : The proportion of hobbyists under 30 is **the same as** the proportion of hobbyists at least 30.

 $H_A$ : The proportion of hobbyists under 30 is **different from** the proportion of hobbyists at least 30.

alpha <- 0.1

### Proceeding with a proportion test regardless

```
stack_overflow_imbalanced %>%

prop_test(
  hobbyist ~ age_cat,
  order = c("At least 30", "Under 30"),
  success = "Yes",
  alternative = "two.sided",
  correct = FALSE
)
```

# A grammar of graphics

Plot type	base-R	ggplot2
Scatter plot	plot(, type = "p")	<pre>ggplot() + geom_point()</pre>
Line plot	plot(, type = "l")	<pre>ggplot() + geom_line()</pre>
Histogram	hist()	<pre>ggplot() + geom_histogram()</pre>
Box plot	<pre>boxplot()</pre>	<pre>ggplot() + geom_boxplot()</pre>
Bar plot	<pre>barplot()</pre>	<pre>ggplot() + geom_bar()</pre>
Pie plot	pie()	<pre>ggplot() + geom_bar() + coord_polar()</pre>

#### A grammar of hypothesis tests

- Allen Downey's There is only one test framework.
- Implemented in R in the infer package.
- generate() makes simulated data.
  - Computationally expensive.
  - Robust against small samples or imbalanced data.

```
null_distn <- dataset %>%
  specify() %>%
  hypothesize() %>%
  generate() %>%
  calculate()
```

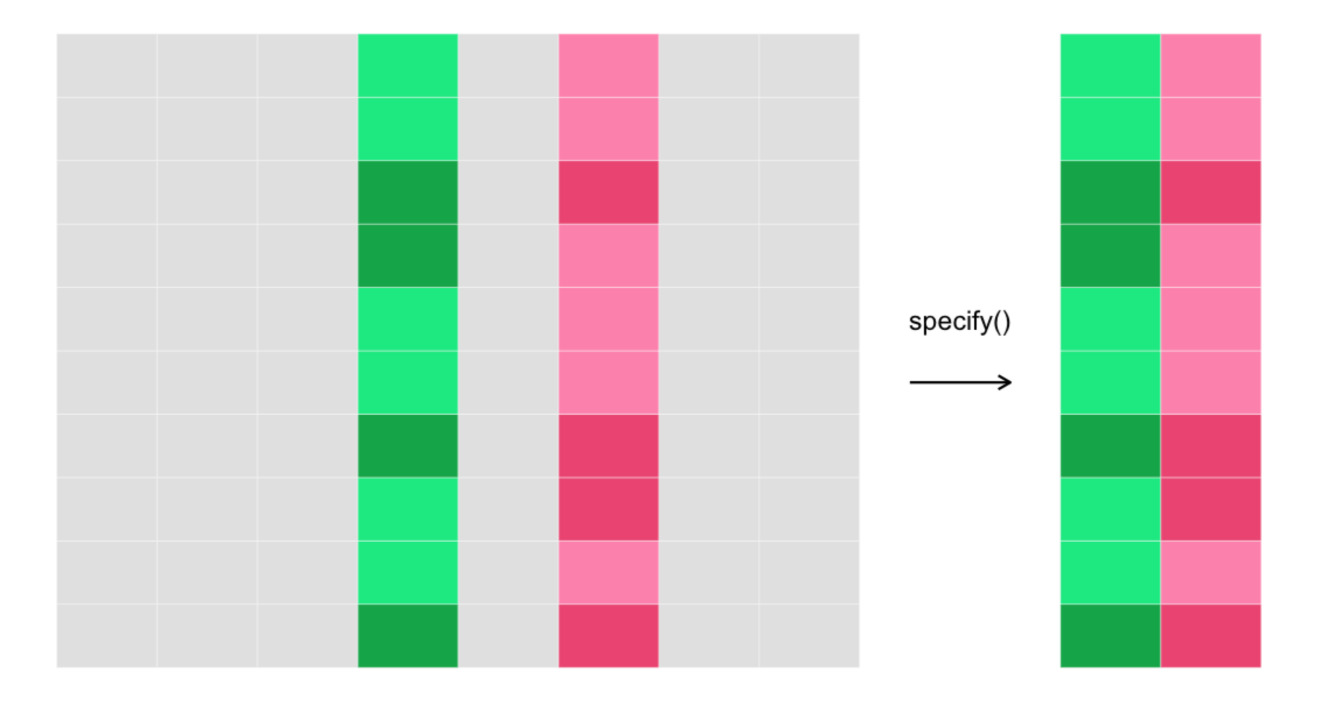
```
obs_stat <- dataset %>%
  specify() %>%
  calculate()
```

get\_p\_value(null\_distn, obs\_stat)

<sup>&</sup>lt;sup>1</sup> Allen Downey teaches "Exploratory Data Analysis in Python".



# Specifying the variables of interest



# specify()

```
specify() selects the variable(s) you want to test.
```

- For 2 sample tests, use
   response ~ explanatory.
- For 1 sample tests use response ~ NULL.

```
stack_overflow_imbalanced %>%
  specify(hobbyist ~ age_cat, success = "Yes")
```

```
Response: hobbyist (factor)
Explanatory: age_cat (factor)
# A tibble: 1,231 x 2
  hobbyist age_cat
  <fct>
          <fct>
1 Yes
          At least 30
          At least 30
2 Yes
          At least 30
3 Yes
4 Yes
          Under 30
          At least 30
5 Yes
6 Yes
          At least 30
          Under 30
7 No
# ... with 1,224 more rows
```

# hypothesize()

hypothesize() declares the type of null hypothesis.

- For 2 sample tests, use "independence" or "point".
- For 1 sample tests, use "point".

```
stack_overflow_imbalanced %>%
  specify(hobbyist ~ age_cat, success = "Yes") %>%
  hypothesize(null = "independence")
```

```
Response: hobbyist (factor)
Explanatory: age_cat (factor)
Null Hypothesis: independence
# A tibble: 1,231 x 2
 hobbyist age_cat
 <fct>
          <fct>
          At least 30
1 Yes
          At least 30
2 Yes
3 Yes
          At least 30
4 Yes
          Under 30
          At least 30
5 Yes
6 Yes
          At least 30
7 No
          Under 30
# ... with 1,224 more rows
```

# Let's practice!

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# Continuing the infer pipeline

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#### Recap: hypotheses and dataset

 $H_0$ : The proportion of hobbyists under 30 is the same as the prop'n of hobbyists at least 30.

 $H_A$ : The proportion of hobbyists under 30 is different from the prop'n of hobbyists at least 30.

```
alpha <- 0.1
```

```
stack_overflow_imbalanced %>%
count(hobbyist, age_cat, .drop = FALSE)
```

#### Recap: workflow

get\_p\_value(null\_distn, observed\_stat)

```
null_distn <- dataset %>%
   specify() %>%
   hypothesize() %>%
   generate() %>%
   calculate()

observed_stat <- dataset %>%
   specify() %>%
   calculate()
```

```
stack_overflow_imbalanced %>%
  specify(hobbyist ~ age_cat, success = "Yes") %>%
  hypothesize(null = "independence")
```

```
Response: hobbyist (factor)
Explanatory: age_cat (factor)
Null Hypothesis: independence
# A tibble: 1,231 x 2
 hobbyist age_cat
          <fct>
 <fct>
1 Yes
         At least 30
2 Yes
        At least 30
3 Yes
        At least 30
4 Yes
         Under 30
5 Yes
        At least 30
6 Yes
         At least 30
7 No
          Under 30
# ... with 1,224 more rows
```

# **Motivating generate()**

 $H_0$ : The proportion of hobbyists under 30 is the same as the prop'n of hobbyists at least 30.

If  $H_0$  is true, then

- In each row, the hobbyist value could have appeared with either age category with equal probability.
- To simulate this, we can permute (shuffle) the hobbyist values while keeping the age categories fixed.

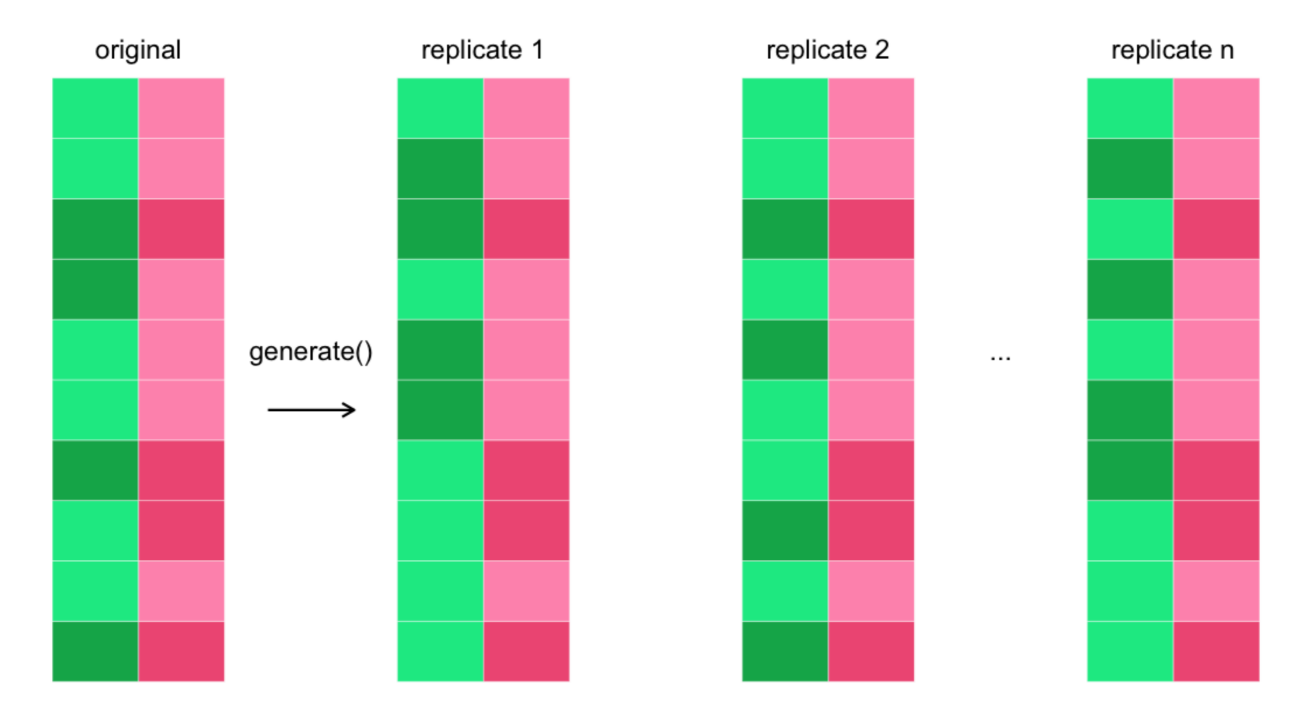
```
stack_overflow_imbalanced
```

```
bind_cols(
   stack_overflow_imbalanced %>%
     select(hobbyist) %>%
     slice_sample(prop = 1),
   stack_overflow_imbalanced %>%
     select(age_cat)
)
```

```
# A tibble: 1,231 x 2
 hobbyist age_cat
 <fct>
          <fct>
1 Yes
       At least 30
2 Yes
      At least 30
       At least 30
3 Yes
4 Yes
       Under 30
5 Yes
     At least 30
     At least 30
6 Yes
7 No
         Under 30
# ... with 1,224 more rows
```

```
# A tibble: 1,231 x 2
hobbyist age_cat
  <fct> <fct>
1 Yes         At least 30
2 Yes         At least 30
3 No         At least 30
4 No         Under 30
5 Yes         At least 30
6 Yes         At least 30
7 Yes         Under 30
# ... with 1,224 more rows
```

# Generating many replicates





# generate()

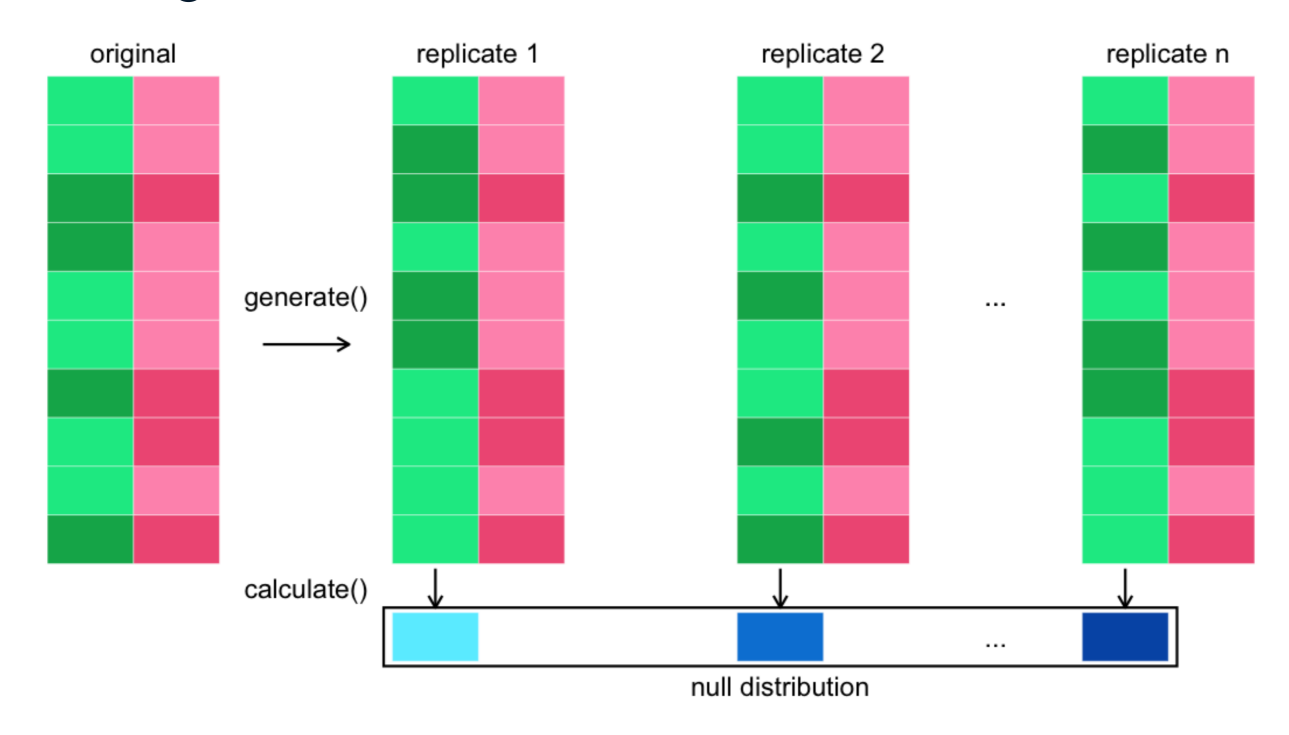
generate() generates simulated data reflecting the null hypothesis.

- For "independence" null hypotheses, set type to "permute".
- For "point" null hypotheses, set type to "bootstrap" or "simulate".

```
stack_overflow_imbalanced %>%
  specify(hobbyist ~ age_cat, success = "Yes") %>%
  hypothesize(null = "independence") %>%
  generate(reps = 5000, type = "permute")
```

```
Response: hobbyist (factor)
Explanatory: age_cat (factor)
Null Hypothesis: independence
# A tibble: 6,155,000 x 3
# Groups: replicate [5,000]
 hobbyist age_cat replicate
 <fct>
         <fct>
                       <int>
         At least 30
1 Yes
2 Yes
         At least 30
3 Yes
         At least 30
4 Yes
         Under 30
5 Yes
         At least 30
6 Yes
         At least 30
         Under 30
7 Yes
# ... with 6,154,993 more rows
```

# Calculating the test statistic





# calculate()

calculate() calculates a distribution of test statistics known as the *null distribution*.

```
null_distn <- stack_overflow_imbalanced %>%
 specify(
    hobbyist ~ age_cat,
    success = "Yes"
 ) %>%
 hypothesize(null = "independence") %>%
 generate(reps = 5000, type = "permute") %>%
 calculate(
    stat = "diff in props",
    order = c("At least 30", "Under 30")
```

```
# A tibble: 5,000 x 2
  replicate
             stat
      <int> <dbl>
         1 0.0896
1
2
         2 0.0896
3
         3 - 0.180
         4 0.157
4
5
         5 0.0896
6
         6 -0.113
         7 0.0221
# ... with 4,993 more rows
```

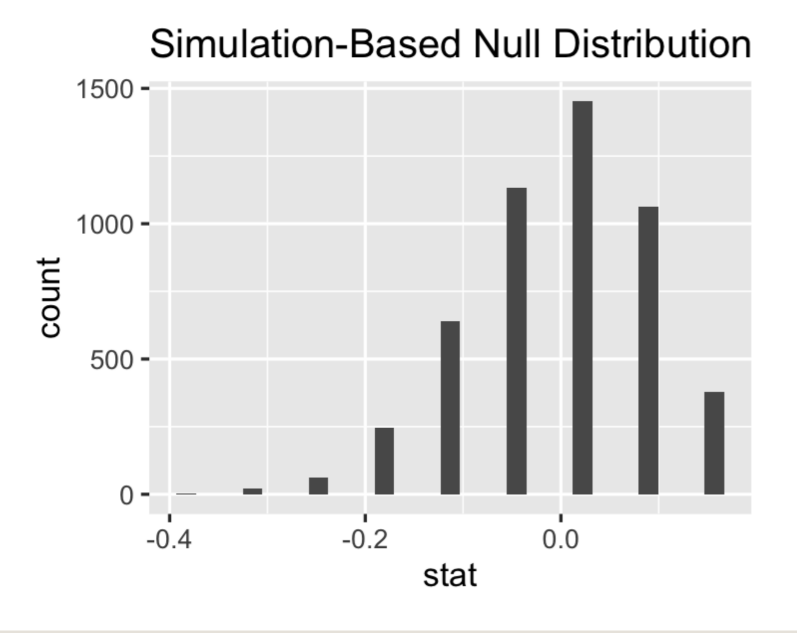
<sup>&</sup>lt;sup>1</sup> The ?calculate help page lists all possible test statistics.



#### Visualizing the null distribution

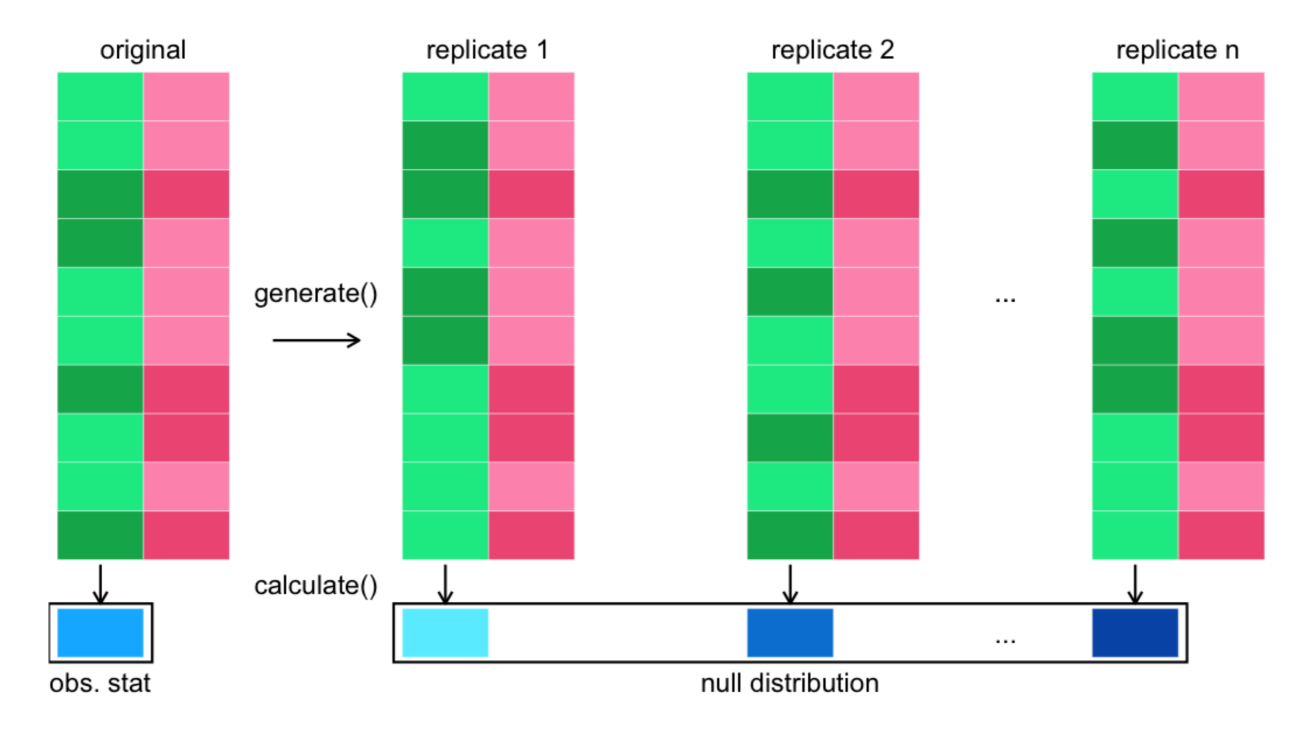
visualize(null\_distn)

null\_distn %>% count(stat)



```
# A tibble: 9 x 2
     stat
              n
    <dbl> <int>
1 - 0.383
2 - 0.315
3 - 0.248
             63
4 -0.180
            246
5 -0.113
          641
6 -0.0454
           1132
  0.0221
           1453
  0.0896
           1063
  0.157
            378
```

# Calculating the test statistic on the original dataset





# Observed statistic: specify() %>% calculate()

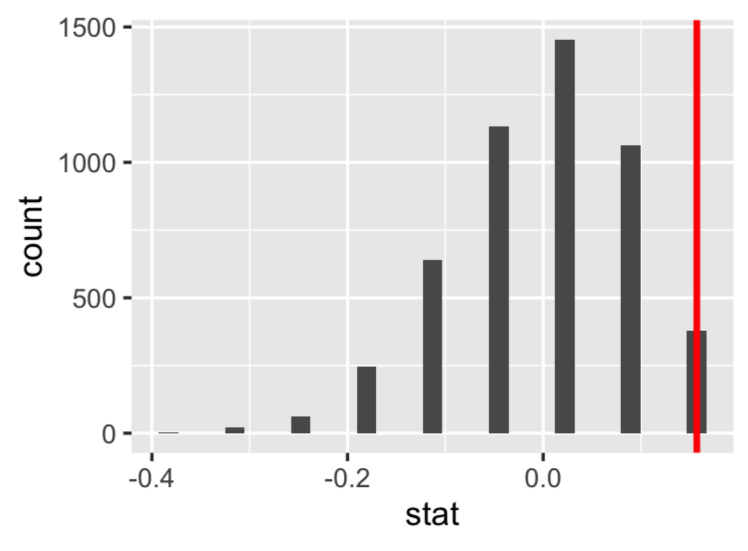
```
obs_stat <- stack_overflow_imbalanced %>%
  specify(hobbyist ~ age_cat, success = "Yes") %>%
  # hypothesize(null = "independence") %>%
  # generate(reps = 5000, type = "permute") %>%
  calculate(
    stat = "diff in props",
    order = c("At least 30", "Under 30")
)
```

```
# A tibble: 1 x 1
    stat
    <dbl>
1 0.157
```

#### Visualizing the null distribution vs the observed stat

```
visualize(null_distn) +
  geom_vline(
    aes(xintercept = stat),
    data = observed_stat,
    color = "red"
)
```

#### Simulation-Based Null Distribution



#### Get the p-value

```
get_p_value(
  null_distn, obs_stat,
  direction = "two sided"  # Not alternative = "two.sided"
)
```

```
# A tibble: 1 x 1
    p_value
    <dbl>
1 0.151
```

# Let's practice!

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# Non-parametric ANOVA and unpaired t-tests

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#### Non-parametric tests

A *non-parametric test* is a hypothesis test that doesn't assume a probability distribution for the test statistic.

There are two types of non-parametric hypothesis test:

- 1. Simulation-based.
- 2. Rank-based.

# t\_test()

```
H_0: \mu_{child} - \mu_{adult} = 0 H_A: \mu_{child} - \mu_{adult} > 0
```

```
library(infer)
stack_overflow %>%

t_test(
    converted_comp ~ age_first_code_cut,
    order = c("child", "adult"),
    alternative = "greater"
)
```

## Calculating the null distribution

## Simulation-based pipeline

```
null_distn <- stack_overflow %>%
  specify(converted_comp ~ age_first_code_cut) %>%
  hypothesize(null = "independence") %>%
  generate(reps = 5000, type = "permute") %>%
  calculate(
    stat = "diff in means",
    order = c("child", "adult")
)
```

### t-test, for comparison

```
library(infer)
stack_overflow %>%
  t_test(
    converted_comp ~ age_first_code_cut,
    order = c("child", "adult"),
    alternative = "greater"
)
```

## Calculating the observed statistic

### Simulation-based pipeline

```
obs_stat <- stack_overflow %>%
  specify(converted_comp ~ age_first_code_cut) %>%
  calculate(
    stat = "diff in means",
    order = c("child", "adult")
)
```

### t-test, for comparison

```
library(infer)
stack_overflow %>%
  t_test(
    converted_comp ~ age_first_code_cut,
    order = c("child", "adult"),
    alternative = "greater"
)
```

## Get the p-value

## Simulation-based pipeline

```
get_p_value(
  null_distn, obs_stat,
  direction = "greater"
)
```

### t-test, for comparison

```
library(infer)
stack_overflow %>%
  t_test(
    converted_comp ~ age_first_code_cut,
    order = c("child", "adult"),
    alternative = "greater"
)
```

## Ranks of vectors

```
x \leftarrow c(1, 15, 3, 10, 6)
```

rank(x)

#### 1 5 2 4 3

A Wilcoxon-Mann-Whitney test (a.k.a. Wilcoxon rank sum test) is (very roughly) a t-test on the ranks of the numeric input.

# Wilcoxon-Mann-Whitney test

```
wilcox.test(
  converted_comp ~ age_first_code_cut,
  data = stack_overflow,
  alternative = "greater",
  correct = FALSE
)
```

```
Wilcoxon rank sum test

data: converted_comp by age_first_code_cut

W = 967298, p-value <2e-16

alternative hypothesis: true location shift is greater than 0
```

<sup>&</sup>lt;sup>1</sup> Also known as the "Wilcoxon rank-sum test" and the "Mann-Whitney U test".



## Kruskal-Wallis test

Kruskal-Wallis test is to Wilcoxon-Mann-Whitney test as ANOVA is to t-test.

```
kruskal.test(
  converted_comp ~ job_sat,
  data = stack_overflow
)
```

```
Kruskal-Wallis rank sum test

data: converted_comp by job_sat
Kruskal-Wallis chi-square = 81, df = 4, p-value <2e-16</pre>
```

# Let's practice!

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# Congratulations!

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# You learned things

### Chapter 1

- Workflow for testing proportions vs. a hypothesized value.
- False negative/false positive errors.

### Chapter 2

- Testing differences in sample means between two groups using t-tests.
- Extending this to more than two groups using ANOVA and pairwise t-tests.

#### Chapter 3

- Testing differences in sample proportions between two groups using proportion tests.
- Using chi-square independence/goodness of fit tests.

### Chapter 4

- Reviewing assumptions of parametric hypothesis tests.
- Examined nonparametric alternatives when assumptions aren't valid

## More courses

#### Inference

Statistical Inference with R skill track

## **Bayesian statistics**

Fundamentals of Bayesian Data Analysis in R

## **Applications**

A/B Testing in R

# Let's practice!

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