## 1.4 Bootstrap CIs

We will look at five different ways to create confidence intervals using the boostrap and discuss which to use when.

- 1. Percentile Bootstrap CI

- 2. Basic Bootstrap CI

  3. Standard Normal Bootstrap CI

  4. Bootstrap t (studentized)

  5. Accelerated Bias-Corrected (BCa) adjusted for skewness.

- 1) When you say "we und bootstrapping I estimate CI" you need to say which one.
- Lhatever you are boutstrapping reeds to be independent.

  3 Boutstrapping is an attempt to simulate replication.

  (think about interpretasion of a CI).

### 1.4.1 Percentile Bootstrap CI

Let  $\hat{\theta}^{(1)}, \ldots, \hat{\theta}^{(B)}$  be bootstrap replicates and let  $\hat{\theta}_{\alpha/2}$  be the  $\alpha/2$  quantile of  $\hat{\theta}^{(1)}, \ldots, \hat{\theta}^{(B)}$ . (bootstryp den of ô). Then, the  $100(1-\alpha)\%$  Percentile Bootstrap CI for  $\theta$  is

In R, if bootstrap.reps =  $c(\hat{\theta}^{(1)}, \dots, \hat{\theta}^{(B)})$ , the percentile CI is rector of bootstap replicates. quantile(bootstrap.reps, c(alpha/2, 1 - alpha/2))

#### Assumptions/usage

- I widely used because simple to implement & explain.

  I use when little bias and skewness in bootstrap dsn.
- (3) Drawback: CI's usually too narrow! (corrage too low)
- (4) BCa intervals usually performs better (nominal corrage).

# 1.4.2 Basic Bootstrap CI (corrects for Lins).

The  $100(1-\alpha)\%$  Basic Bootstrap CI for  $\theta$  is

$$100(1-\alpha)\%$$
 Basic Bootstrap CI for  $\theta$  is
$$\begin{pmatrix} \hat{\theta} - \begin{bmatrix} \hat{\theta} \\ -\alpha/2 \end{bmatrix} - \hat{\theta} \end{pmatrix} \qquad \hat{\theta} - \begin{bmatrix} \hat{\theta} \\ \alpha/2 \end{bmatrix} - \hat{\theta} \end{bmatrix}$$
estimate from original data set.

$$\Rightarrow \left(2\hat{\theta} - \hat{\theta}_{1-d/2}, 2\hat{\theta} - \hat{\theta}_{d/2}\right).$$

### Assumptions/usage

## 1.4.3 Standard Normal Bootstrap CI

If our statistic is of the form got we have a CLT, From the CLT,

$$\frac{\hat{\theta} - E(\hat{\theta})}{Se(\hat{\theta})} \sim N(0,1).$$

So, the  $100(1-\alpha)\%$  Standard Normal Bootstrap CI for  $\theta$  is

$$\hat{\theta} \pm Z_{1-\alpha/2}$$
 Se  $(\hat{\theta})$ 

from Lootstrap samples

 $sd(bootstrap samples) = sd(\hat{\theta}^{(1)},...,\hat{\theta}^{(0)}).$ 

Assumptions/usage

(1) 
$$\hat{\theta} \sim Normal \left( E(\hat{\theta}), Se(\hat{\theta})^2 \right) = BlG assumption if  $\hat{\theta}$  is not a sample mean!$$

(a) 
$$\hat{\theta}$$
 is unbiased  $\Rightarrow \hat{E}\hat{\theta} = \theta$  (can use bigs correction w/ this method too)  $\Rightarrow$  see later code.

13

### 1.4.4 Bootstrap t CI (Studentized Bootstrap)

Even if the distribution of  $\hat{\theta}$  is Normal and  $\hat{\theta}$  is unbiased for  $\theta$ , the Normal distribution is not exactly correct for z. (because we estimate se( $\hat{\theta}$ )).

$$t^{*} = \frac{\hat{\theta} - E(\hat{\theta})}{\hat{se}(\hat{\theta})}$$
  $\wedge t_{n}$ ? Additionally, the distribution of  $\hat{se}(\hat{\theta})$  is unknown.

 $\Rightarrow$  The bootstrap t interval does not use a Student t distribution as the reference distribuion, instead we estimate the distribution of a "t type" statistic by resampling.

The 
$$100(1-\alpha)\%$$
 Boostrap  $t$  CI is quentiles of the bootstrap "t-type" statistic  $(\hat{\theta} - t_{-\alpha/2}^{*}, se(\hat{\theta}))$  of  $t_{-\alpha/2}^{*}$  se( $\hat{\theta}$ ).

Overview

+-type statistic: 
$$t^{(i)} = \frac{\hat{\theta}^{(i)} - \hat{\theta}}{\hat{se}(\hat{\theta}^{(i)})}$$
,...,  $t^{(B)} = \frac{\hat{\theta}^{(B)} - \hat{\theta}}{\hat{se}(\hat{\theta}^{(B)})}$ .

To astimate the "t style distribution" for  $\hat{\theta}$  bootstrap estinate of se of  $\hat{\theta}$  based on  $1^{th}$ 

To estimate the "t style distribution" for  $\hat{\theta}$ , Lootstrap sample.

a. french replicate b=1,..., B

a) simple of replacement from 
$$X$$
  

$$\chi^{(b)} = (\chi_1^{(b)}, \dots, \chi_n^{(b)}).$$

b) Compute ô Cb).

C) for each toplicate 
$$r = 1, ..., R$$

i) sample  $w/$  replacement from  $\chi^{(b)}$ 
 $\chi^{(b)(r)} = (\chi^{(5)(r)}, ..., \chi^{(b)(r)})$ 

ii)  $\hat{\theta}^{(5)(r)}$  based on  $\chi^{(5)(r)}$ .

d) compute 
$$\hat{Se}(\hat{\theta}^{(\omega)}) = sd(\hat{\theta}^{(\omega(1)}, \dots, \hat{\theta}^{(b)(R)}).$$

DOUBLE BOOT STRAPI WOAH.

e) compute 
$$t = -\frac{1}{5}$$
 =  $\frac{\hat{\theta}^{(b)} - \hat{\theta}}{\hat{se}(\hat{\theta}^{(b)})}$ .

### Assumptions/usage

(1) Requires small bias and skowness in Soutstrap den.

\*\*X 2) Computationally intensive

\*\*3 Need & independent sec(8).

1.4 Bootstrap CIs

### 1.4.5 BCa CIs

Modified version of percentile intervals that adjusts for bias of estimator and skewness of the sampling distribution.

This method automatically selects a transformation so that the normality assumption holds.

Idea:

Assume there exists a monotonically inscasing function 
$$g$$
 and constant  $a,b$  st. 
$$U = \frac{g(\hat{\theta}) - g(\hat{\theta})}{1 + ag(\hat{\theta})} + b \sim N(O(1)).$$
 where  $1 + ag(\hat{\theta}) \geq 0$ .

The BCa method uses bootstrapping to estimate the bias and skewness then modifies which percentiles are chosen to get the appropriate confidence limits for a given data set.

In summary,

Assumptions / Usage:

(D) Better theoretical & practical performance than percentile method (butter onesage).

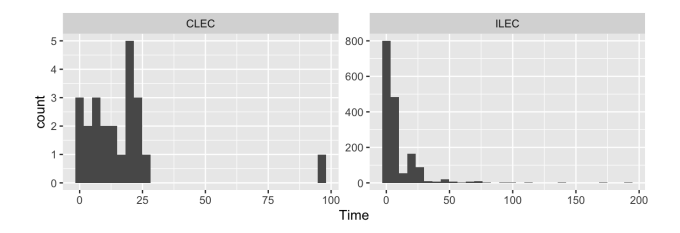
(2) Hardler to explain.

## Your Turn

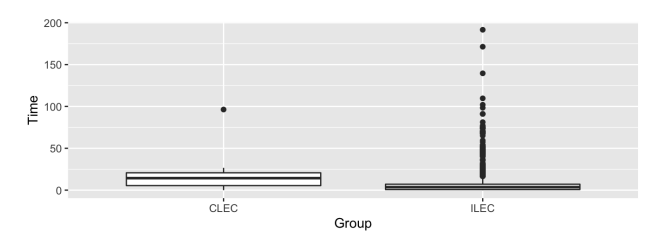
We will consider a telephone repair example from Hesterberg (2014). Verizon has repair times, with two groups, CLEC and ILEC, customers of the "Competitive" and "Incumbent" local exchange carrier.

```
verizon is regard by law to serve both at the same speed.
 library(resample) # package containing the data
 data(Verizon)
 head(Verizon)
 ##
       Time Group
 ## 1 17.50
             ILEC
       2.40
 ## 2
             ILEC
 ## 3
      0.00 ILEC
 ## 4 0.65 ILEC
 ## 5 22.23 ILEC
 ## 6
       1.20 ILEC
 Verizon %>%
   group by(Group) %>%
   summarize(mean = mean(Time), sd = sd(Time), min = min(Time), max =
  max(Time)) %>%
   kable()
                                          sd min
                   Group
                              mean
                                                  max
                                                       23
                                                 96.32
                   CLEC 16.509130 19.50358
                                                       1664
                   ILEC
                          8.411611 14.69004
                                              0 191.60
```

```
ggplot(Verizon) +
  geom_histogram(aes(Time)) +
  facet wrap(.~Group, scales = "free")
```



ggplot(Verizon) +
 geom\_boxplot(aes(Group, Time))



## 1.5 Bootstrapping CIs

There are many bootstrapping packages in R, we will use the boot package. The function boot generates R resamples of the data and computes the desired statistic(s) for each sample. This function requires 3 arguments:

- 1. data = the data from the original sample (data.frame or matrix).
- 2. statistic = a function to compute the statistic from the data where the first argument is the data and the second argument is the indices of the obervations in the boostrap sample.
- 3. R = the number of bootstrap replicates.

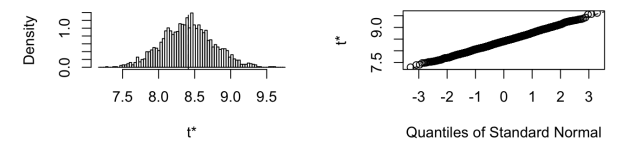
```
library(boot) # package containing the bootstrap function

mean_func <- function(x, idx) {
    mean(x[idx])
}

ilec_times <- Verizon[Verizon$Group == "ILEC",]$Time
boot.ilec <- boot(ilec_times, mean_func, 2000)

plot(boot.ilec)</pre>
```

### Histogram of t



If we want to get Bootstrap CIs, we can use the boot.ci function to generate the 5 different nonparamteric bootstrap confidence intervals.

```
boot.ci(boot.ilec, conf = .95, type = c("perc", "basic", "norm",
    "bca"))

## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 2000 bootstrap replicates
##
## CALL:
## boot.ci(boot.out = boot.ilec, conf = 0.95, type = c("perc", "basic",
```

```
"norm", "bca"))
##
##
## Intervals :
## Level
              Normal
                                   Basic
## 95%
       (7.719, 9.114) (7.709, 9.119)
##
## Level
             Percentile
                                     BCa
       (7.704, 9.114) (7.752, 9.164)
## 95%
## Calculations and Intervals on Original Scale
## we can do some of these on our own
## normal
mean(boot.ilec\$t) + c(-1, 1)*qnorm(.975)*sd(boot.ilec\$t)
          bootstap scaples of ô
## [1] 7.709670 9.104182
in boot package, Normal bootstap CI ## normal is bias corrected
2*mean(ilec_times) - (mean(boot.ilec$t) - c(-1,
 1)*qnorm(.975)*sd(boot.ilec$t))
## [1] 7.719039 9.113551
## percentile
quantile(boot.ilec$t, c(.025, .975))
##
       2.5%
               97.5%
## 7.707656 9.111150
2*mean(ilec_times) - quantile(boot.ilec$t, c(.975, .025))
      97.5%
                2.5%
## 7.712071 9.115565
```

To get the studentized bootstrap CI, we need our statistic function to also return the variance of  $\hat{\theta}$ .

```
mean var func <- function(x, idx) {</pre>
  c(mean(x[idx]), var(x[idx])/length(idx))
}
boot.ilec_2 <- boot(ilec_times, mean_var_func, 2000)</pre>
boot.ci(boot.ilec 2, conf = .95, type = "stud")
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 2000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = boot.ilec_2, conf = 0.95, type = "stud")
## Intervals :
## Level
           Studentized
## 95%
       (7.733, 9.231)
## Calculations and Intervals on Original Scale
```

Which CI should we use?

## 1.6 Bootstrapping for the difference of two means

Given iid draws of size n and m from two populations, to compare the means of the two groups using the bootstrap,

The function two.boot in the simpleboot package is used to bootstrap the difference between univariate statistics. Use the bootstrap to compute the shape, bias, and bootstrap sample error for the samples from the Verizon data set of CLEC and ILEC customers.

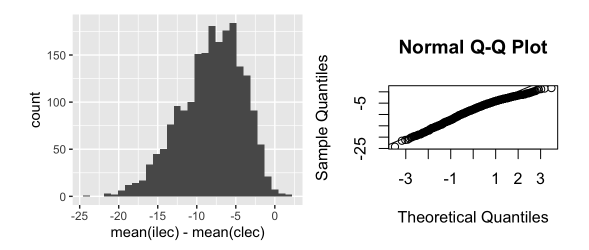
```
library(simpleboot)

clec_times <- Verizon[Verizon$Group == "CLEC",]$Time

diff_means.boot <- two.boot(ilec_times, clec_times, "mean", R = 2000)

ggplot() +
    geom_histogram(aes(diff_means.boot$t)) +
    xlab("mean(ilec) - mean(clec)")

qqnorm(diff_means.boot$t)
qqline(diff_means.boot$t)</pre>
```



# Your turn: estimate the bias and se of the sampling distribution

Which confidence intervals should we use?

# Your turn: get the chosen CI using boot.ci

Is there evidence that

$$H_0: \mu_1 - \mu_2 = 0 \ H_a: \mu_1 - \mu_2 < 0$$

is rejected?

# 2 Parametric Bootstrap

In a nonparametric bootstrap, we

In a parametric bootstrap,

For both methods,

# 2.1 Bootstrapping for linear regression

Consider the regression model  $Y_i = oldsymbol{x}_i^T oldsymbol{eta} + \epsilon_i, i = 1, \ldots, n ext{ with } \epsilon_i \overset{iid}{\sim} N(0, \sigma^2).$ 

Two approaches for bootstrapping linear regression models –

1.

2.

### 2.1.1 Bootstrapping the residuals

- 1. Fit the regression model using the original data
- 2. Compute the residuals from the regression model,

$$\hat{oldsymbol{\epsilon}}_i = y_i - \hat{oldsymbol{y}}_i = y_i - oldsymbol{x}_i^T \hat{oldsymbol{eta}}, \quad i = 1, \dots, n$$

- 3. Sample  $\hat{\epsilon}_1^*, \dots, \hat{\epsilon}_n^*$  with replacement from  $\hat{\epsilon}_1, \dots, \hat{\epsilon}_n$ .
- 4. Create the bootstrap sample

$$y_i^* = oldsymbol{x}_i^T \hat{oldsymbol{eta}} + \epsilon_i^*, \quad i = 1, \dots, n$$

- 5. Estimate  $\hat{\boldsymbol{\beta}}^*$
- 6. Repeat steps 2-4 B times to create B bootstrap estimates of  $\hat{\beta}$ .

#### **Assumptions:**

## 2.1.2 Paired bootstrapping

Resample  $z_i^* = (y_i, \boldsymbol{x}_i)^*$  from the empirical distribution of the pairs  $(y_i, \boldsymbol{x}_i)$ .

### **Assumptions:**

## 2.1.3 Which to use?

- 1. Standard inferences -
- 2. Bootstrapping the residuals -

3. Paired bootstrapping -

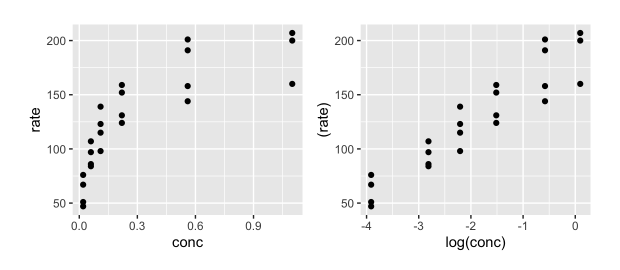
# Your Turn

This data set is the Puromycin data in R. The goal is to create a regression model about the rate of an enzymatic reaction as a function of the substrate concentration.

```
head(Puromycin)
##
     conc rate
                 state
## 1 0.02
            76 treated
## 2 0.02
            47 treated
## 3 0.06
            97 treated
## 4 0.06
           107 treated
## 5 0.11
           123 treated
## 6 0.11
           139 treated
dim(Puromycin)
## [1] 23
          3
ggplot(Puromycin) +
  geom_point(aes(conc, rate))
```

ggplot(Puromycin) +

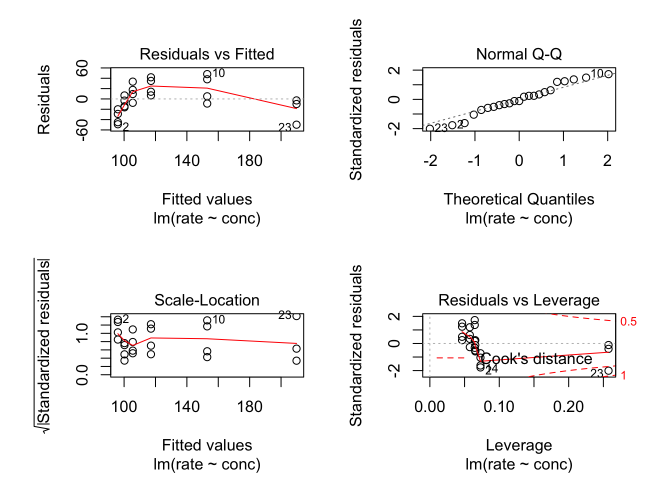
geom\_point(aes(log(conc), (rate)))

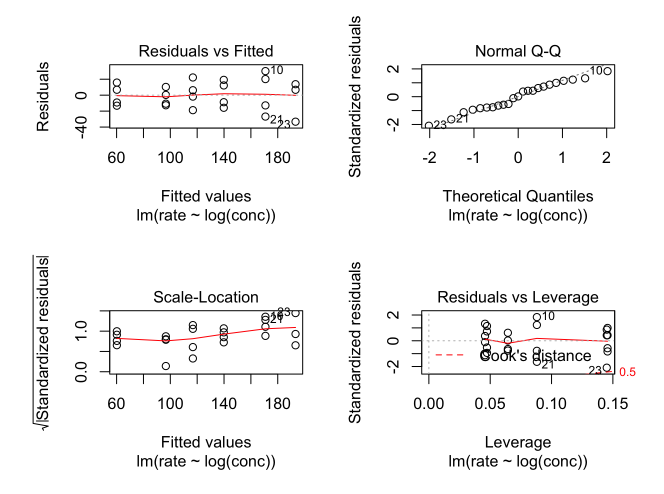


### 2.1.4 Standard regression

```
m0 <- lm(rate ~ conc, data = Puromycin)
plot(m0)
summary(m0)
##
## Call:
## lm(formula = rate ~ conc, data = Puromycin)
## Residuals:
               1Q Median
                               3Q
                                      Max
## -49.861 -15.247 -2.861 15.686 48.054
##
## Coefficients:
      Estimate Std. Error t value Pr(>|t|)
## (Intercept) 93.92
                           8.00 11.74 1.09e-10 ***
## conc
               105.40 16.92 6.23 3.53e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 28.82 on 21 degrees of freedom
## Multiple R-squared: 0.6489, Adjusted R-squared: 0.6322
## F-statistic: 38.81 on 1 and 21 DF, p-value: 3.526e-06
confint(m0)
##
                 2.5 % 97.5 %
## (Intercept) 77.28643 110.5607
## conc
             70.21281 140.5832
m1 <- lm(rate ~ log(conc), data = Puromycin)</pre>
plot(m1)
summary(m1)
##
## Call:
## lm(formula = rate ~ log(conc), data = Puromycin)
##
```

```
## Residuals:
              1Q Median
##
      Min
                              3Q
                                     Max
## -33.250 -12.753 0.327 12.969 30.166
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                           6.332
                                   30.02 < 2e-16 ***
## (Intercept) 190.085
## log(conc)
                33.203
                           2.739 12.12 6.04e-11 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 17.2 on 21 degrees of freedom
## Multiple R-squared: 0.875, Adjusted R-squared: 0.869
## F-statistic: 146.9 on 1 and 21 DF, p-value: 6.039e-11
confint(m1)
##
                  2.5 % 97.5 %
## (Intercept) 176.91810 203.2527
## log(conc) 27.50665 38.8987
```





### 2.1.5 Paired bootstrap

```
# Your turn
library(boot)

reg_func <- function(dat, idx) {
    # write a regression function that returns fitted beta
}

# use the boot function to get the bootstrap samples

# examing the bootstrap sampling distribution, make histograms

# get confidence intervals for beta_0 and beta_1 using boot.ci</pre>
```

### 2.1.6 Bootstrapping the residuals

```
# Your turn
library(boot)

reg_func_2 <- function(dat, idx) {
    # write a regression function that returns fitted beta
    # from fitting a y that is created from the residuals
}

# use the boot function to get the bootstrap samples
# examing the bootstrap sampling distribution, make histograms
# get confidence intervals for beta_0 and beta_1 using boot.ci</pre>
```