The sign test

Packages

```
library(tidyverse)
library(smmr)
```

smmr is new. See later how to install it.

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Duality between confidence intervals and hypothesis tests

- Tests and CIs really do the same thing, if you look at them the right way. They are both telling you something about a parameter, and they use same things about data.
- To illustrate, some data (two groups):

```
my_url <- "http://ritsokiguess.site/datafiles/duality.txt"
twogroups <- read_delim(my_url," ")</pre>
```

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The data

twogroups

```
# A tibble: 15 x 2
       y group
   <dbl> <dbl>
 1
      10
      11
      11
      13
 5
      13
      14
7
      14
8
      15
9
      16
10
      13
11
      13
12
      14
              2
13
      17
              2
14
      18
15
      19
              2
```

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95% CI (default)

for difference in means, group 1 minus group 2:

13.00000 15.66667

```
t.test(y ~ group, data = twogroups)
```

Welch Two Sample t-test

```
data: y by group
t = -2.0937, df = 8.7104, p-value = 0.0668
alternative hypothesis: true difference in means between group
95 percent confidence interval:
   -5.5625675   0.2292342
sample estimates:
mean in group 1 mean in group 2
```

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90% CI

```
t.test(y ~ group, data = twogroups, conf.level = 0.90)
```

Welch Two Sample t-test

mean in group 1 mean in group 2

13.00000 15.66667

```
data: y by group
t = -2.0937, df = 8.7104, p-value = 0.0668
alternative hypothesis: true difference in means between group
90 percent confidence interval:
   -5.010308 -0.323025
sample estimates:
```

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Hypothesis test

Null is that difference in means is zero:

```
t.test(y ~ group, mu=0, data = twogroups)
```

Welch Two Sample t-test

13.00000 15.66667

```
data: y by group
t = -2.0937, df = 8.7104, p-value = 0.0668
alternative hypothesis: true difference in means between group
95 percent confidence interval:
   -5.5625675   0.2292342
sample estimates:
mean in group 1 mean in group 2
```

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Comparing results

Recall null here is $H_0: \mu_1-\mu_2=0$. P-value 0.0668.

- 95% CI from -5.6 to 0.2, contains 0.
- 90% CI from -5.0 to -0.3, does not contain 0.
- At $\alpha=0.05$, would not reject H_0 since P-value >0.05.
- At $\alpha = 0.10$, would reject H_0 since P-value < 0.10.

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Test and CI

Not just coincidence. Let $C=100(1-\alpha)$, so C% gives corresponding CI to level- α test. Then following always true. (Symbol \iff means "if and only if".)

Test decision		Confidence interval
Reject H_0 at level α	\Leftrightarrow	C% CI does not contain
		H_0 value
Do not reject ${\cal H}_0$ at level	\iff	$C\%$ CI contains H_0 value
α		

Idea: "Plausible" parameter value inside CI, not rejected; "Implausible" parameter value outside CI, rejected.

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The value of this

- If you have a test procedure but no corresponding CI:
- you make a CI by including all the parameter values that would not be rejected by your test.
- Use:
 - $\alpha = 0.01$ for a 99% CI.
 - $\sim \alpha = 0.05$ for a 95% CI.
 - $\sim \alpha = 0.10$ for a 90% CI, and so on.

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Testing for non-normal data

- The IRS ("Internal Revenue Service") is the US authority that deals with taxes (like Revenue Canada).
- One of their forms is supposed to take no more than 160 minutes to complete. A citizen's organization claims that it takes people longer than that on average.
- Sample of 30 people; time to complete form recorded.
- \bullet Read in data, and do t-test of $H_0: \mu = 160$ vs. $H_a: \mu > 160.$
- For reading in, there is only one column, so can pretend it is delimited by anything.

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Read in data

```
my_url <- "http://ritsokiguess.site/datafiles/irs.txt"
irs <- read_csv(my_url)
irs</pre>
```

```
A tibble: 30 x 1
   Time
   <dbl>
     91
2
    64
3
    243
    167
5
    123
6
    65
7
    71
8
    204
9
    110
10
    178
# i 20 more rows
```

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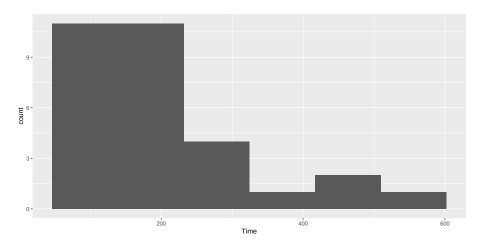
Test whether mean is 160 or greater

One Sample t-test

Reject null; mean (for all people to complete form) greater than 160.

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But, look at a graph



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Comments

- Skewed to right.
- Should look at median, not mean.

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- But how to test whether the median is greater than 160?
- Idea: if the median really is 160 (H_0 true), the sampled values from the population are equally likely to be above or below 160.
- If the population median is greater than 160, there will be a lot of sample values greater than 160, not so many less. Idea: test statistic is number of sample values greater than hypothesized median.

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Getting a P-value for sign test 1/3

- How to decide whether "unusually many" sample values are greater than 160? Need a sampling distribution.
- ullet If H_0 true, pop. median is 160, then each sample value independently equally likely to be above or below 160.
- So number of observed values above 160 has binomial distribution with n=30 (number of data values) and p=0.5 (160 is hypothesized to be median).

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Getting P-value for sign test 2/3

Count values above/below 160:

```
irs %>% count(Time > 160)
```

• 17 above, 13 below. How unusual is that? Need a binomial table.

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Getting P-value for sign test 3/3

 \bullet R function dbinom gives the probability of eg. exactly 17 successes in a binomial with n=30 and p=0.5 :

```
dbinom(17, 30, 0.5)
```

[1] 0.1115351

• but we want probability of 17 *or more*, so get all of those, find probability of each, and add them up:

```
tibble(x=17:30) %>%
  mutate(prob=dbinom(x, 30, 0.5)) %>%
  summarize(total=sum(prob))
```

```
# A tibble: 1 x 1
  total
  <dbl>
1 0.292
```

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Aside

pbinom gives the cumulative probability (prob. of less than or equal than the first input):

```
pbinom(17, 30, 0.5) # prob of <= 17
```

[1] 0.8192027

and hence (note first input):

[1] 0.2923324

This last is $P(X \ge 17) = P(X > 16)$.

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Using my package smmr

- I wrote a package smmr to do the sign test (and some other things).
 Installation is a bit fiddly:
 - ▶ Install devtools (once) with

```
install.packages("devtools")
```

• then install smmr using devtools (once):

```
library(devtools)
install_github("nxskok/smmr")
```

• Then load it:

```
library(smmr)
```

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smmr for sign test

 smmr's function sign_test needs three inputs: a data frame, a column and a null median:

```
sign_test(irs, Time, 160)
$above_below
below above
   13 17
$p_values
 alternative p_value
       lower 0.8192027
       upper 0.2923324
   two-sided 0.5846647
```

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Comments (1/4)

- Testing whether population median *greater than* 160, so want *upper-tail* P-value 0.2923. Same as before.
- Also get table of values above and below; this too as we got.

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Comments (2/4)

P-values are:

Test	P-value
\overline{t}	0.0392
Sign	0.2923

• These are very different: we reject a mean of 160 (in favour of the mean being bigger), but clearly *fail* to reject a median of 160 in favour of a bigger one.

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Comments 3/4

• Why is that? Obtain mean and median:

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Comments (4/4)

- The mean is pulled a long way up by the right skew, and is a fair bit bigger than 160.
- The median is quite close to 160.
- We ought to be trusting the sign test and not the t-test here (median and not mean), and therefore there is no evidence that the "typical" time to complete the form is longer than 160 minutes.
- Having said that, there are clearly some people who take a lot longer than 160 minutes to complete the form, and the IRS could focus on simplifying its form for these people.
- In this example, looking at any kind of average is not really helpful; a better question might be "do an unacceptably large fraction of people take longer than (say) 300 minutes to complete the form?": that is, thinking about worst-case rather than average-case.

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Confidence interval for the median

- The sign test does not naturally come with a confidence interval for the median.
- So we use the "duality" between test and confidence interval to say: the (95%) confidence interval for the median contains exactly those values of the null median that would not be rejected by the two-sided sign test (at $\alpha=0.05$).

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For our data

- The procedure is to try some values for the null median and see which ones are inside and which outside our CI.
- smmr has pval_sign that gets just the 2-sided P-value:

```
pval_sign(160, irs, Time)
```

[1] 0.5846647

• Try a couple of null medians:

```
pval_sign(200, irs, Time)
```

[1] 0.3615946

```
pval_sign(300, irs, Time)
```

[1] 0.001430906

So 200 inside the 95% CI and 300 outside.

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Doing a whole bunch

• Choose our null medians first:

```
(d <- tibble(null_median=seq(100,300,20)))</pre>
```

```
# A tibble: 11 x 1
   null median
          <dbl>
            100
            120
 3
            140
            160
 5
            180
 6
            200
            220
            240
            260
10
            280
11
            300
```

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... and then

"for each null median, run the function pval_sign for that null median and get the P-value":

```
d %>% rowwise() %>%
 mutate(p_value = pval_sign(null_median, irs, Time))
 A tibble: 11 x 2
# Rowwise:
  null_median p_value
         <dbl> <dbl>
           100 0.000325
           120 0.0987
           140 0.200
           160 0.585
5
           180 0.856
6
           200 0.362
7
           220 0.0428
8
           240 0.0161
9
           260 0.00522
10
           280 0.00143
11
           300 0.00143
```

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Make it easier for ourselves

11

```
d %>% rowwise() %>%
 mutate(p_value = pval_sign(null_median, irs, Time)) %>%
 mutate(in_out = ifelse(p_value > 0.05, "inside", "outside"))
# A tibble: 11 \times 3
# Rowwise:
  null_median p_value in_out
        <dbl> <dbl> <chr>
          100 0.000325 outside
2
          120 0.0987 inside
3
          140 0.200 inside
          160 0.585 inside
5
          180 0.856 inside
6
          200 0.362 inside
7
          220 0.0428 outside
8
          240 0.0161 outside
          260 0.00522 outside
10
          280 0.00143 outside
```

300 0.00143 outside

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confidence interval for median?

- 95% CI to this accuracy from 120 to 200.
- Can get it more accurately by looking more closely in intervals from 100 to 120, and from 200 to 220.

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A more efficient way: bisection

• Know that top end of CI between 200 and 220:

```
lo <- 200
hi <- 220
```

• Try the value halfway between: is it inside or outside?

```
try <- (lo + hi) / 2
try</pre>
```

[1] 210

```
pval_sign(try,irs,Time)
```

[1] 0.09873715

• Inside, so upper end is between 210 and 220. Repeat (over):

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... bisection continued

```
lo <- try
try <- (lo + hi) / 2
try
```

[1] 215

```
pval_sign(try, irs, Time)
```

[1] 0.06142835

- 215 is inside too, so upper end between 215 and 220.
- Continue until have as accurate a result as you want.

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Bisection automatically

 A loop, but not a for since we don't know how many times we're going around. Keep going while a condition is true:

```
10 = 200
hi = 220
while (hi - lo > 1) {
  try = (hi + lo) / 2
  ptry = pval_sign(try, irs, Time)
  print(c(try, ptry))
  if (ptry \ll 0.05)
    hi = try
  else
    lo = try
```

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The output from this loop

```
      [1]
      210.00000000
      0.09873715

      [1]
      215.00000000
      0.06142835

      [1]
      217.50000000
      0.04277395

      [1]
      216.25000000
      0.04277395

      [1]
      215.62500000
      0.04277395
```

• 215 inside, 215.625 outside. Upper end of interval to this accuracy is 215.

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Using smmr

• smmr has function ci_median that does this (by default 95% CI):

```
ci_median(irs, Time)
```

[1] 119.0065 214.9955

- Uses a more accurate bisection than we did.
- Or get, say, 90% CI for median:

```
ci_median(irs, Time, conf.level=0.90)
```

[1] 123.0031 208.9960

• 90% CI is shorter, as it should be.

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Bootstrap

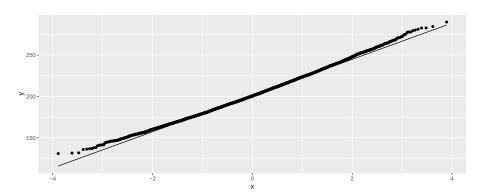
- but, was the sample size (30) big enough to overcome the skewness?
- Bootstrap, again:

```
tibble(sim = 1:1000) %>%
  rowwise() %>%
  mutate(my_sample = list(sample(irs$Time, replace = TRUE))) %
  mutate(my_mean = mean(my_sample)) %>%
  ggplot(aes(x=my_mean)) + geom_histogram(bins=10) -> g
```

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With normal quantile plot

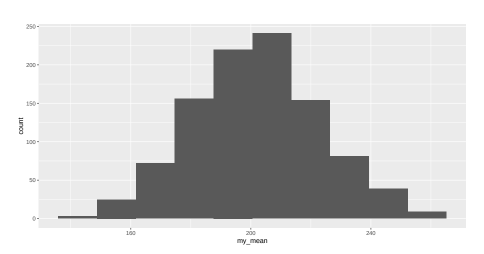
```
tibble(sim = 1:10000) %>%
  rowwise() %>%
  mutate(my_sample = list(sample(irs$Time, replace = TRUE))) %
  mutate(my_mean = mean(my_sample)) %>%
  ggplot(aes(sample = my_mean)) + stat_qq() + stat_qq_line()
```



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The sampling distribution

g



Comments

- A little skewed to right, but not nearly as much as I was expecting.
- The t-test for the mean might actually be OK for these data, if the mean is what you want.
- In actual data, mean and median very different; we chose to make inference about the median.
- Thus for us it was right to use the sign test.

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