BB503/BB602 - R Training - Week XI

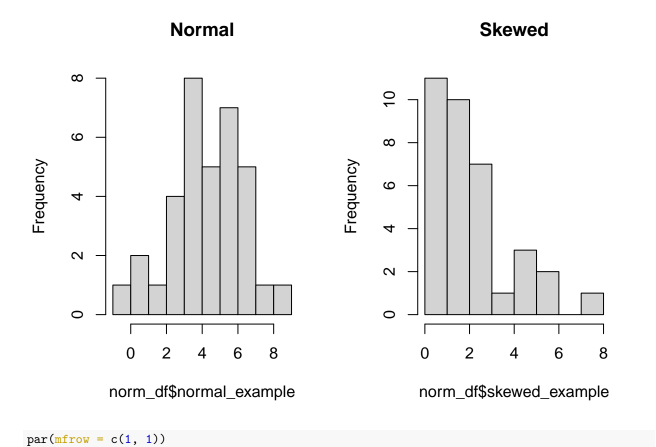
Ege Ulgen

Assessing Normality

We'll be using "Normal" dataset for this exercise. This dataset contains 2 continuous variables where one is an example of normally distributed data and the other one is an example of skewed data.

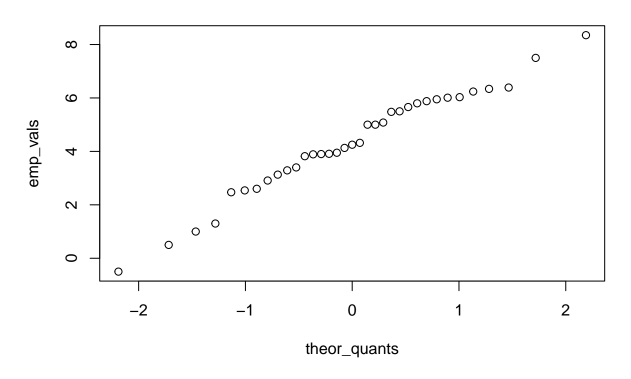
Histogram

```
par(mfrow = c(1, 2))
hist(norm_df$normal_example, main = "Normal")
hist(norm_df$skewed_example, main = "Skewed")
```



Quantile-Quantile Plot

```
# manually
emp_vals <- sort(norm_df$normal_example)
theor_quants <- qnorm((seq_along(emp_vals) - .5) / length(emp_vals))
plot(theor_quants, emp_vals)</pre>
```

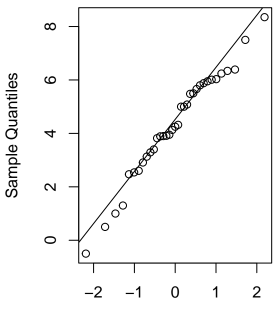


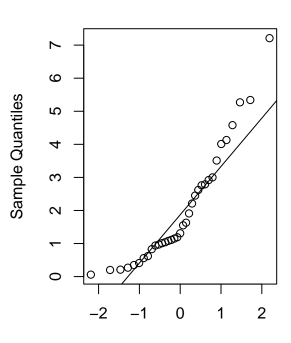
```
# using base R function 'qqnorm'
par(mfrow = c(1, 2))
qqnorm(norm_df$normal_example)
qqline(norm_df$normal_example)

qqnorm(norm_df$skewed_example)
qqline(norm_df$skewed_example)
```

Normal Q-Q Plot

Normal Q-Q Plot

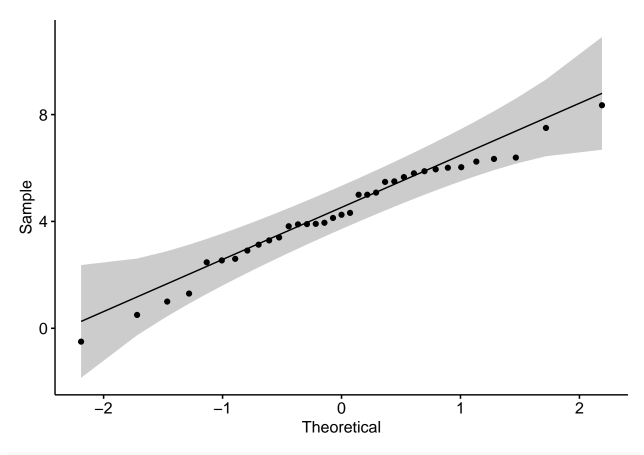




Theoretical Quantiles

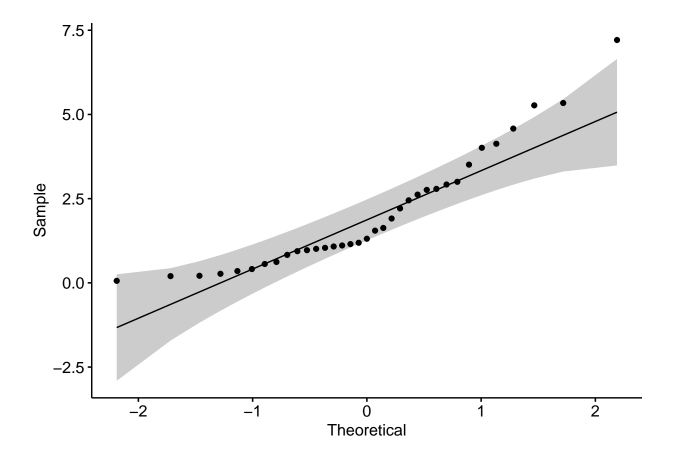
Theoretical Quantiles

```
par(mfrow = c(1, 1))
# using 'ggqqplot' from 'ggpubr'
# install.packages("ggpubr")
library(ggpubr)
## Loading required package: ggplot2
ggqqplot(norm_df, "normal_example")
## Warning: The following aesthetics were dropped during statistical transformation: sample
## i This can happen when ggplot fails to infer the correct grouping structure in
##
## i Did you forget to specify a `group` aesthetic or to convert a numerical
     variable into a factor?
## The following aesthetics were dropped during statistical transformation: sample
## i This can happen when ggplot fails to infer the correct grouping structure in
     the data.
## i Did you forget to specify a `group` aesthetic or to convert a numerical
     variable into a factor?
```



ggqqplot(norm_df, "skewed_example")

```
## Warning: The following aesthetics were dropped during statistical transformation: sample
## i This can happen when ggplot fails to infer the correct grouping structure in
## the data.
## i Did you forget to specify a `group` aesthetic or to convert a numerical
## variable into a factor?
## The following aesthetics were dropped during statistical transformation: sample
## i This can happen when ggplot fails to infer the correct grouping structure in
## the data.
## i Did you forget to specify a `group` aesthetic or to convert a numerical
## variable into a factor?
```



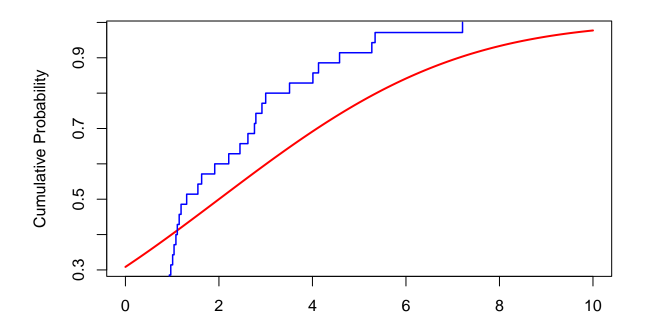
Shapiro-Wilk Test of Normality

```
H_0: the population is normally distributed \\ shapiro.test(norm_df$normal_example)
```

```
##
## Shapiro-Wilk normality test
##
## data: norm_df$normal_example
## W = 0.975, p-value = 0.58
shapiro.test(norm_df$skewed_example)
##
```

```
##
## Shapiro-Wilk normality test
##
## data: norm_df$skewed_example
## W = 0.885, p-value = 0.0016
```

Kolmogorov-Smirnov Tests



```
# does x come from the selected distribution with the specified parameters?
ks.test(norm_df$normal_example, pnorm, mean = mean(norm_df$normal_example), sd = sd(norm_df$normal_example)
## Warning in ks.test.default(norm_df$normal_example, pnorm, mean =
## mean(norm_df$normal_example), : ties should not be present for the
## Kolmogorov-Smirnov test
    Asymptotic one-sample Kolmogorov-Smirnov test
##
## data: norm_df$normal_example
## D = 0.0948, p-value = 0.91
## alternative hypothesis: two-sided
\# Do x and y come from the same distribution?
ks.test(norm_df$normal_example, norm_df$skewed_example)
##
##
   Exact two-sample Kolmogorov-Smirnov test
## data: norm_df$normal_example and norm_df$skewed_example
## D = 0.571, p-value = 1.3e-05
## alternative hypothesis: two-sided
```

Non-parametric Tests

Wilcoxon Rank Sum Test

We'll work on the ToothGrowth dataset for this exercise. The response is the length of odontoblasts (cells responsible for tooth growth) in 60 guinea pigs. Each animal received vitamin C by one of two delivery methods, orange juice (OJ) or ascorbic acid (VC). We'll compare the mean lengths between VC and OJ.

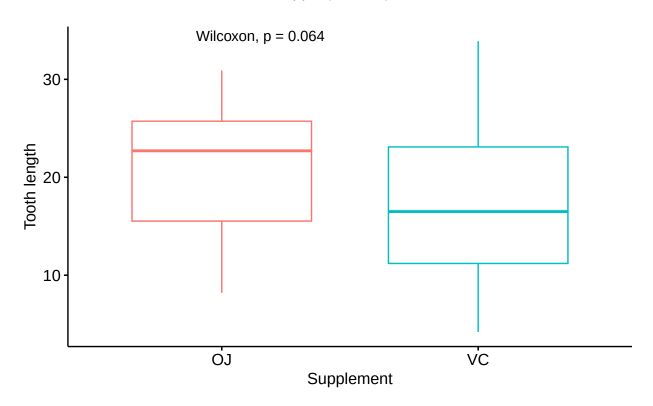
```
?ToothGrowth
head(ToothGrowth, 3)
##
      len supp dose
## 1
     4.2
            VC 0.5
## 2 11.5
            VC
               0.5
## 3 7.3
            VC
               0.5
# Hypothesis: mean tooth lengths of VC and OJ are different
summary(ToothGrowth$len[ToothGrowth$supp == "VC"])
##
      Min. 1st Qu.
                    Median
                               Mean 3rd Qu.
                                                Max.
##
       4.2
              11.2
                       16.5
                               17.0
                                       23.1
                                                33.9
summary(ToothGrowth$len[ToothGrowth$supp == "OJ"])
##
      Min. 1st Qu.
                    Median
                               Mean 3rd Qu.
                                                Max.
       8.2
              15.5
                       22.7
                                                30.9
##
                               20.7
                                       25.7
g <- ggboxplot(data = ToothGrowth,</pre>
               x = "supp", y = "len",
               color = "supp",
               xlab = "Supplement", ylab = "Tooth length")
g
```

supp 🖨 OJ 🖨 VC 30 Tooth length 20 10 ÖJ VC Supplement # Normal distribution? shapiro.test(ToothGrowth\$len[ToothGrowth\$supp == "VC"]) ## Shapiro-Wilk normality test ## ## ## data: ToothGrowth\$len[ToothGrowth\$supp == "VC"] ## W = 0.966, p-value = 0.43 shapiro.test(ToothGrowth\$len[ToothGrowth\$supp == "OJ"]) ## ## Shapiro-Wilk normality test ## ## data: ToothGrowth\$len[ToothGrowth\$supp == "OJ"] ## W = 0.918, p-value = 0.024 # Wilcox test (Mann-Whitney U test) wilcox.test(len~supp, data = ToothGrowth) ## Warning in wilcox.test.default(x = DATA[[1L]], y = DATA[[2L]], ...): cannot ## compute exact p-value with ties ## Wilcoxon rank sum test with continuity correction ##

data: len by supp ## W = 576, p-value = 0.064

```
## alternative hypothesis: true location shift is not equal to 0
g + stat_compare_means(method = "wilcox")
```





Kruskal-Wallis Rank Sum Test

We'll use the diet dataset which contains information on 78 people using one of three diets.

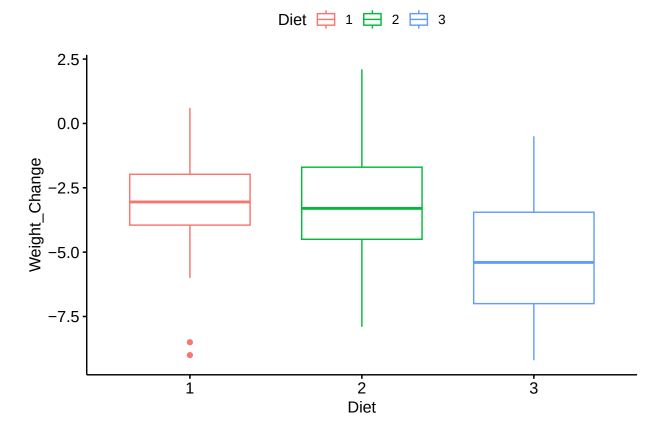
```
diet_df <- read.csv("../data/Diet_R.csv")
head(diet_df)</pre>
```

```
##
     Person gender Age Height pre.weight Diet weight6weeks
                                                           60.0
## 1
         25
                 NA
                     41
                            171
                                         60
                                                2
## 2
         26
                            174
                                        103
                                                2
                                                          103.0
                 NA
                     32
## 3
          1
                     22
                            159
                                         58
                                                1
                                                           54.2
## 4
           2
                  0
                     46
                            192
                                         60
                                                           54.0
                                                1
## 5
                     55
                            170
                                         64
                                                1
                                                           63.3
## 6
                  0
                     33
                            171
                                                           61.1
```

```
# turn categorical variables into factors
diet_df$Diet <- as.factor(diet_df$Diet)
diet_df$gender <- as.factor(diet_df$gender)

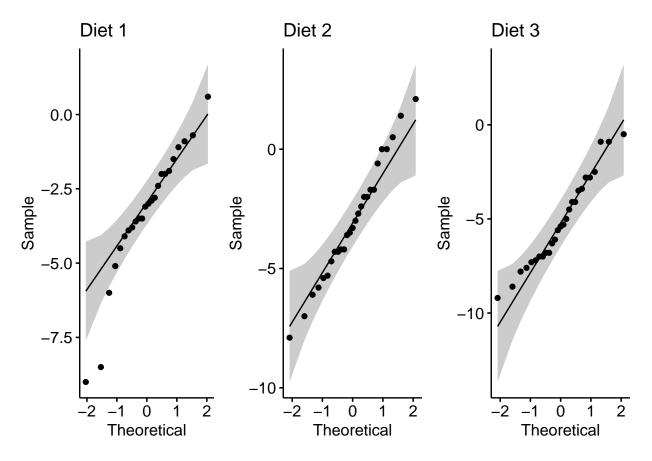
# create new variable
diet_df$Weight_Change <- diet_df$weight6weeks - diet_df$pre.weight</pre>
```

```
summary(diet_df)
                    gender
##
                                                              pre.weight
        Person
                                   Age
                                                 Height
                                                                            Diet
##
                       :43
                                     :16.0
                                                                            1:24
    Min.
          : 1.0
                             Min.
                                                    :141
                                                            Min. : 58.0
##
    1st Qu.:20.2
                   1
                       :33
                              1st Qu.:32.2
                                             1st Qu.:164
                                                            1st Qu.: 66.0
                                                                            2:27
##
    Median:39.5
                   NA's: 2
                             Median:39.0
                                             Median:170
                                                            Median : 72.0
                                                                            3:27
##
    Mean
           :39.5
                             Mean
                                     :39.2
                                             Mean
                                                    :171
                                                            Mean
                                                                   : 72.5
    3rd Qu.:58.8
                              3rd Qu.:46.8
                                             3rd Qu.:175
                                                            3rd Qu.: 78.0
##
##
    Max.
           :78.0
                             Max.
                                     :60.0
                                             Max.
                                                    :201
                                                            Max.
                                                                   :103.0
##
     weight6weeks
                    Weight Change
##
                    Min.
    Min.
           : 53.0
                           :-9.20
    1st Qu.: 61.9
                    1st Qu.:-5.55
   Median: 69.0
                    Median :-3.60
##
##
    Mean : 68.7
                    Mean :-3.84
    3rd Qu.: 73.8
                    3rd Qu.:-2.00
##
    Max.
           :103.0
                    Max.
                           : 2.10
g <- ggboxplot(diet_df, x = "Diet", y = "Weight_Change", color = "Diet")
g
```



We'll compare weight changes of the three diet groups. Let's check the normality of weight changes of the three groups:

- ## Warning: The following aesthetics were dropped during statistical transformation: sample
- ## i This can happen when ggplot fails to infer the correct grouping structure in
- ## the data.
- ## i Did you forget to specify a `group` aesthetic or to convert a numerical
- ## variable into a factor?
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- ## i This can happen when ggplot fails to infer the correct grouping structure in
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- ## i Did you forget to specify a `group` aesthetic or to convert a numerical
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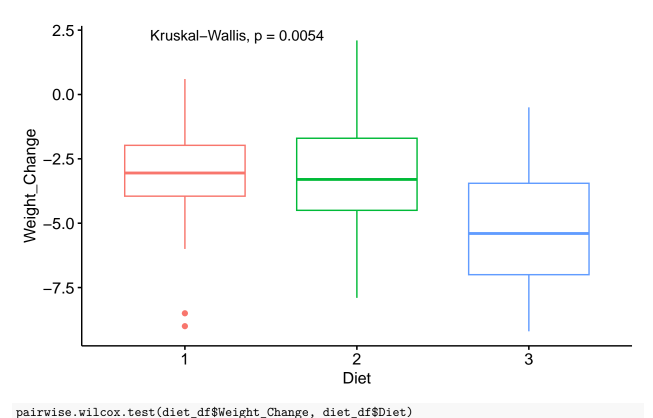


The normality assumption of ANOVA is not met, we'll use the Kruskal-Wallis test instead: kruskal.test(Weight_Change~Diet, data = diet_df)

```
##
## Kruskal-Wallis rank sum test
##
## data: Weight_Change by Diet
## Kruskal-Wallis chi-squared = 10.4, df = 2, p-value = 0.0054
```

g + stat_compare_means()

Diet 😑 1 😑 2 😑 3



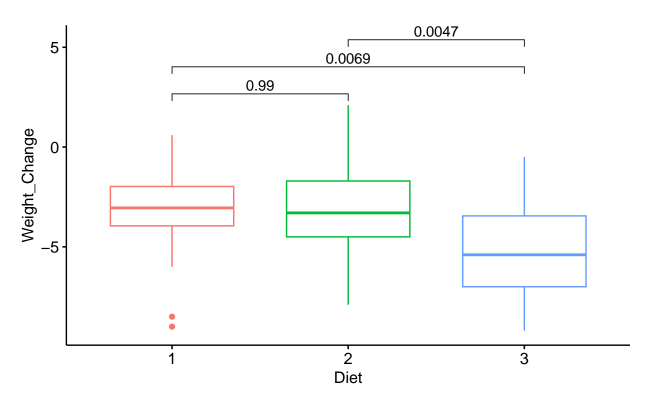
```
## Warning in wilcox.test.default(xi, xj, paired = paired, ...): cannot compute
## exact p-value with ties
## Warning in wilcox.test.default(xi, xj, paired = paired, ...): cannot compute
## exact p-value with ties
## Warning in wilcox.test.default(xi, xj, paired = paired, ...): cannot compute
## exact p-value with ties
##
  Pairwise comparisons using Wilcoxon rank sum test with continuity correction
##
##
## data: diet_df$Weight_Change and diet_df$Diet
##
##
     1
          2
## 2 0.99 -
## 3 0.01 0.01
## P value adjustment method: holm
g + stat_compare_means(comparisons = list(c(1, 2), c(1, 3), c(2, 3)), method = "wilcox")
## [1] FALSE
```

Warning in wilcox.test.default(c(-3.8, -6, -0.7000000000003, -2.9, -2.8, :

cannot compute exact p-value with ties

```
## Warning in wilcox.test.default(c(-3.8, -6, -0.700000000000003, -2.9, -2.8, :
## cannot compute exact p-value with ties
## Warning in wilcox.test.default(c(0, 0, 2.1, -2, -1.7, -4.3, -7,
## -0.600000000000001, : cannot compute exact p-value with ties
```





Multiple Testing Correction

```
?p.adjust
p_bonf <- p.adjust(p_plain, method = "bonferroni")</pre>
p_holm <- p.adjust(p_plain, method = "holm")</pre>
p_fdr <- p.adjust(p_plain, method = "fdr")</pre>
head(sort(p_bonf), 10)
## gene_2124 gene_896 gene_2600
                                    gene_766 gene_829 gene_2851 gene_703
## 8.4847e-09 4.6888e-06 2.5701e-05 4.8125e-05 6.9542e-05 1.2413e-04 1.7291e-04
## gene_2386 gene_2645 gene_2002
## 1.8074e-04 2.0762e-04 2.2060e-04
head(sort(p holm), 10)
## gene_2124 gene_896 gene_2600 gene_766 gene_829 gene_2851 gene_703
## 8.4847e-09 4.6873e-06 2.5684e-05 4.8078e-05 6.9451e-05 1.2393e-04 1.7257e-04
## gene_2386 gene_2645 gene_2002
## 1.8032e-04 2.0707e-04 2.1995e-04
head(sort(p_fdr), 10)
## gene_2124 gene_896 gene_2600 gene_766 gene_829 gene_2851 gene_703
## 8.4847e-09 2.3444e-06 8.5669e-06 1.2031e-05 1.3908e-05 2.0689e-05 2.2060e-05
## gene_2002 gene_2386 gene_2645
## 2.2060e-05 2.2060e-05 2.2060e-05
sum(p_bonf <= 0.05)
## [1] 103
sum(p_holm \le 0.05)
## [1] 103
sum(p_fdr <= 0.05) # less conservative</pre>
## [1] 695
```