

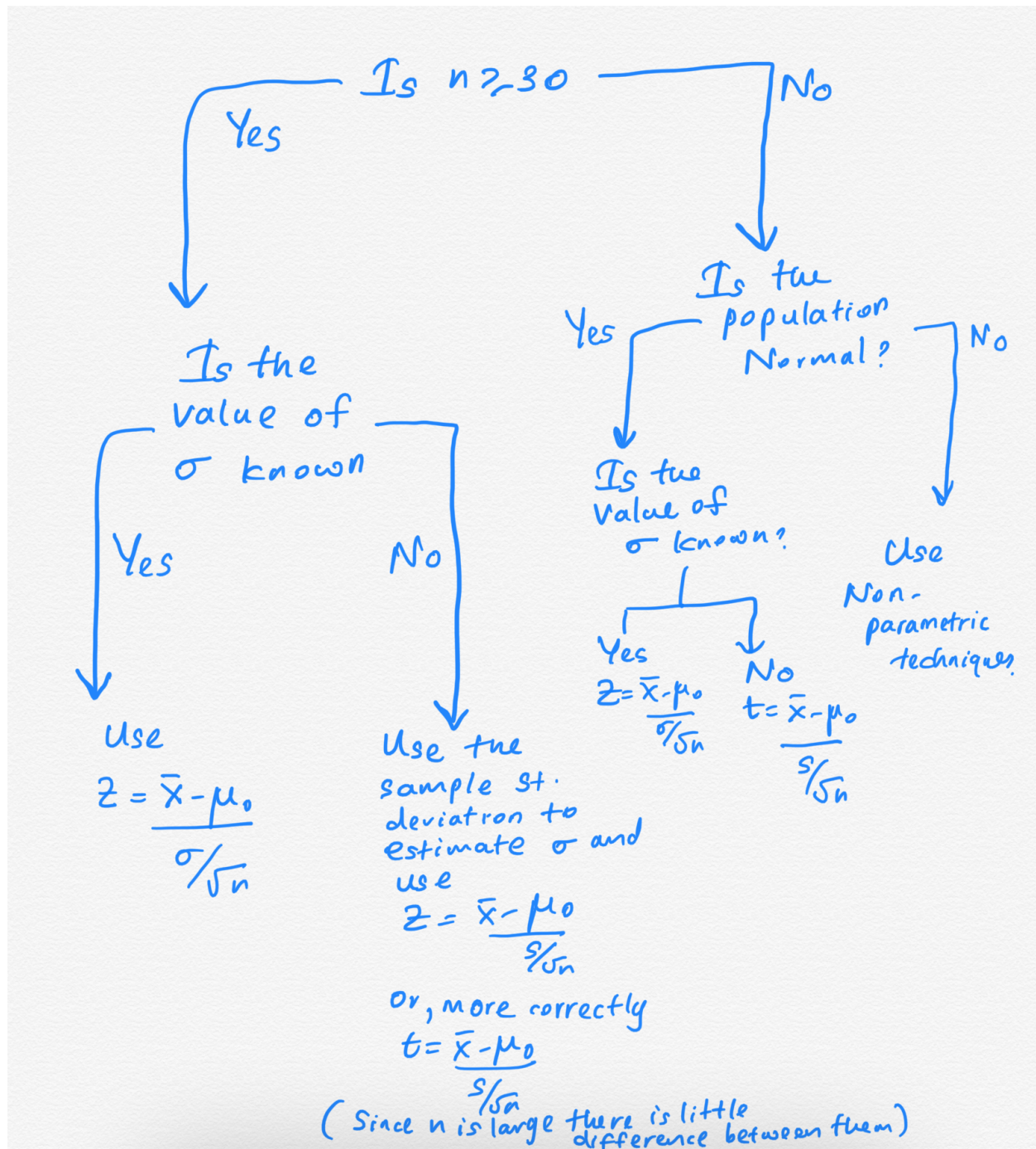
Hypothesis Testing: Comparing Means

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1. One Sample - mean



1.2 Parametric

1.2.1 Z-test (σ known)

As reported by the US National Centre for Health Statistics, the mean serum high density (HDL) cholesterol of female 20 - 29 years old is 53. Dr Jack Hall claims that the HDL Cholesterol level of female 20 - 29 years old is greater than 53. He uses the following data, randomly gathered from 22 individuals.

```
HDL <- c(65, 47, 51, 54, 70, 55, 44, 48, 36, 53, 45, 34, 59, 45, 54, 50, 40, 60, 53, 53, 54, 55)
```

It is known from past research that the distribution of the HDL cholesterol is normally distributed and the corresponding population variance is 81. Test the claim that the HDL level is greater than 53 at $\alpha = 0.01$ level of significance.

```
HDL.df <- data.frame(HDL=HDL)
ggplot(HDL.df, aes(y=HDL, x="")) +
  geom_boxplot(outlier.shape = NA, fill="forestgreen", alpha=0.5) +
  geom_jitter(alpha=0.5) + labs(x = "")
```

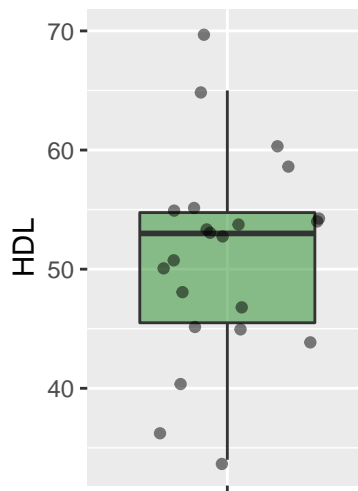


Figure 1: Distribution of HDL level

Hypothesis

H0:

H1:

μ -

```

z.test <- function(data, mu, var, alternative){
  z = (mean(data) - mu) / (sqrt(var / length(data)))
  if(alternative == "greater"){
    1-pnorm(z)

  } else if (alternative == "less"){

    pnorm(z)

  } else {

    pnorm(-1*abs(z)) * 2

  }
}
z.test(HDL.df$HDL, 53, 81, "greater")

```

```
[1] 0.8342875
```

Decision:

Conclusion:

1.2.2 t-test (σ unknown)

A chemist wants to measure the bias in a pH meter. She uses the meter to measure the pH in 14 neutral substances (pH=7) and obtains the data below.

```
ph <- c( 7.01, 7.04, 6.97, 7.00, 6.99, 6.97, 7.04, 7.04, 7.01, 7.00, 6.99, 7.04, 7.07, 6.97)
```

Is there sufficient evidence to support the claim that the pH meter is not correctly calibrated at the $\alpha = 0.05$ level of significance?

Answer:

```

ph.df <- data.frame(pH=ph)
ggplot(ph.df, aes(y=pH, x="")) +
  geom_boxplot(outlier.shape = NA) +
  geom_jitter(alpha=0.5) +
  labs(x = "")

```

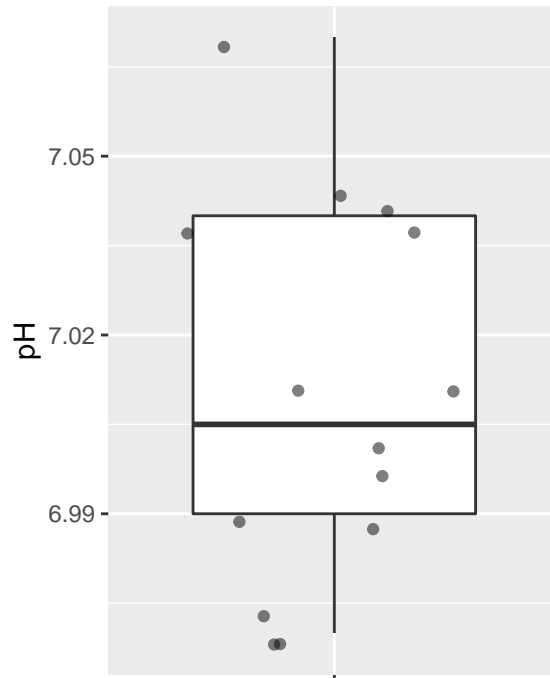


Figure 2: Distribution of pH values

In this case, we have only sixteen observations, meaning that the Central Limit Theorem does not apply. With a small sample, we should only use the t-test if we can reasonably assume that the population is normally distributed. Hence, we must first verify that pH is normally distributed.

```
ggplot(ph.df,
  aes(sample=pH))+
  stat_qq() + stat_qq_line()+labs(x="Theoretical Quantiles", y="Sample Quantiles")
```

```
shapiro.test(ph.df$pH)
```

Shapiro-Wilk normality test

```
data:  ph.df$pH
W = 0.91603, p-value = 0.1927
```

Hypothesis to be tested:

H0: Data are normally distributed.

H1: Data are not normally distributed.

According to the Shapiro-Wilk normality test p-value, $0.19 > 0.05$. Hence, we do not reject H0 at the 0.05 level of significance. We can conclude data are normally distributed.

Now we can proceed with the t.test.

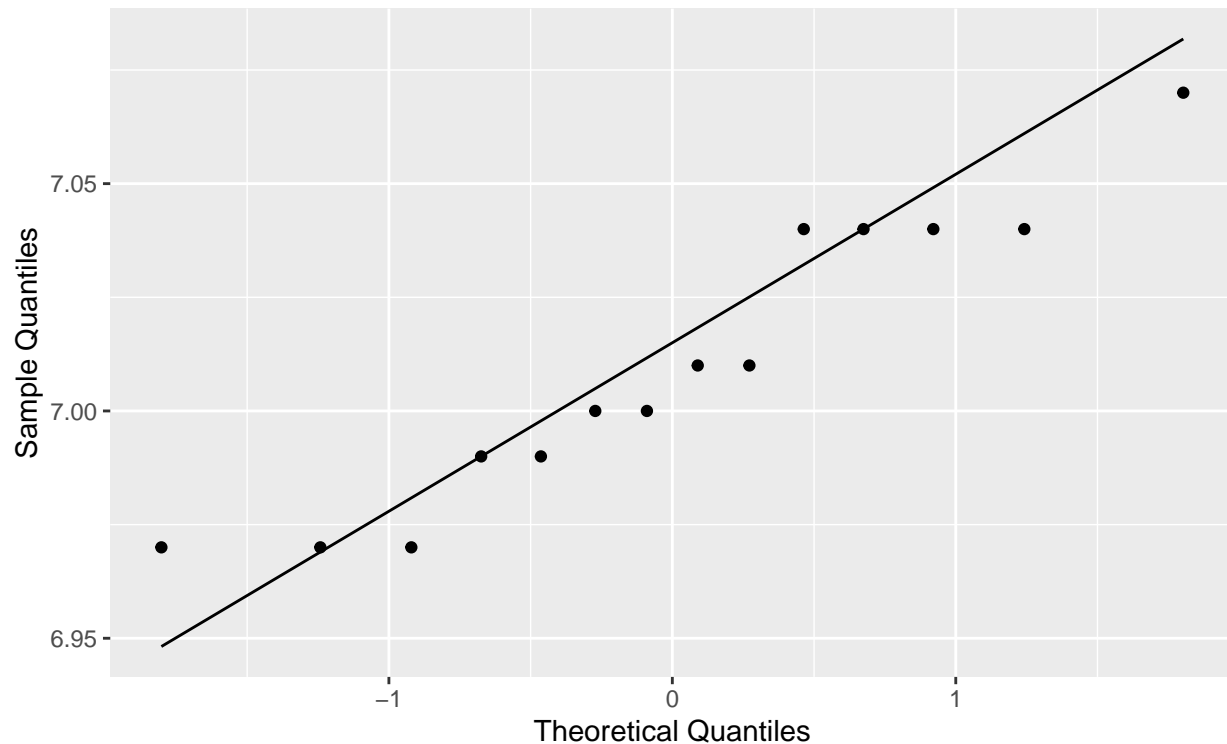


Figure 3: Normal probability plot of pH values

Hypothesis to be tested.

H0: $\mu = 7$

H1: $\mu \neq 7$

μ - Population mean pH value (in neutral substances).

t.test syntax

```
t.test(x, y = NULL,
       alternative = c("two.sided", "less", "greater"),
       mu = 0, paired = FALSE, var.equal = FALSE,
       conf.level = 0.95, ...)
```

```
t.test(ph.df$pH, alternative = "two.sided", mu=7)
```

One Sample t-test

```
data:  ph.df$pH
t = 1.1832, df = 13, p-value = 0.2579
alternative hypothesis: true mean is not equal to 7
95 percent confidence interval:
 6.991742 7.028258
sample estimates:
mean of x
 7.01
```

Decision: $p\text{-value} (0.258) > \alpha = 0.05$. Hence, we do not reject H_0 .

Conclusion: We do not have enough evidence to conclude that the population mean pH level is different from 7 at the 0.05 level of significance.

2. Two sample - mean

2.1 Dependent (paired)

Approach 1

A dietician hopes to reduce a person's cholesterol level by using a special diet supplemented with a combination of vitamin pills. Twenty (20) subjects were pre-tested and then placed on diet for two weeks. Their cholesterol levels were checked after the two week period. The results are shown below. Cholesterol levels are measured in milligrams per decilitre.

- i) Test the claim that the Cholesterol level before the special diet is greater than the Cholesterol level after the special diet at $\alpha = 0.01$ level of significance.
- ii) Construct 99% confidence interval for the difference in mean cholesterol levels. Assume that the cholesterol levels are normally distributed both before and after.

```
id <- 1:20
before <- c(210, 235, 208, 190, 172, 244, 211, 235, 210,
            190, 175, 250, 200, 270, 222, 203, 209, 220, 250, 280)
after <- c(190, 170, 210, 188, 173, 195, 228, 200, 210, 184,
           196, 208, 211, 212, 205, 221, 240, 250, 230, 220)
cholesterol_1 <- data.frame(id=id, before=before, after=after)
head(cholesterol_1)
```

	id	before	after
1	1	210	190
2	2	235	170
3	3	208	210
4	4	190	188
5	5	172	173
6	6	244	195

```
cholesterol_2 <- pivot_longer(cholesterol_1, before:after, "type", "value")
head(cholesterol_2)
```

```
# A tibble: 6 x 3
      id type  value
  <int> <chr> <dbl>
1     1 before  210
2     1 after   190
3     2 before  235
4     2 after   170
5     3 before  208
6     3 after   210
```

```
ggplot(data= cholesterol_2, aes(x=type, y=value)) +
  geom_boxplot(outlier.shape = NA, aes(fill=type), alpha=0.5) +
  geom_jitter(aes(fill=type))
```

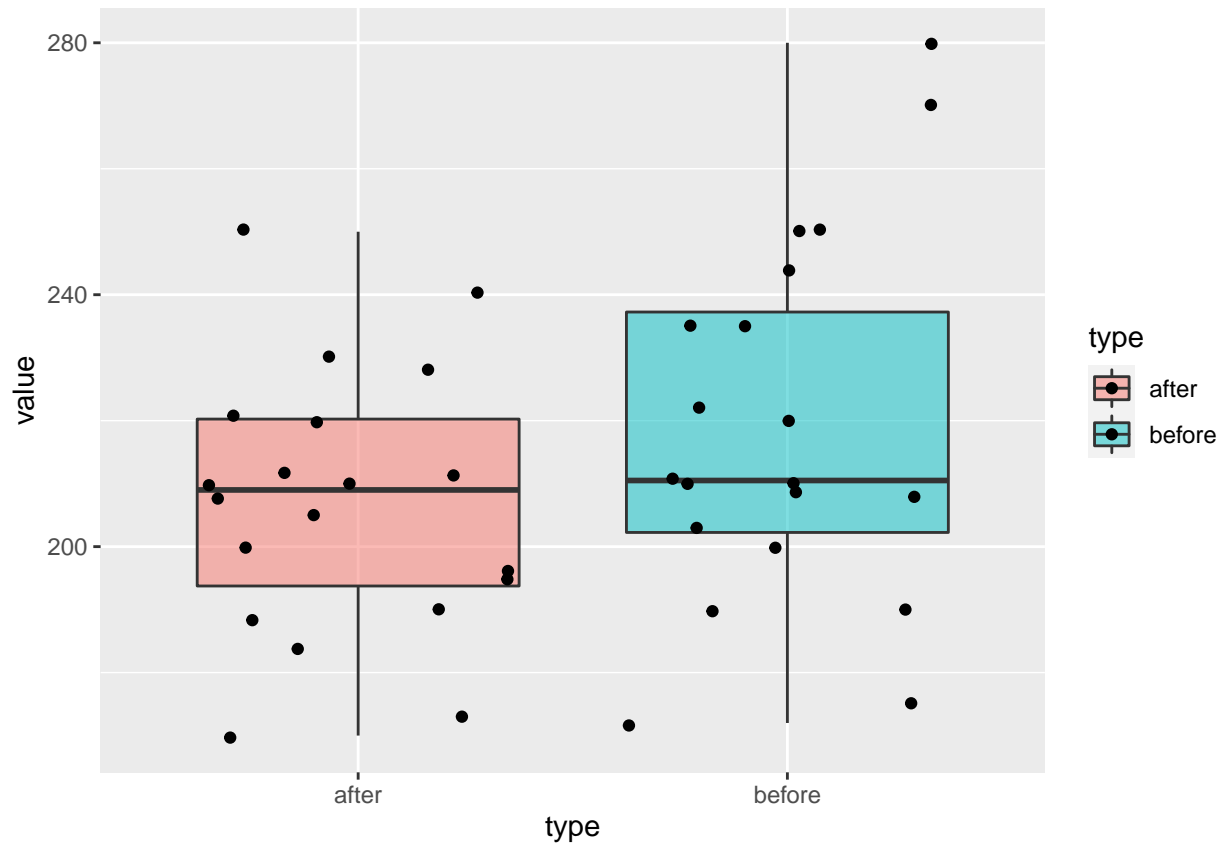


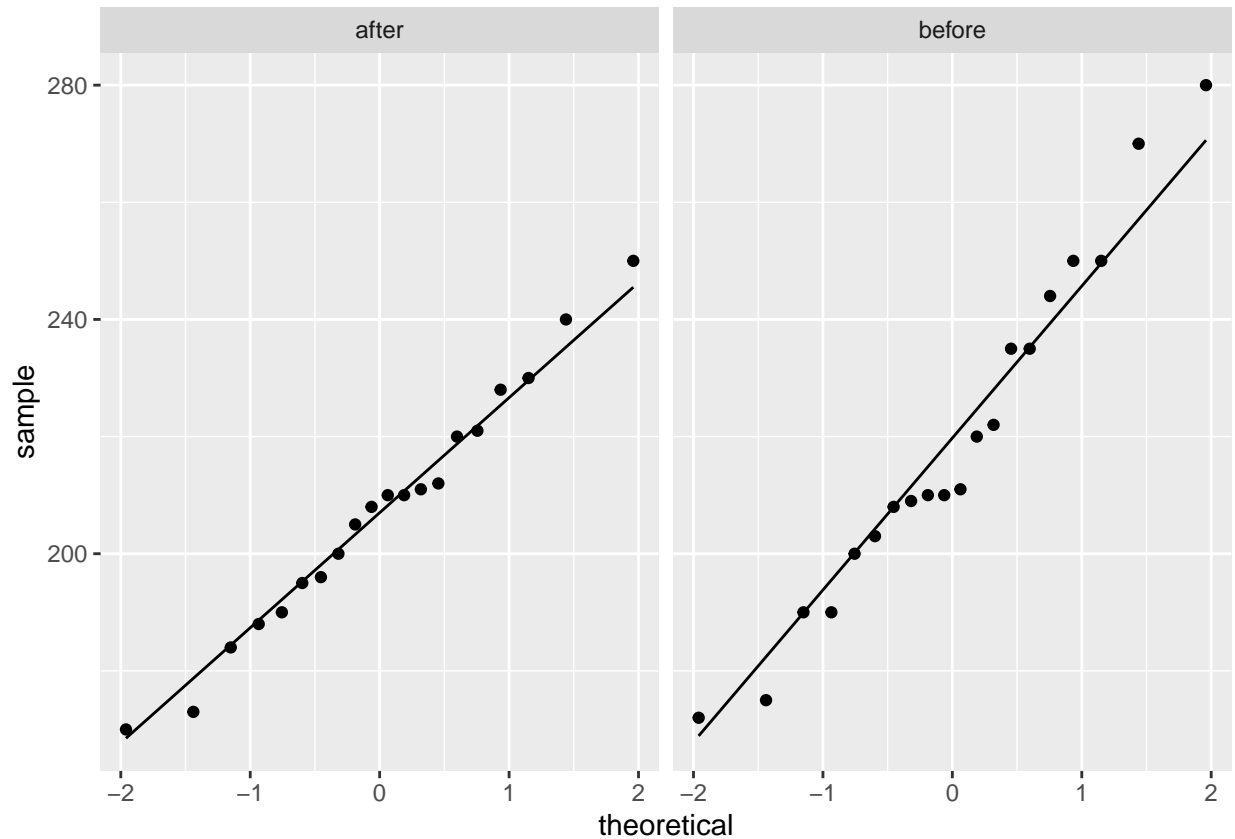
Figure 4: Distribution of cholesterol levels after and before the special diet

```
cholesterol_2 %>%
  group_by(type) %>%
  summarize(mean = round(mean(value), 2),
            sd = round(sd(value), 2))
```

```
# A tibble: 2 x 3
  type    mean    sd
<chr> <dbl> <dbl>
1 after  207.  21.0
2 before 219.  29.3
```

2.1.1 Testing for Normality

```
ggplot(data = cholesterol_2, aes(sample = value)) +
  stat_qq() +
  stat_qq_line() +
  facet_grid(. ~ type)
```

2.1.2 Paired t-test

Hypothesis:

$H_0: \mu_{before} \leq \mu_{after}$

$H_0: \mu_{before} > \mu_{after}$

μ_{before} - population mean cholesterol level before the special diet

μ_{after} - population mean cholesterol level after the special diet

```
t.test(before, after, data=cholesterol_1, "greater", paired=TRUE)
```

Paired t-test

data: before and after

t = 1.7754, df = 19, p-value = 0.04593

alternative hypothesis: true difference in means is greater than 0

95 percent confidence interval:

0.3167385 Inf

sample estimates:

mean of the differences

12.15

Decision: _____

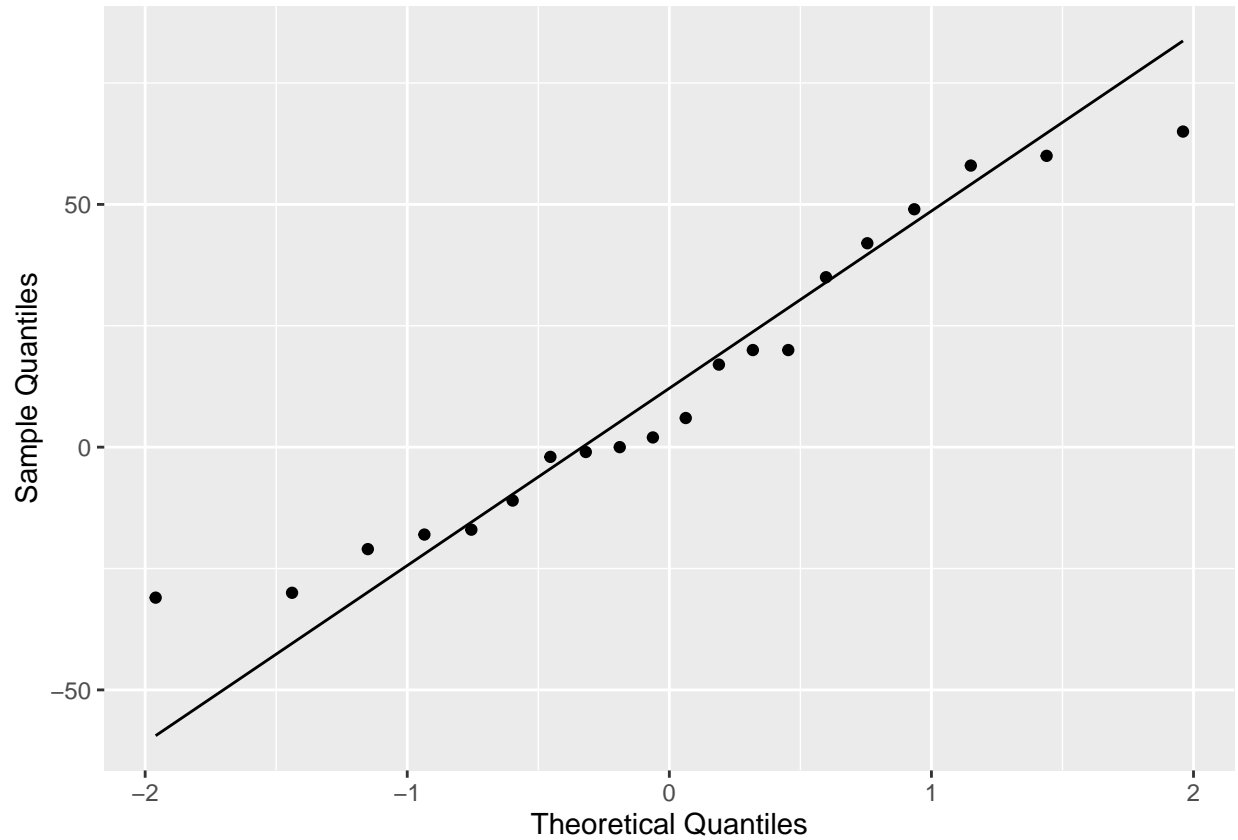
Conclusion: _____

Approach 2

```
approach2_tbl <- tibble(diff = cholesterol_1$before - cholesterol_1$after)
```

2.1.3 Testing for Normality

```
ggplot(approach2_tbl,  
  aes(sample=diff))+  
  stat_qq() + stat_qq_line()+  
  labs(x="Theoretical Quantiles", y="Sample Quantiles")
```



```
shapiro.test(approach2_tbl$diff)
```

Shapiro-Wilk normality test

data: approach2_tbl\$diff
W = 0.93729, p-value = 0.213

H0: $\mu_d \leq 0$

H0: $\mu_d > 0$,

where: $\mu_d = \mu_{before} - \mu_{after}$

```
t.test(x = approach2_tbl$diff, alternative = c("greater"), mu=0)
```

One Sample t-test

```
data: approach2_tbl$diff
t = 1.7754, df = 19, p-value = 0.04593
alternative hypothesis: true mean is greater than 0
95 percent confidence interval:
 0.3167385      Inf
sample estimates:
mean of x
    12.15
```

Decision: _____

Conclusion: _____

2.1.4 Confidence intervals

To obtain confidence intervals

```
t.test(before, after, data=cholesterol_1, "two.sided", paired=TRUE)
```

Paired t-test

```
data: before and after
t = 1.7754, df = 19, p-value = 0.09185
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 -2.173539 26.473539
sample estimates:
mean of the differences
    12.15
```

95% CI for $\mu_{before} - \mu_{after}$: _____

```
t.test(before, after, data=cholesterol_1, "two.sided", paired=TRUE, conf.level = 0.99)
```

Paired t-test

```
data: before and after
t = 1.7754, df = 19, p-value = 0.09185
alternative hypothesis: true difference in means is not equal to 0
```

99 percent confidence interval:
 -7.428709 31.728709
 sample estimates:
 mean of the differences
 12.15

99% CI for $\mu_{before} - \mu_{after}$: _____

2.2 Independent

```
birthwt <- as_tibble(MASS::birthwt)
head(birthwt)
```

```
# A tibble: 6 x 10
  low  age  lwt  race smoke  ptl  ht  ui  ftv  bwt
<int> <int> <int> <int> <int> <int> <int> <int> <int> <int>
1     0   19  182     2     0     0     0     1     0  2523
2     0   33  155     3     0     0     0     0     3  2551
3     0   20  105     1     1     0     0     0     1  2557
4     0   21  108     1     1     0     0     1     2  2594
5     0   18  107     1     1     0     0     1     0  2600
6     0   21  124     3     0     0     0     0     0  2622
```

?birthwt

smoke: smoking status during pregnancy.

(0 - No, 1 - Yes)

Is there a significant difference in birth weights between mothers who smoked during pregnancy and those who did not?

Data Wrangling

```
birthwt <- as_tibble(MASS::birthwt)

# Rename variables
birthwt <- birthwt %>%
  rename(smoking.status = smoke,
         birthwt.grams = bwt)

# Change factor level names
birthwt <- birthwt %>%
  mutate_at(c("smoking.status"),
            ~ recode_factor(.x, `0` = "no", `1` = "yes"))
head(birthwt)
```

```
# A tibble: 6 x 10
  low  age  lwt  race smoking.status  ptl  ht  ui  ftv birthwt.grams
<int> <int> <int> <int> <fct>          <int> <int> <int> <int>          <int>
1     0   19  182     2 no              0     0     1     0          2523
2     0   33  155     3 no              0     0     0     3          2551
```

3	0	20	105	1	yes	0	0	0	1	2557
4	0	21	108	1	yes	0	0	1	2	2594
5	0	18	107	1	yes	0	0	1	0	2600
6	0	21	124	3	no	0	0	0	0	2622

```
ggplot(birthwt, aes(x=smoking.status, y=birthwt.grams))+
  geom_boxplot(outlier.shape=NA, aes(fill=smoking.status), alpha=0.05) +
  geom_jitter(aes(colour=smoking.status)) +
  scale_colour_manual(values = c("#d95f02", "#7570b3"))
```

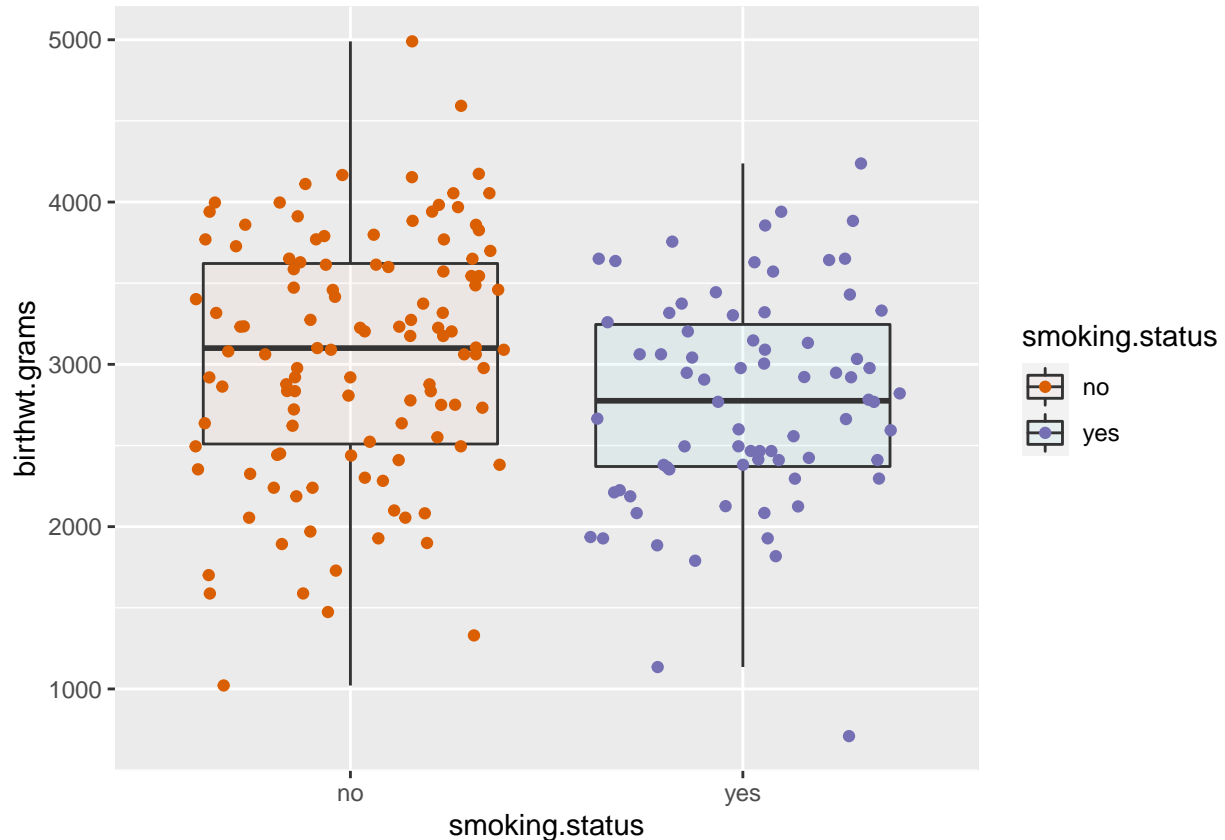


Figure 5: Distribution of infants birth weight by mothers' smoking status

```
birthwt %>%
  group_by(smoking.status) %>%
  summarize(mean = round(mean(birthwt.grams), 1),
            sd = round(sd(birthwt.grams), 1),
            max. = round(max(birthwt.grams), 1),
            min = round(min(birthwt.grams), 1),
            missing= sum(is.na(birthwt.grams)),
            count= sum(is.na(birthwt.grams)==FALSE))
```

```
# A tibble: 2 x 7
  smoking.status mean    sd max.  min missing count
  <fct>          <dbl> <dbl> <dbl> <dbl>   <int> <int>
```

1	no	3056.	753.	4990	1021	0	115
2	yes	2772.	660.	4238	709	0	74

```
se <- function(data){
  sd(data)/sqrt(length(data))
}
```

```
birthwt %>%
  group_by(smoking.status) %>%
  summarize(mean = round(mean(birthwt.grams), 1),
            sd = round(sd(birthwt.grams), 1),
            max. = round(max(birthwt.grams), 1),
            min = round(min(birthwt.grams), 1),
            missing= sum(is.na(birthwt.grams)),
            count= sum(is.na(birthwt.grams)==FALSE),
            se = se(birthwt.grams))
```

```
# A tibble: 2 x 8
  smoking.status mean    sd max.   min missing count    se
  <fct>          <dbl> <dbl> <dbl> <dbl>   <int> <int> <dbl>
1 no            3056.  753. 4990  1021     0    115  70.2
2 yes            2772.  660. 4238   709     0     74  76.7
```

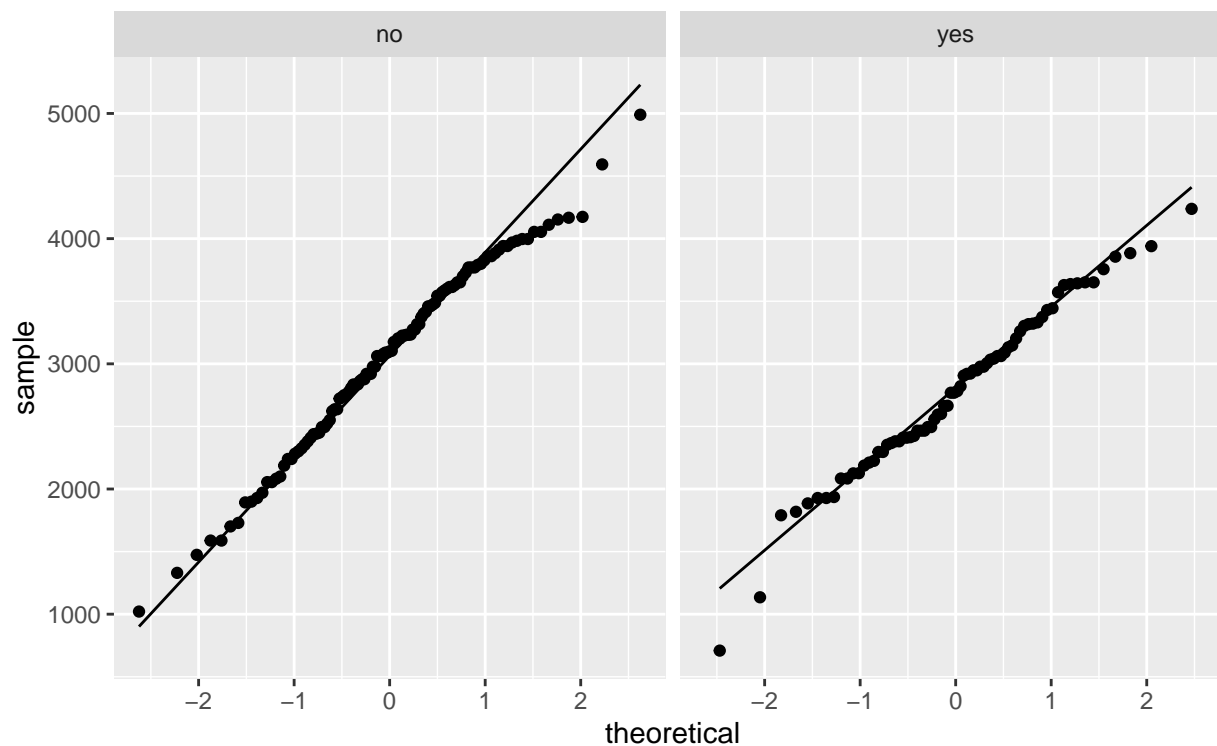
```
birthwt %>%
  group_by(smoking.status) %>%
  summarize(num.obs = n(),
            mean.birthwt = round(mean(birthwt.grams), 0),
            sd.birthwt = round(sd(birthwt.grams), 0),
            se.birthwt = round(sd(birthwt.grams) / sqrt(num.obs), 0))
```

`summarise()` ungrouping output (override with `.groups` argument)

```
# A tibble: 2 x 5
  smoking.status num.obs mean.birthwt sd.birthwt se.birthwt
  <fct>          <int>      <dbl>      <dbl>      <dbl>
1 no            115      3056      753      70
2 yes            74      2772      660      77
```

2.2.1 Testing for Normality

```
ggplot(data = birthwt, aes(sample = birthwt.grams)) +
  stat_qq() +
  stat_qq_line() +
  facet_grid(. ~ smoking.status)
```



```
mother_yes_birthwt <- birthwt %>% filter(smoking.status=="yes")
dim(mother_yes_birthwt)
```

```
[1] 74 10
```

```
shapiro.test(mother_yes_birthwt$birthwt.grams)
```

Shapiro-Wilk normality test

data: mother_yes_birthwt\$birthwt.grams
W = 0.98296, p-value = 0.4195

Hypothesis:

H0:

H1:

Decision: _____

Conclusion: _____

```
mother_no_birthwt <- birthwt %>% filter(smoking.status=="no")
dim(mother_no_birthwt)
```

```
[1] 115 10
```

```
shapiro.test(mother_no_birthwt$birthwt.grams)
```

Shapiro-Wilk normality test

```
data: mother_no_birthwt$birthwt.grams  
W = 0.98694, p-value = 0.3337
```

Hypothesis:

H0:

H1:

Decision: _____

Conclusion: _____

2.2.2 Equality of variance

The equality of variances between two samples can be tested using the F test.

Hypothesis:

H0: _____

H1: _____

σ_1^2 -

σ_2^2 -

```
var.test(birthwt.grams ~ smoking.status, data = birthwt,  
         alternative = "two.sided")
```

F test to compare two variances

```
data: birthwt.grams by smoking.status  
F = 1.3019, num df = 114, denom df = 73, p-value = 0.2254  
alternative hypothesis: true ratio of variances is not equal to 1  
95 percent confidence interval:  
 0.8486407 1.9589574  
sample estimates:  
ratio of variances  
      1.301927
```

2.2.3 How can we assess whether the mean difference is statistically significant?

Hypothesis

H0: _____

H1: _____

where,

μ_1 -

μ_2 -


```
t.test(birthwt.grams ~ smoking.status, data = birthwt,  
       alternative = c("two.sided"),  
       var.equal = TRUE)
```

Two Sample t-test

```
data: birthwt.grams by smoking.status  
t = 2.6529, df = 187, p-value = 0.008667  
alternative hypothesis: true difference in means is not equal to 0  
95 percent confidence interval:  
 72.75612 494.79735  
sample estimates:  
mean in group no mean in group yes  
 3055.696      2771.919
```

3. Other test functions

- `fisher.test` Fisher's exact test for counts
- `t.test(data)` 1 sample t test
- `t.test(data1,data2)` 2 sample t test
- `t.test(pre,post,paired=TRUE)` paired sample t test
- `wilcox.test(data)` Wilcoxon test
- `cor.test(data1,data2)` correlation test
- `chisq.test(data)` Chi square test
- `shapiro.test(data)` Shapiro test
- `aov()` ANOVA
- etc.