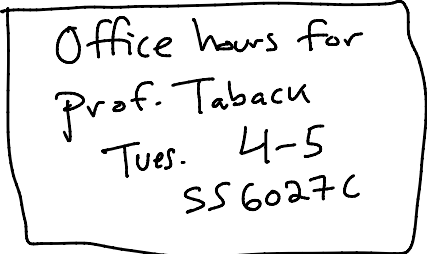


STA305/1004 - Class 4

September 17, 2019

Today's Class



Office hours for
Prof. Taback
Tues. 4-5
SS6027C

- ▶ Hypothesis testing via randomization
- ▶ Two-sample t-test
- ▶ Paired t-test

Example: Wheat Yield

- ▶ Assigning treatments randomly avoids any pre-experimental bias.
- ▶ 12 playing cards, 6 red, 6 black were shuffled (7 times??) and dealt
- ▶ 1st card black \rightarrow 1st plot gets B
- ▶ 2nd card red \rightarrow 2nd plot gets A
- ▶ 3rd card black \rightarrow 3rd plot gets B
- ▶ Completely randomized design

Wheat Yield Example

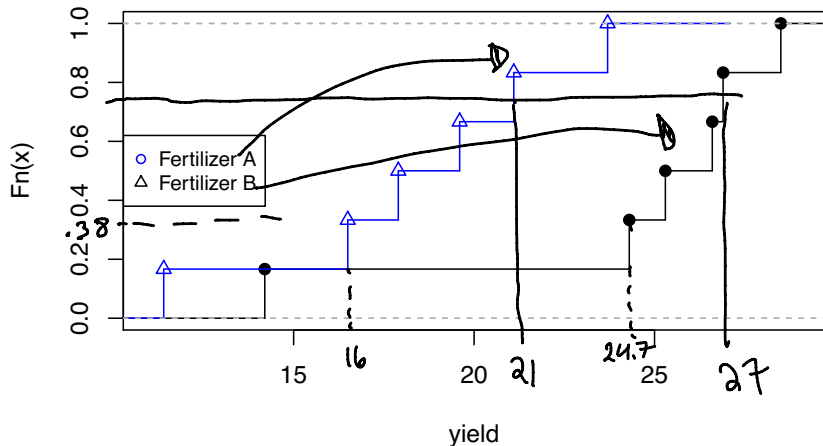
treatment assignment for this plot
Wheat yield.

B 26.9	A 11.4	B 26.6	A 23.7	B 25.3	B 28.5
B 14.2	A 17.9	A 16.5	A 21.1	B 24.3	A 19.6

- ▶ Evidence that fertilizer type is a source of yield variation?
- ▶ Evidence about differences between two populations is generally measured by comparing summary statistics across two sample populations.
- ▶ A statistic is any computable function of the observed data.

Wheat Yield Study

Empirical CDF Fertilizer



Wheat Yield Study

fert A

```
summary(yA); sd(yA); quantile(yA, prob=c(0.25, 0.75))
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
##    11.40  16.85   18.75   18.37  20.73   23.70
## [1] 4.234934
```

```
##      25%    75%
##    16.850 20.725
```

```
summary(yB); sd(yB); quantile(yB, prob=c(0.25, 0.75))
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
##    14.20  24.55   25.95   24.30  26.82   28.50
```

```
## [1] 5.151699
```

```
##      25%    75%
##    24.550 26.825
```

.4
25th
75th
.8

fert B.

IQR
= 20.725 -
16.850

Results

```
mean(yA) - mean(yB)
```

```
## [1] -5.933333
```

- ▶ So there is a moderate/large difference in mean yield for these fertilizers.
 - ▶ Would you recommend B over A for future plantings?
 - ▶ Do you think these results generalize to a larger population?
 - ▶ Could the result be due to chance?
-

Hypothesis Testing Via Randomization

- ▶ Are the observed differences in yield due to fertilizer type?
- ▶ Are the observed differences in yield due to plot-to-plot variation?

Hypothesis Testing Via Randomization

Hypothesis tests:

- ▶ H_0 (null hypothesis): Fertilizer type does not affect yield.
- ▶ H_1 (alternative hypothesis): Fertilizer type does affect yield.
- ▶ A statistical hypothesis evaluates the compatibility of H_0 with the data

Test Statistics and Null Distributions

We can evaluate H_0 by answering:

- ▶ Is a mean difference of -5.93 plausible/probable if H_0 true?
- ▶ Is a mean difference of -5.93 large compared to experimental noise?

Test Statistics and Null Distributions

- ▶ Compare $\bar{y}_a - \bar{y}_b = -5.93$ (observed difference in the experiment) to values of $\bar{y}_a - \bar{y}_b$ that could have been observed if H_0 were true.
- ▶ Hypothetical values of $\bar{y}_a - \bar{y}_b$ that could have been observed under H_0 are referred to as samples from the null distribution.

Test Statistics and Null Distributions

- ▶ $\bar{y}_a - \bar{y}_b$ is a function of the outcome of the experiment.
- ▶ If a different experiment were performed then we would obtain a different value of $\bar{y}_a - \bar{y}_b$.

Test Statistics and Null Distributions

- ▶ In this experiment we observed $\bar{y}_a - \bar{y}_b = -5.93$.
- ▶ If there was no difference between fertilizers then what other possible values of $\bar{y}_a - \bar{y}_b$ could have been observed?

Experimental Procedure and Potential Outcomes

The cards were shuffled and we were dealt B, R, B, R, ...

B	A	B	A	B	B
B	A	A	A	B	A



Under this treatment assignment we observed the yields:

B 26.9	A 11.4	B 26.6	A 23.7	B 25.3	B 28.5
B 14.2	A 17.9	A 16.5	A 21.1	B 24.3	A 19.6

↑
observed
data

Experimental Procedure and Potential Outcomes

Another potential treatment assignment under H_0 is:

B	A	B	B	A	A
A	B	B	A	A	B

The yields obtained under this assignment are:

B 26.9	A 11.4	B 26.6	B 23.7	A 25.3	A 28.5
A 14.2	B 17.9	B 16.5	A 21.1	A 24.3	B 19.6

This data could occur if the experiment were run again.

Experimental Procedure and Potential Outcomes

Observed treatment assignment:

plot
Observed

1	2	3	4	5	6
B 26.9	A 11.4	B 26.6	A 23.7	B 25.3	B 28.5
B 14.2	A 17.9	A 16.5	A 21.1	B 24.3	A 19.6

Potential treatment assignment:

hypothetical
assignment

7	8	9	10	11	12
B 26.9	A 11.4	B 26.6	B 23.7	A 25.3	A 28.5
A 14.2	B 17.9	B 16.5	A 21.1	A 24.3	B 19.6

plot 4
A 23.7
B 23.7

This is
true
 $\therefore H_0$ is
true

Respond at [PollEv.com/nathantaback](https://pollen.com/nathantaback)

Text NATHANTABACK to 37607 once to join, then A, B, C, or D

What is the difference between the observed and potential treatment assignments?

$H_0: \mu_A = \mu_B$
 μ_A, μ_B means
for fert. A, B
 $N = 72$

The wheat yields of each plot are different in observed and potential assignments, but the treatments are the same.

A

16%

The wheat yields of each plot are the same in observed and potential assignments, but the treatments are different.

B

~~67%~~ 67% ✓

There is no difference since it has been assumed that H_0 is true.

C

12%

The observed treatment assignment is statistically significant, but the potential treatment assignment is not.

D

40%

Experimental Procedure and Potential Outcomes

- ▶ Under this hypothetical assignment the mean difference is:

```
yA <- c(11.4, 25.3, 28.5, 14.2, 21.1, 24.3)
yB <- c(26.9, 26.6, 23.7, 17.9, 16.5, 19.6)
mean(yA) - mean(yB)
```

```
## [1] -1.066667
```

This represents an outcome of the experiment in a universe where:

1. The treatment assignment is B, A, B, B, A, A, A, B, B, A, A, B
2. H_0 is true (i.e., $\mu_A = \mu_B$, where μ_A, μ_B are the mean yields of fertilizers A and B).

The Null distribution

- ▶ What potential outcomes **could** we see if H_0 is true?
- ▶ Compute $\bar{y}_a - \bar{y}_b$ for each possible treatment assignment.

The Null Distribution

- ▶ For each treatment assignment compute

$$\delta_i = \bar{y}_a - \bar{y}_b, i = 1, 2, \dots, 924.$$

- ▶ $\{\delta_1, \delta_2, \dots, \delta_{924}\}$ enumerates all pre-randomisation outcomes assuming no treatment effect.
- ▶ Since each treatment assignment is equally likely under the null distribution, a probability distribution of experimental results if H_0 is true can be described as

$$\begin{aligned}\hat{F}(y) &= \frac{\#(\delta_i \leq y)}{924} \\ &= \frac{\sum_{k=1}^{\binom{12}{6}} I(\delta_k \leq y)}{\binom{12}{6}}\end{aligned}$$

This is called the randomisation distribution.

Randomization Distribution

- ▶ The yield is not random since the plots were not chosen randomly.
- ▶ Their assignment to treatments is random.
- ▶ The basis for building a probability distribution for $\bar{y}_a - \bar{y}_b$ comes from the randomization of fertilizers to plots.

Randomization Distribution

- ▶ This randomization results in 6 plots getting fertilizer A and the remaining 6 plots receiving fertilizer B.
- ▶ This is one of $\binom{12}{6} = 924$ equally likely randomizations that could have occurred.

Experimental Procedure and Potential Outcomes

This represents an outcome of the experiment in a universe where:

1. H_0 is true.
2. The yield will be the same regardless of which fertilizer a plot received.

For example a plot that had a yield of 26.9 given fertilizer B would have the same yield if the plot received fertilizer A if H_0 is true.

A and B are labels for treatment that are exchangeable assuming H_0 is true.

R Code for Randomization Distribution

```
yA <- c(11.4,23.7,17.9,16.5,21.1,19.6)
yB <- c(26.9,26.6,25.3,28.5,14.2,24.3)
fert <- c(yA,yB); N <- choose(12,6)
res <- numeric(N) # store the results
index <- combn(1:12,6) #Generate N treatment assignments
for (i in 1:N)
{res[i] <- mean(fert[index[,i]])-mean(fert[-index[,i]])}
index[,1:2] #output first two randomizations
```

```
##      [,1] [,2]
## [1,]    1    1
## [2,]    2    2
## [3,]    3    3
## [4,]    4    4
## [5,]    5    5
## [6,]    6    7
```

```
res[1:2] #output first two mean diffs
```

```
## [1] -5.933333 -3.500000
```

yield's under trt. A
" B

"

6 plots labelled A

6 plots
labelled B.

1:4
D A A A A A A
B B B B B B

A A A A A B
B A B B B B

Computational Note: Vectorized Coding/Functional Programming

```
m1 <- function(){ # Vectorized version
  f <- function(index){mean(fert[index]) - mean(fert[-index])}
  index <- combn(1:12,6)
  y <- lapply(1:924, function(x) index[,x])
  z <- lapply(y,f)
  return(z)
}
```

Vectorized
version of
for loop.

```
system.time(m1())
```

```
##      user      system elapsed
##    0.007      0.000      0.007
```

```
m2 <- function() { #non-vectorized version
  res <- numeric(N) # store the results
  index <- combn(1:12,6) #Generate N treatment assignments
  for (i in 1:N)
    {res[i] <- mean(fert[index[,i]]) - mean(fert[-index[,i]])}
  return(res)
}
```

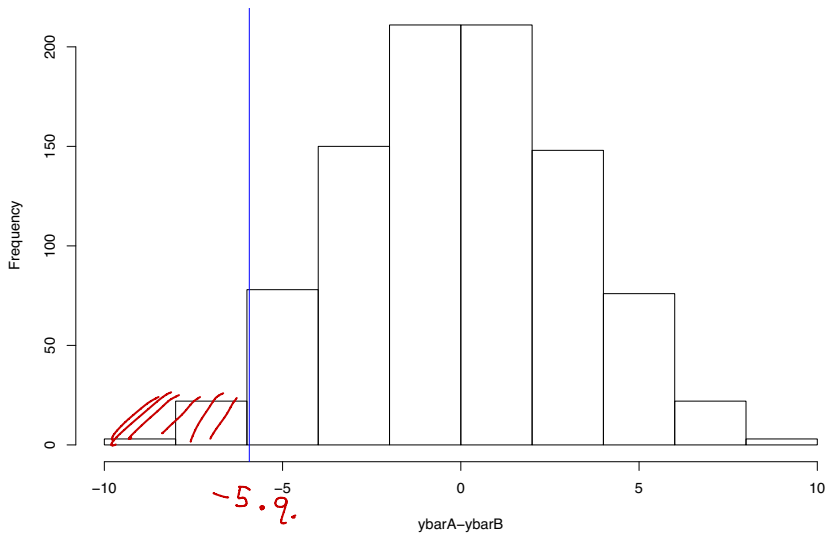
for loop.

```
system.time(m2())
```

```
##      user      system elapsed
##    0.013      0.000      0.013
```


Randomization Distribution

Randomization Distribution of difference in means



Hypothesis Testing

- ▶ Is there any contradiction between H_0 and the observed data?
- ▶ A **P-value** is the probability, under the null hypothesis of obtaining a more extreme than the observed result.

$$\text{P-value} = P(\delta \leq -5.93) = \hat{F}(-5.93)$$

- ▶ A small P-value implies evidence **against** null hypothesis.
- ▶ If the P-value is large does this imply that the null is true?

No!

Randomization Test

- ▶ Assume H_0 is true.
- ▶ Calculate the difference in means for every possible way to split the data into two samples of size 6.
- ▶ This would result in $\binom{12}{6} = 924$ differences.
- ▶ Calculate the probability of observing a value as extreme or more extreme than the observed value of the test statistic (*P-value*).
- ▶ If the P-value is small then there are two possible explanations:
 1. An unlikely value of the statistic has occurred, or
 2. The assumption that H_0 is true is incorrect.
- ▶ If the P-value is large then the hypothesis test is inconclusive.

Computing the P-value

The observed value of the test statistic is -5.93. So, the p-value is

```
# of times values from the mean randomization distribution  
# less than observed value  
sum(res<=observed)
```

```
## [1] 26
```

```
N # Number of randomizations
```

```
## [1] 924
```

```
pval <- sum(res<=observed)/N # Randomization p value  
round(pval,2)
```

```
## [1] 0.03
```

Interpretation of P-value

- ▶ A p-value of 0.03 can be interpreted as: assume there is no difference in yield between fertilizers A and B then the proportion of randomizations that would produce an observed mean difference between A and B of at most -5.93 is 0.03.
- ▶ In other words, under the assumption that there is no difference between A and B only 3% of randomizations would produce an extreme or more extreme difference than the observed mean difference.
- ▶ Therefore it's unlikely (if we consider 3% unlikely) that an observed mean difference as extreme or more extreme than -5.93 would be observed if

$$\mu_A = \mu_B.$$

evidence that fert. A is diff. than fert. B.

Two-Sided Randomization P value

- ▶ If we are using a two-sided alternative then how do we calculate a p-value?
- ▶ The randomization distribution may not be symmetric so there is no justification for simply doubling the probability in one tail.

Let

$$\bar{t} = \left(1 / \binom{N}{N_A} \right) \sum_{i=1}^{\binom{N}{N_A}} t_i$$

be the mean of the randomization distribution then we can define the two-sided p-value as

$$N = 12$$
$$N_A = 6$$

$$P(|T - \bar{t}| \geq |t^* - \bar{t}| | H_0) = \sum_{i=1}^{\binom{N}{N_A}} \frac{I(|t_i - \bar{t}| \geq |t^* - \bar{t}|)}{\binom{N}{N_A}},$$

$I(|t_i - \bar{t}| \geq |t^* - \bar{t}|)$
 $\left. \begin{array}{l} = 1 \\ 0 \end{array} \right\} \begin{array}{l} |t_i - \bar{t}| \geq \\ |t^* - \bar{t}| \\ \text{o.w} \end{array}$

The probability of obtaining an observed value of the test statistic as far, or farther, from the mean of the randomization distribution.

Two-Sided Randomization P value

```
yA <- c(11.4,23.7,17.9,16.5,21.1,19.6)
yB <- c(26.9,26.6,25.3,28.5,14.2,24.3)
fert <- c(yA,yB) #pool data
N <- choose(12,6) # randomizations
res <- numeric(N) # store the results
index <- combn(1:12,6)
for (i in 1:N)
{
  res[i] <- mean(fert[index[,i]])-mean(fert[-index[,i]])
}
tbar <- mean(res)
pval <- sum(abs(res-tbar)>=abs(observed-tbar))/N
round(pval,2)
```

```
## [1] 0.06
```

for mlt
for computing
P-Value.

Randomization Test

- ▶ We could calculate the difference in means for every possible way to split the data into two samples of size 6.
- ▶ This would result in $\binom{12}{6} = 924$ differences.
- ▶ If there were 30 observations split evenly into two groups then there are $\binom{30}{15} = 155,117,520$ differences.
- ▶ So unless the sample sizes are small these exhaustive calculations are not practical.

Randomization Test

Instead we can create a permutation resample (Monte Carlo Sampling).

1. Draw 6 observations from the pooled data without replacement. (fert A)
2. The remaining 6 observations will be the second sample (fert B)
3. Calculate the difference in means of the two samples
4. Repeat 1-3 at least 250000 times.
5. P-value is the fraction of times the random statistics exceeds the original statistic.

Estimate P-value via Monte Carlo Sampling

If M test statistics, t_i , $i = 1, \dots, M$ are randomly sampled from the permutation distribution, a one-sided Monte Carlo p value for a test of $H_0 : \mu_T = 0$ versus $H_1 : \mu_T > 0$ is

$$\hat{p} = \frac{1 + \sum_{i=1}^M I(t_i \geq t^*)}{M + 1} = \begin{cases} 1 & t_i > t^* \\ 0 & \text{o.w} \end{cases}$$

Including the observed value t^* there are $M + 1$ test statistics.

Estimate P-value via Monte Carlo Sampling

Monte Carlo Sampling.

```
N <- 250000 # number of times to repeat this process
result <- numeric(N) # space to save random diffs.
for (i in 1:N)
{ #sample of size 6, from 1 to 12, without replacement
  index <- sample(12,size=6,replace=F)
  result[i] <- mean(fert[index]) - mean(fert[-index])
}

#store observed mean difference
observed <- mean(yA) - mean(yB)

#P-value - mean - results will vary
pval <- (sum(result <= observed)+1)/(N+1)
round(pval,4)
```

```
## [1] 0.0279
```

Basic Decision Theory

Data {

	<i>Truth</i>	
	H_0 True	H_0 False
Accept H_0	correct	type II error
Reject H_0	type I error	correct

P-value = $P(\text{test statistic} \geq \text{observed value of test statistic})$

$$\alpha = P(\text{type I error})$$

$$\beta = P(\text{type II error})$$

$$1 - \beta = \text{power}$$

The Randomization P-value

Properties of the randomization test.

- ▶ An achievable P-value of the randomization test must be a multiple of $\frac{k}{\binom{12}{6}} = \frac{k}{924}$, where $k = 1, 2, \dots, 924$.
- ▶ If we choose a significance level of $\alpha = \frac{k}{924}$ that is one of the achievable P-values then $P(\text{type I error}) = \alpha$.
- ▶ The randomization test is an exact test.
- ▶ If α is not chosen to be one of the achievable P-values but $\alpha = \frac{k}{924}$ is the largest achievable P-value less than α then $P(\text{type I error}) < \alpha$.

Choosing a Test Statistic

A test statistic should be able to differentiate between H_0 and H_a in ways that are scientifically relevant.

Other Test Statistics

- ▶ Other test statistics could be used instead of $T = \bar{Y}_A - \bar{Y}_B$ to measure the effectiveness of fertilizer A.
- ▶ The difference in group medians

$$\text{median}(Y_A) - \text{median}(Y_B)$$

or trimmed means are examples of other test statistics.

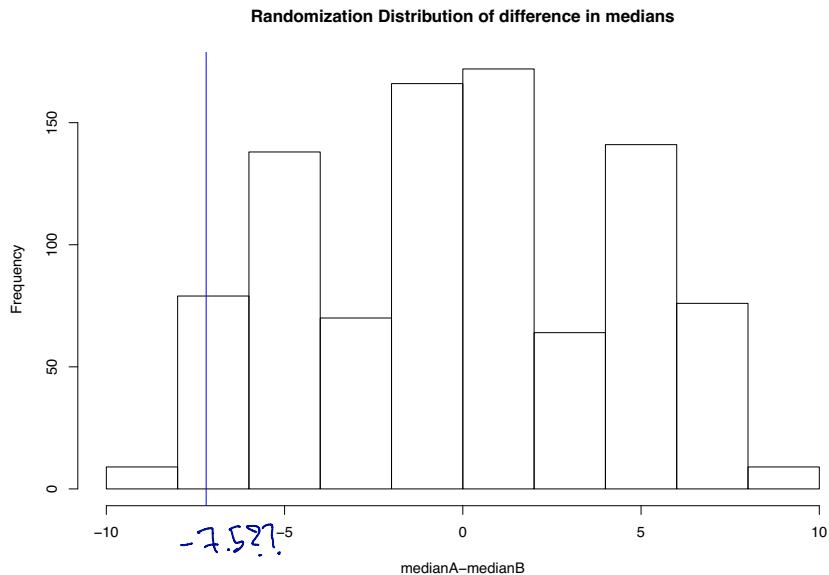
Other Test Statistics

The randomization distribution of the difference in group medians can be obtained by modifying the R code used for the difference in group means.

```
fert <- c(yA,yB) #pool data —
N <- choose(12,6) —
res <- numeric(N) # store the results —
index <-combn(1:12,6) # Generate N treatment assignments —
for (i in 1:N)
{
  res[i] <- median(fert[index[,i]])-median(fert[-index[,i]])
}
```

↑
Instead of mean
use median.

Other Test Statistics



Other Test Statistics

The p-value of the randomization test can be calculated

```
# of times values from the median randomization  
# distribution less than observed value  
sum(res<=observed)
```

```
## [1] 36
```

```
N # Number of randomizations
```

```
## [1] 924
```

```
pval <- sum(res<=observed)/N # Randomization p value  
round(pval,2)
```

```
## [1] 0.04
```

$$= 36 / 924$$

$$H_0: \tilde{\mu}_A = \tilde{\mu}_B$$

where $\tilde{\mu}_A, \tilde{\mu}_B$
are the medians of yield
under A and B.

The two-sample t-test

If the two wheat yield samples are independent random samples from a normal distribution with means μ_A and μ_B but the same variance then the statistic

$$\bar{y}_A - \bar{y}_B \sim N(\mu_A - \mu_B, \sigma^2(1/n_A + 1/n_B)).$$

So,

$$\frac{\bar{y}_A - \bar{y}_B - \delta}{\sigma \sqrt{(1/n_A + 1/n_B)}} \sim N(0, 1),$$

where $\delta = \mu_A - \mu_B$.

If we substitute

$$S^2 = \frac{\sum_{i=1}^{n_A} (y_{iA} - \bar{y}_A)^2 + \sum_{i=1}^{n_B} (y_{iB} - \bar{y}_B)^2}{n_A + n_B - 2}$$

Sample Variance

for σ^2 then

$$\frac{\bar{y}_A - \bar{y}_B - \delta}{s \sqrt{(1/n_A + 1/n_B)}} \sim t_{n_A + n_B - 2},$$

d.f.

is called the two sample t-statistic.

The two-sample t-test

In the wheat yield example $H_0 : \mu_A = \mu_B$ and suppose that $H_1 : \mu_A < \mu_B$. The p-value of the test is obtained by calculating the observed value of the two sample t-statistic under H_0 .

$$t^* = \frac{\bar{y}_A - \bar{y}_B}{s\sqrt{(1/n_A + 1/n_B)}} = \frac{18.37 - 24.3}{4.72\sqrt{(1/6 + 1/6)}} = -2.18$$

The p-value is $P(t_{18} < -2.18) = 0.03$.

The calculation was done in R.

```
s <- sqrt((5*var(yA)+5*var(yB))/10)
tstar <- (mean(yA)-mean(yB))/(s*sqrt(1/6+1/6)); round(tstar,2)
```

```
## [1] -2.18
```

```
pval <- pt(tstar,10); round(pval,5)
```

```
## [1] 0.02715
```

→ CDF t_{10}

The two-sample t-test

In R the command to run a two-sample t-test is `t.test()`.

```
t.test(yA,yB,var.equal = TRUE,alternative = "less")
```

```
##  
## Two Sample t-test  
##  
## data: yA and yB  
## t = -2.1793, df = 10, p-value = 0.02715  
## alternative hypothesis: true difference in means is less than 0  
## 95 percent confidence interval:  
## -Inf -0.9987621  
## sample estimates:  
## mean of x mean of y  
## 18.36667 24.30000
```

Handwritten: $H_a: \mu_A < \mu_B$.

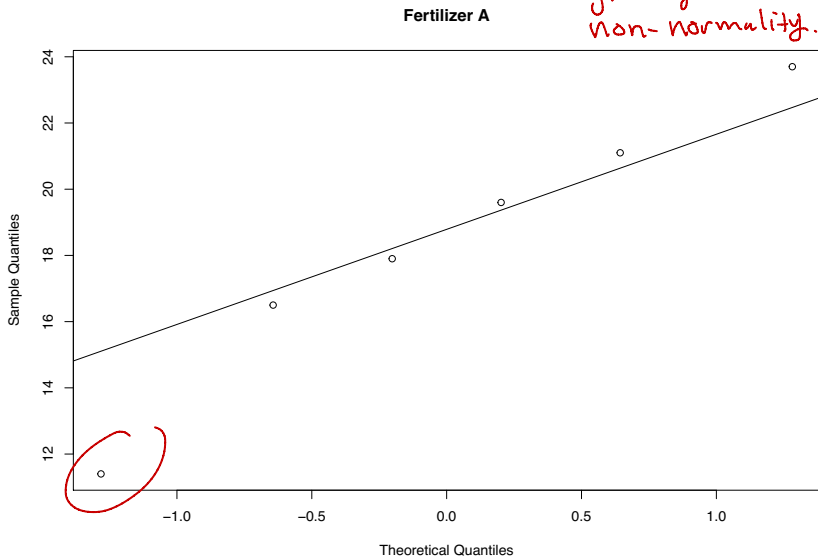
Handwritten: one-sided.

The two-sample t-test

The assumption of normality can be checked using normal quantile plots, although the t-test is robust against non-normality.

```
qqnorm(yA, main = "Fertilizer A");qqline(yA)
```

accuracy of
P-value is not
greatly affected by
non-normality.



The two-sample t-test

```
qqnorm(yB, main = "Fertilizer B"); qqline(yB)
```

QQ plots assess normality of yields under treatment.



Two-Sample t-test versus Randomization Test

- ▶ The p-value from the randomization test and the p-value from two-sample t-test are almost identical.
- ▶ The randomization test does not depend on normality or independence.

Two-Sample t-test versus Randomization Test

- ▶ The randomization test does depend on Fisher's concept that after randomization, if the null hypothesis is true, the two results obtained from each particular plot will be exchangeable.
- ▶ The randomization test tells you what you could say if exchangeability were true.

Paired Comparisons

- ▶ Increase precision by making comparisons within matched pairs of experimental material.
- ▶ Randomize within a pair.

Boy's Shoe Experiment

- ▶ Two materials to make boy's shoes, A and B, are tested to evaluate if B is more sturdy compared to A.
- ▶ During the experimental test some boys scuffed their shoes more than others.
- ▶ Each boy's two shoes were subjected to the same treatment by having each boy wear both materials.
- ▶ Working with 10 differences $B-A$ most of the boy-to-boy variation could be eliminated.
- ▶ Called a randomized paired comparison design.

Boy's Shoe Experiment

- ▶ Toss a coin to randomize material to L/R foot of a boy.
- ▶ Head: Material A used on right foot.
- ▶ Null hypothesis: amount of wear associated with material A and B are the same.
- ▶ So labelling given to a pair of results only affects the sign of the difference.

Randomized paired comparison

SS6027C.

```
library(BHH2)
data(shoes.data)
shoes.data
```

##	boy	matA	sideA	matB	sideB
## 1	1	13.2	L	14.0	R
## 2	2	8.2	L	8.8	R
## 3	3	10.9	R	11.2	L
## 4	4	14.3	L	14.2	R
## 5	5	10.7	R	11.8	L
## 6	6	6.6	L	6.4	R
## 7	7	9.5	L	9.8	R
## 8	8	10.8	L	11.3	R
## 9	9	8.8	R	9.3	L
## 10	10	13.3	L	13.6	R

matA - matB

Diff
13.2 - 14.0
8.2 - 8.8

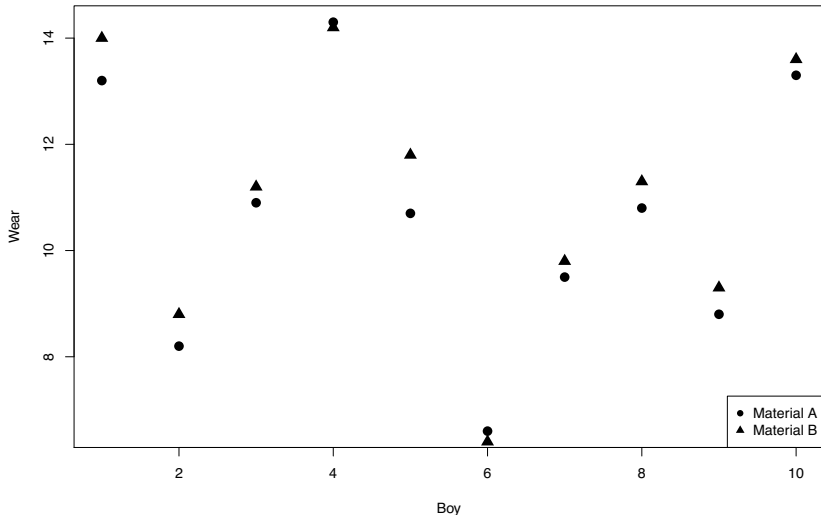
⋮

⋮

ave diff.

Randomized paired comparison

```
plot(shoes.data$boy,shoes.data$matA,pch=16,cex=1.5,  
     xlab="Boy",ylab="Wear")  
points(shoes.data$boy,shoes.data$matB,pch=17,cex=1.5)  
legend("bottomright",legend=c("Material A", "Material B"),pch=c(16,17))
```



Randomized paired comparison

```
diff <- shoes.data$matA-shoes.data$matB  
meandiff <- mean(diff); meandiff
```

```
## [1] -0.41
```

```
shoe.dat2 <- data.frame(shoes.data,diff)  
shoe.dat2
```

```
##      boy matA sideA matB sideB diff  
## 1     1 13.2     L 14.0     R -0.8  
## 2     2  8.2     L  8.8     R -0.6  
## 3     3 10.9     R 11.2     L -0.3  
## 4     4 14.3     L 14.2     R  0.1  
## 5     5 10.7     R 11.8     L -1.1  
## 6     6  6.6     L  6.4     R  0.2  
## 7     7  9.5     L  9.8     R -0.3  
## 8     8 10.8     L 11.3     R -0.5  
## 9     9  8.8     R  9.3     L -0.5  
## 10    10 13.3     L 13.6     R -0.3
```

Boy's Shoe Experiment

- ▶ The sequence of coin tosses is one of $2^{10} = 1024$ equiprobable outcomes.
- ▶ To test H_0 the average difference of -0.41 observed observed can be compared with the other 1023 averages by calculating the average difference for each of 1024 arrangements of signs in:

$$\bar{d} = \frac{\pm 0.8 \pm 0.6 \cdots \pm 0.3}{10}$$

Randomized paired comparison

```
N <- 2^(10) # number of treatment assignments
res <- numeric(N) #vector to store results
LR <- list(c(-1,1)) # difference is multiplied by -1 or 1
# generate all possible treatment assign
trtassign <- expand.grid(rep(LR, 10))
```

```
for(i in 1:N){
res[i] <- mean(as.numeric(trtassign[i,])*diff)
}
trtassign[1:2,]
```

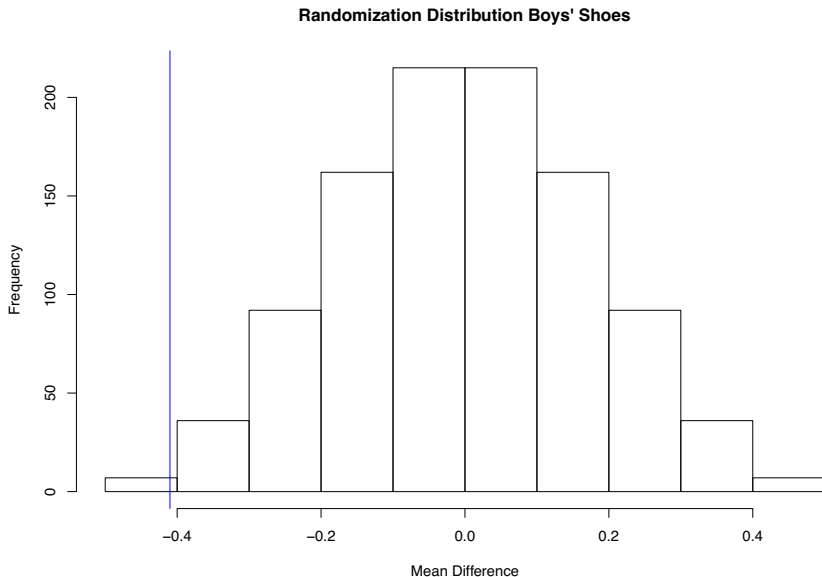
```
##   Var1 Var2 Var3 Var4 Var5 Var6 Var7 Var8 Var9 Var10
## 1   -1  -1  -1  -1  -1  -1  -1  -1  -1  -1
## 2    1  -1  -1  -1  -1  -1  -1  -1  -1  -1
```

```
res[1:2]
```

```
## [1] 0.41 0.25
```

Randomized paired comparison

```
hist(res, xlab="Mean Difference",main="Randomization Distribution Boys'  
abline(v = meandiff,col="blue")
```



Randomized paired comparison

```
sum(res<=meandiff) # number of differences le observed diff
```

```
## [1] 7
```

```
sum(res<=meandiff)/N # p-value
```

```
## [1] 0.006835938
```

Paired t-test

If we assume that the differences -0.8, -0.6, -0.3, 0.1, -1.1, 0.2, -0.3, -0.5, -0.5, -0.3 are a random sample from a normal distribution then the statistic

$$t = \frac{\bar{d}}{s_{\bar{d}}/\sqrt{10}} \sim t_{10-1},$$

where, $s_{\bar{d}}$ is the sample standard deviation of the paired differences. The p-value for testing if $\bar{D} < 0$ is

$$P(t_9 < t).$$

Paired t-test

In general if there are n differences then

$$t = \frac{\bar{d}}{s_{\bar{d}}/\sqrt{n}} \sim t_{n-1},$$

where, $s_{\bar{d}}$ is the sample standard deviation of the paired differences. The p-value for testing if $\bar{D} < 0$ is

$$P(t_{n-1} < t).$$

NB: This is the same as a one-sample t-test of the differences.

Paired t-test

In R a paired t-test can be obtained by using the command `t.test()` with `paired=T`.

```
t.test(shoes.data$matA,shoes.data$matB,paired = TRUE,  
       alternative = "less")
```

```
##  
## Paired t-test  
##  
## data: shoes.data$matA and shoes.data$matB  
## t = -3.3489, df = 9, p-value = 0.004269  
## alternative hypothesis: true difference in means is less than 0  
## 95 percent confidence interval:  
##      -Inf -0.1855736  
## sample estimates:  
## mean of the differences  
##                -0.41
```

Paired t-test

This is the same as a one-sample t-test on the difference.

```
# same as a one-sample t-test on the diff  
t.test(diff, alternative = "less")
```

```
##  
## One Sample t-test  
##  
## data: diff  
## t = -3.3489, df = 9, p-value = 0.004269  
## alternative hypothesis: true mean is less than 0  
## 95 percent confidence interval:  
##      -Inf -0.1855736  
## sample estimates:  
## mean of x  
##      -0.41
```

Paired t-test

```
qqnorm(diff); qqline(diff)
```

