Assignment 1 - Report

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```
install.packages("tinytex", repos = "https://cran.r-project.org")
##
## The downloaded binary packages are in
## /var/folders/lt/3tf47cnj2n5f1w0d_h6xmyvr0000gn/T//RtmpJS2HVF/downloaded_packages
```

Exercise 1

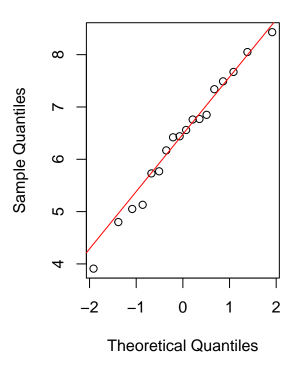
First we load and read the necessary data set

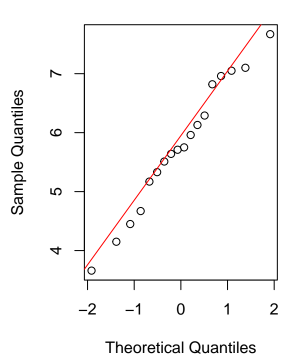
```
data = read.delim("cholesterol.txt", sep=' ')
```

a) Make some relevant plots of this data set, comment on normality. Investigate whether the columns Before and After8weeks are correlated. In order to investigate the normality of the data set, Q-Q plots are created below for both the Before and After8weeks columns. As in both plots the data points closely follow the diagonal red line, the data is approximating a normal distribution. While some minor deviations may be present in the tails, the overall pattern suggests that the normality assumption is reasonable.

Q-Q Plot for Before

Q-Q Plot for After 8 Weeks

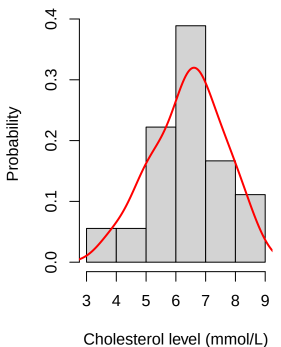


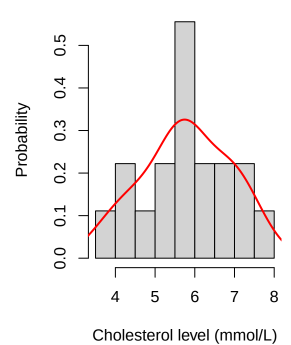


To further explore the normality assumption, histograms below were plotted for both 'Before' and 'After8weeks'. The histograms exhibit a roughly bell-shaped distribution, which supports the assumption of normality.

Distribution for Cholesterol level Before margarine

Distribution for Cholesterol level After margarine





However, to address normality more formally, a Shapiro-Wilk test is conducted, as this test is suitable to test on normality for small data sets. For the test the null hypothesis is as follows:

H0: The data is normally distributed.

The W-statistic measures how closely the data aligns with a normal distribution, ranging from 0 to 1, where values closer to 1 indicate a stronger likelihood of normality. Considering the results for *Before* and *After8weeks*, both W-values are close to 1. Additionally, with a 95% confidence level, both p-values exceed 0.05, meaning that we fail to reject H0. These findings provide strong evidence that the data in both columns can be considered to be normally distributed.

```
shapiro.test(data$Before)

##

## Shapiro-Wilk normality test

##

## data: data$Before

## W = 0.9819, p-value = 0.9675

shapiro.test(data$After8weeks)

##

## Shapiro-Wilk normality test

##

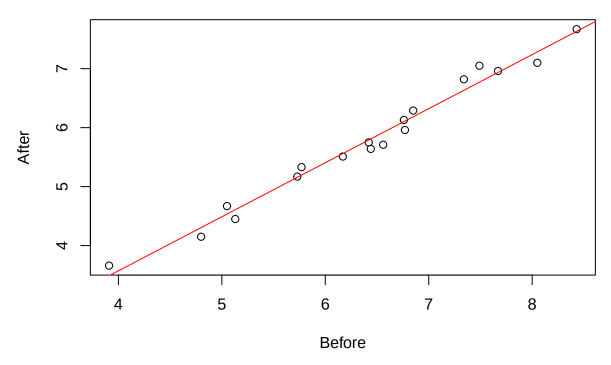
## data: data$After8weeks

##

## 0.97733, p-value = 0.9183
```

In order to investigate the relationship between the columns of *Before* and *After8weeks* a scatter plot is created below. The scatter plot demonstrates a strong positive correlation between 'Before' and 'After8weeks' cholesterol levels. The data points align closely with the red regression line, suggesting that individuals with higher cholesterol levels before the diet intervention also tend to have higher cholesterol levels after 8 weeks. This indicates that while cholesterol levels may have decreased, there remains a strong relationship between pre- and post-diet measurements.

Regression for Before and After8weeks



To quantify this correlation, the Pearson's correlation coefficient is calculated below. A high Pearson correlation (close to 1) indicates a strong positive relationship between the two columns. The correlation coefficient exhibits a value of approximately 0.99, confirming the strong positive relationship. Additionally, the p-value is smaller than 0.05, indicating that the correlation is statistically significant.

```
cor.test(data$Before, data$After8weeks, method = "pearson")
```

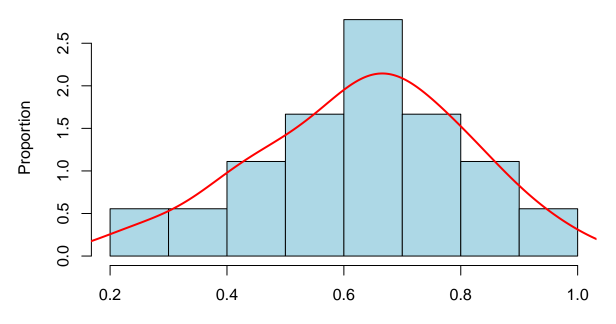
```
##
## Pearson's product-moment correlation
##
## data: data$Before and data$After8weeks
## t = 29.428, df = 16, p-value = 2.321e-15
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.9751289 0.9966788
## sample estimates:
## cor
## 0.9908885
```

b) Apply a couple of relevant tests (at least two tests, see Lectures 2–3) to verify whether the diet with low fat margarine has an effect (argue whether the data are paired or not). Is a permutation test applicable? Is the Mann-Whitney test applicable? As the cholesterol data was measured on the same population at different times, we consider the data to be paired. In this case it is possible to conduct a T-test for paired samples. However, in order to This test assumes that the mean difference of the two populations is normally distributed, thus, a Shapiro-Wilk test is conducted first to investigate the distribution.

```
difference = data$Before - data$After8weeks
shapiro.test(difference)
##
##
   Shapiro-Wilk normality test
##
## data: difference
## W = 0.98501, p-value = 0.9869
Explain that the data is paired:
We are doing a T-test
The null hypothesis is as follows:
H0: The margarine diet has no effect, i.e. the mean cholesterol levels Before and After8weeks are
the same.
# HO: The margarine diet has no effect, i.e. the mean cholesterol levels Before and After 8 we
# H1: The margarine diet reduces cholesterol levels --> mean_before > mean_after
# Paired t-test
# Assumption: the differences between Before and After should be normally distributed
# Outcome: if p < 0.05, reject HO
t.test(data$Before, data$After8weeks, paired = TRUE, alternative = "greater")
##
## Paired t-test
##
## data: data$Before and data$After8weeks
## t = 14.946, df = 17, p-value = 1.639e-11
## alternative hypothesis: true mean difference is greater than 0
## 95 percent confidence interval:
## 0.5556906
## sample estimates:
## mean difference
         0.6288889
##
# Visualizing distribution of the differences
difference = data$Before - data$After8weeks
shapiro.test(difference)
##
   Shapiro-Wilk normality test
##
## data: difference
## W = 0.98501, p-value = 0.9869
hist(difference, probability = TRUE,
     main = 'Data distribution of differences between Before and After8weeks',
     col = 'lightblue',
```

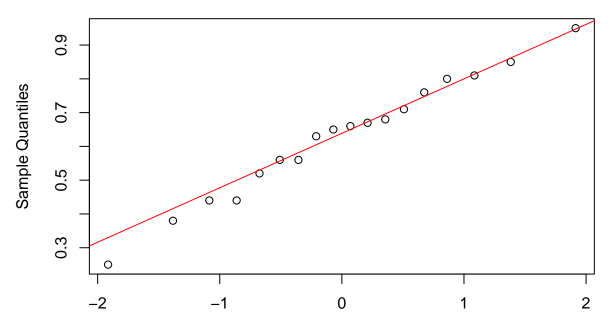
```
xlab = 'Difference (Before - After8weeks)',
ylab = 'Proportion')
lines(density(difference), col = 'red', lwd = 2)
```

Data distribution of differences between Before and After8weeks



Difference (Before – After8weeks)

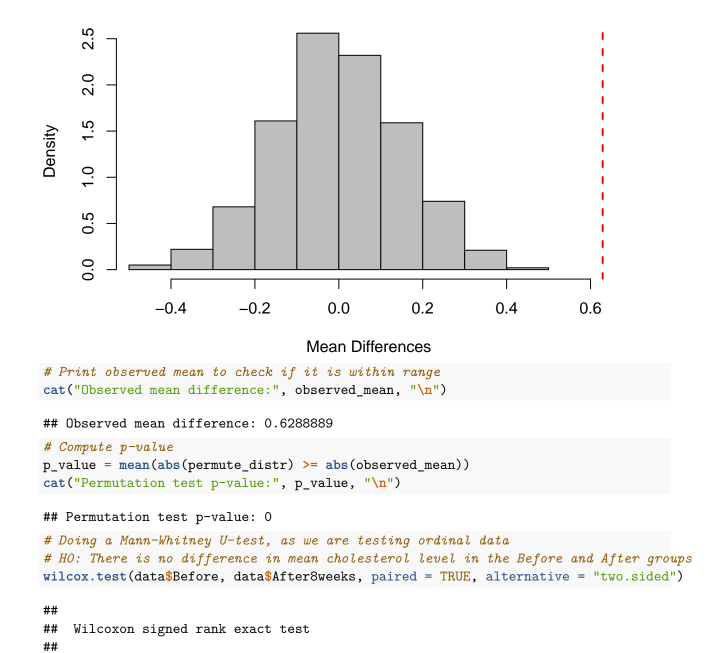
QQ-plot for differences



Theoretical Quantiles

```
# Doing permutation test:
# HO: there is no difference between the Before and After8weeks groups
diff = data$Before - data$After8weeks
n_permutations = 1000
observed_mean = mean(diff)
permute_test = function(diff) {
 permuted_diff <- diff * sample(c(-1, 1), length(diff), replace = TRUE) # Randomly flip sign</pre>
 return(mean(permuted_diff))
}
set.seed(42)
permute_distr = replicate(n_permutations, permute_test(diff))
# Histogram of the permutation distribution
hist(permute_distr, probability = TRUE, col = "gray",
     main = "Permutation Test Distribution", xlab = "Mean Differences",
     xlim = range(c(permute_distr, observed_mean))) # Adjust x-axis limits
# Add a red line at observed mean difference
abline(v = observed_mean, col = "red", lwd = 2, lty = 2)
```

Permutation Test Distribution



```
## alternative hypothesis: true location shift is not equal to 0
# H1: cholesterol levels are lower after 8 weeks
wilcox.test(data$Before, data$After8weeks, paired = TRUE, alternative = "greater")
##
## Wilcoxon signed rank exact test
##
```

data: data\$Before and data\$After8weeks

data: data\$Before and data\$After8weeks

V = 171, p-value = 7.629e-06

```
## V = 171, p-value = 3.815e-06
## alternative hypothesis: true location shift is greater than 0
```

```
# calculating 97% CI for mu using t-score
n = length(data$After8weeks)
sample_mean = mean(data$After8weeks)
sample_sd = sd(data$After8weeks)
critical_value = qt(1-0.015, df=17)
standard_error = sample_sd / sqrt(n)

left_bound = sample_mean - critical_value * standard_error
right_bound = sample_mean + critical_value * standard_error

cat("97% Confidence Interval for mu: [", left_bound, ",", right_bound, "]\n")
```

c) Let X1,...,X18 be the column After8weeks. Assume $X1,...,X18 \sim N(mu, sigma^2)$ (irrespective of your conclusion in a)) with unknown and 2 . Construct a 97%-CI for based on normality. Next, construct a bootstrap 97%-CI for and compare it to the above CI.

```
## 97% Confidence Interval for mu: [ 5.16385 , 6.393928 ]
```

```
# calculating 97% CI for mu with bootstrapping
bootstrap_ci = function(x, conf_level = 0.97, B = 10000) {
    alpha = 1 - conf_level
    Bstats = lapply(1:B, FUN = function(i) {
        boot_sample = sample(x, size = length(x), replace = TRUE)
        mean(boot_sample)
    })
    Bstats = unlist(Bstats)
    quantile(Bstats, prob = c(alpha/2, 1-alpha/2))
}
set.seed(42)
bootstrap_ci(data$After8weeks)
```

```
## 1.5% 98.5%
## 5.229992 6.320008
```

d) Using a bootstrap test with test statistic T=max(X1,...,X18), determine those [3,12] (if there are any) for which H0:X1,...,X18 Unif[3,] is not rejected. Can the Kolmogorov-Smirnov test be also applied for this question? If yes, apply it; if not, explain why not.

```
median(data$After8weeks)
```

e) Using an appropriate test, verify whether the median cholesterol level after 8 weeks of low fat diet is less than 6. Next, design and perform a test to check whether the fraction of the cholesterol levels after 8 weeks of low fat diet less than 4.5 is at most 25%.

```
## [1] 5.73
wilcox.test(data$After8weeks, mu = 6, alternative = "less")
## Warning in wilcox.test.default(data$After8weeks, mu = 6, alternative = "less"):
## cannot compute exact p-value with ties
##
##
   Wilcoxon signed rank test with continuity correction
##
## data: data$After8weeks
## V = 67.5, p-value = 0.223
## alternative hypothesis: true location is less than 6
# Count how many values in After8weeks are less than 4.5
count_below_4.5 = sum(data$After8weeks < 4.5)</pre>
percentage_below_4.5 = (count_below_4.5 / length(data$After8weeks)) * 100
cat("Percentage of cholesterol levels below 4.5:", percentage_below_4.5, "%\n")
## Percentage of cholesterol levels below 4.5: 16.66667 %
# HO: The fraction of cholesterol levels below 4.5 is at most 25%
# H1: The fraction is greater than 25%
# if the p-value is small (<0.05), we reject HO and conclude that the fraction is significantly
binom.test(count_below_4.5, length(data$After8weeks), p = 0.25, alternative = "greater")
##
##
   Exact binomial test
##
## data: count_below_4.5 and length(data$After8weeks)
## number of successes = 3, number of trials = 18, p-value = 0.8647
## alternative hypothesis: true probability of success is greater than 0.25
## 95 percent confidence interval:
## 0.04702488 1.00000000
## sample estimates:
## probability of success
##
                0.1666667
```

Exercise 2: Crops

Section a

We want to investigate whether two factors County and Related (and possibly their interaction) influence the crops by performing relevant ANOVA model(s), without taking Size into account. So we create and test 3 separate Null Hypotheses with a two-way ANOVA and a one-way ANOVA on the additive model: H_(01): no main effect of factor County, H_(02): no main effect of factor Related and H_(03): no interactions between factors County and Related

```
## Analysis of Variance Table
##
## Response: Crops
##
                  Df
                        Sum Sq Mean Sq F value Pr(>F)
                   2
## County
                       8841441 4420721 0.7644 0.4766
## Related
                                        0.4113 0.5274
                   1
                       2378957 2378957
## County:Related 2
                       1497573
                                748786
                                         0.1295 0.8792
## Residuals
                  24 138805865 5783578
```

From this table we can see the following:

For County: The p-value (0.476) is greater than 0.05, meaning we fail to reject the null hypothesis H_(01), suggesting that there is no significant effect of the County on the Crops variable.

For Related: The p-value (0.527) is also greater than 0.05, meaning we fail to reject the null hypothesis $H_{-}(02)$, which suggests that there is no significant effect of whether the landlord and tenant are related on the Crops variable.

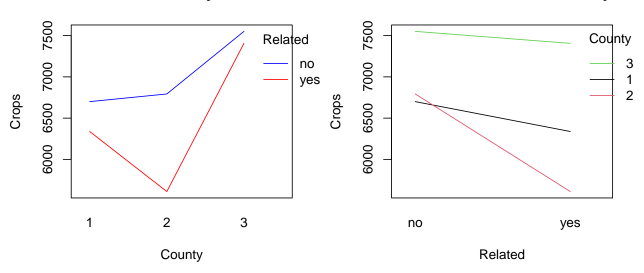
For both: The p-value (0.879) is much greater than 0.05, meaning we fail to reject the null hypothesis $H_{-}(03)$, implying there is no significant interaction between County and Related on the Crops variable.

```
##
## Call:
## lm(formula = Crops ~ County * Related, data = crops_data)
##
## Residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
## -3120.4 -1744.7 -176.9
                            2064.2
##
## Coefficients:
                      Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                        6700.0
                                    1075.5
                                             6.230 1.94e-06 ***
## County2
                                                      0.952
                          93.0
                                    1521.0
                                             0.061
## County3
                         851.2
                                    1521.0
                                             0.560
                                                      0.581
## Relatedyes
                        -362.0
                                    1521.0
                                           -0.238
                                                      0.814
## County2:Relatedyes
                        -820.6
                                    2151.0
                                           -0.381
                                                      0.706
## County3:Relatedyes
                         217.0
                                    2151.0
                                             0.101
                                                      0.920
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2405 on 24 degrees of freedom
## Multiple R-squared: 0.08393,
                                    Adjusted R-squared:
## F-statistic: 0.4398 on 5 and 24 DF, p-value: 0.8163
```

The above model summary table aligns with the ANOVA p-values as both show that none of the predictors (County, Related, or their interaction) are significant in either table.

Interaction: County & Related

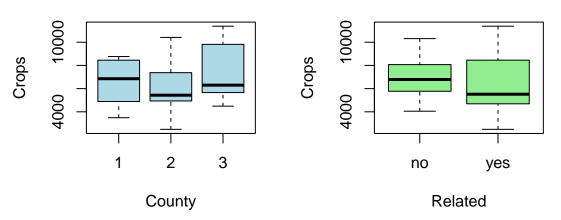
Interaction: Related & County



In the above interaction plots the lines seem parallel, therefore interaction seems to not be present, verifying the two-way anova results.

Effect of County on Crops

Effect of Related on Crops



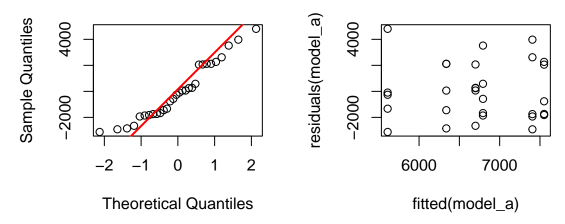
ANOVA shows no significant effect of County or Related on crop yield. High p-values suggest no strong differences, consistent with the boxplot, where distributions overlap, medians are close, and no outliers appear.

Finally, we have to check the model assumptions

[1] "These are the p-values of the Shapiro-Wilk test for model_a: p = 0.099, W = 0.941"

The Q-Q plot shows deviations from normality, particularly in the tails, but the overall trend follows the theoretical quantiles. The residuals vs. fitted plot suggests no strong patterns, indicating an approximately random distribution of residuals. The Shapiro-Wilk test (W = 0.941, p = 0.099) fails to reject the null hypothesis of normality at the 0.05 level. Given the small sample size (n = 30), results should be interpreted with caution, as minor departures from normality can impact statistical inference.

Normal Q-Q Plot



To estimate crop yields for County 3 when the landlord and tenant are unrelated, we use the emmeans function to calculate the adjusted mean yield. This estimation is based on model_a, which incorporates the County-Size interaction. The emmeans function provides the estimated marginal means, accounting for the effects of County and Related while adjusting for interactions.

Estimated crops for County 3 (Landlord and Tenant NOT related): 7551.2

Section b

We define 3 different models:

 Model_county_size: This model examines how crop yields are influenced by County, Related, and Size, with an additional focus on the interaction between County and Size. It does not include an interaction term for Related.

```
## Analysis of Variance Table
##
## Response: Crops
##
                \mathsf{Df}
                                          F value
                                                     Pr(>F)
                       Sum Sq
                                Mean Sq
                 1 119569344 119569344 135.6241 4.01e-11 ***
## Size
                 2
## County
                       767179
                                  383589
                                           0.4351
                                                    0.65242
## Related
                 1
                      1381334
                                 1381334
                                            1.5668
                                                    0.22325
                 2
## Size:County
                     9528654
                                 4764327
                                           5.4040
                                                    0.01192 *
## Residuals
                23
                    20277325
                                  881623
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

2. Model_related_size: This model evaluates how County, Related, and Size affect crop yields, and it adds an interaction term for Related and Size to test if the effect of Size on crop yields depends on whether the landlord and tenant are related.

```
## Analysis of Variance Table
##
## Response: Crops
##
                 Df
                                                      Pr(>F)
                       Sum Sq
                                 Mean Sq
                                          F value
## Size
                  1 119569344 119569344 100.8587 4.521e-10 ***
## Related
                  1
                      1380585
                                 1380585
                                            1.1645
                                                      0.2913
```

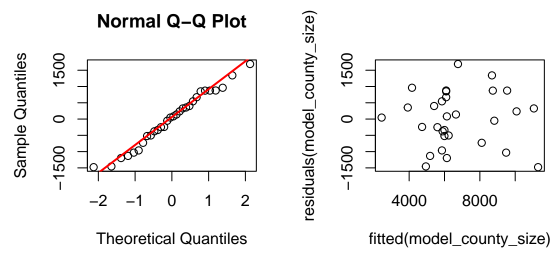
```
767927
                                            0.3239
                                                       0.7264
## County
                  2
                                  383964
## Size:Related
                  1
                      1353666
                                 1353666
                                            1.1418
                                                       0.2959
## Residuals
                 24
                     28452313
                                 1185513
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

3. Model_additive: This model assumes that the effect of each factor on crop yield is independent of the others. It does not test for any interaction effects, only the individual contributions of County, Related, and Size to crop yields.

```
## Analysis of Variance Table
##
## Response: Crops
                                                   Pr(>F)
##
              Df
                    Sum Sq
                             Mean Sq
                                       F value
## Size
                119569344 119569344 100.2897 3.114e-10 ***
## County
              2
                    767179
                               383589
                                        0.3217
                                                   0.7278
                                        1.1586
                                                   0.2920
## Related
              1
                   1381334
                              1381334
## Residuals 25
                  29805979
                              1192239
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

We have tested interaction models as well as purely additive. The interaction Size-Related and the individual effect of County and Related are insignificant(all 3 have p-values>0.5). Therefore, the best model is model_county_size, since it shows the significance of Size and of the interaction Size-County.

Finally, we can check this model's assumptions.



[1] "These are the p-values of the Shapiro-Wilk test for model_county_size: p = 0.733, W = 0.733 The Shapiro-Wilk test for model_county_size residuals (p = 0.733) suggests no significant devi-

ation from normality, supported by the QQ-plot's linear pattern.

Section c

##

Call:

```
## lm(formula = Crops ~ Size * County + Related, data = crops_data)
##
## Residuals:
##
        Min
                   1Q
                        Median
                                     3Q
                                              Max
  -1477.64
             -517.10
                         63.89
                                 639.62
                                         1690.29
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 2461.014
                              929.764
                                         2.647
                                                0.01441 *
## Size
                    22.704
                                4.765
                                        4.764 8.38e-05 ***
## County2
                -4214.050
                             1447.242
                                       -2.912
                                                0.00785 **
## County3
                -1284.813
                             1302.578
                                       -0.986
                                                0.33422
## Relatedyes
                  -239.099
                              347.916
                                       -0.687
                                                0.49881
## Size:County2
                    26.590
                                8.091
                                         3.286
                                                0.00323 **
## Size:County3
                    8.916
                                6.398
                                         1.394
                                                0.17676
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 938.9 on 23 degrees of freedom
## Multiple R-squared: 0.8662, Adjusted R-squared:
## F-statistic: 24.81 on 6 and 23 DF, p-value: 5.851e-09
```

The coefficient for Size is 22.704 (p<0.001), meaning that in County 1 (the reference level), each unit increase in Size leads to an expected 22.7-unit increase in Crops. This effect is statistically significant, confirming a strong positive influence. County 2 has a negative coefficient of -4214.050 (p<0.01), meaning crop yields there are 4214 units lower than in County 1 for the same Size. County 3's coefficient (-1284.813, p=0.334) is not statistically significant, so we cannot conclude a strong difference from County 1. Related has a coefficient of -239.099 (p=0.499), indicating no significant impact on Crops. The Size:County2 interaction is 26.590 (p=0.003), meaning that the effect of Size on Crops in County 2 is stronger than in County 1, with a total increase of 49.3 units per Size unit. The Size:County3 interaction (8.916, p=0.176) is not statistically significant, so we cannot confidently conclude a difference from County 1. The model explains 86.6% of the variation in Crops (R²=0.866), indicating strong explanatory power.

confint(model county size)

```
##
                        2.5 %
                                    97.5 %
## (Intercept)
                   537.651576
                               4384.37722
## Size
                    12.845887
                                  32.56210
## County2
                 -7207.896850 -1220.20244
## County3
                 -3979.399914
                               1409.77389
                  -958.817141
## Relatedyes
                                 480.61942
## Size:County2
                     9.852682
                                  43.32644
## Size:County3
                    -4.319019
                                  22.15129
```

The confidence intervals confirm that Size has a strong, statistically significant positive effect, and that County 2 and the Size:County2 interaction also have significant effects. In contrast, the confidence intervals for County 3, Related, and Size:County3 include zero, meaning there is no strong evidence that these factors significantly influence Crops.

Section d

```
## County Related Size emmean SE df lower.CL upper.CL
## 2 yes 165 6141 345 23 5428 6855
##
## Confidence level used: 0.95
The predicted yield crops for a farm from County 2 of size 165, with related landlord and tenant is 6141, with a 95% CI: (5428, 6855)
## Prediction Variance: 118896.9
## Residual Variance (sigma^2): 881622.8
## Total Variance: 1000520
```

The fact that the prediction variance is much smaller than the residual variance suggests that most of the uncertainty is due to the residual variation (random noise or factors not captured by the model), rather than the instability of the model's coefficient estimates.

Exercise 3

Here, we will explore the effect of different additives on the yield of peas with a primary focus on nitrogen (N). Other additives are potassium (K) and phosphorus (P). The data is obtained through the MASS package library (npk) and represents 6 blocks of soil, each containing 4 plots. Each *block* contains exactly two of each additive and no plot can have two of the same additive.

We start by loading the necessary npk dataset from the MASS package.

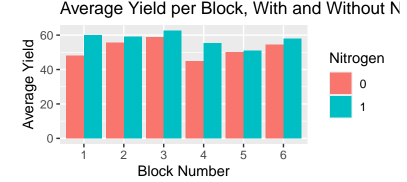
Section a

To optimize possible future instances of a random plot distribution process we create a data frame similar to that of the npk data such that it may be utilized similarly.

```
random_distributed[sample(idx, 2), "K"] <- 1
}</pre>
```

Section b

From the npk data a bar graph is generated to report the average yield per block in the presence (1) and absence (0) of N.



From the graph we can note that there seems to be a correlation between a higher yield and the presence of the additive N in the plot soil. We will further investigate this hypothesis with different tests such as a full two-way ANOVA. The block factor is not of interest but is applied to group plot tests and may introduce variability within the samples.

Section c

A full two-way ANOVA is conducted with the response variable *yield* and the two factors *block* and N. We start with a main effects model.

```
##
               Df Sum Sq Mean Sq F value Pr(>F)
                                     3.395 0.0262 *
## block
                 5
                    343.3
                            68.66
## N
                 1
                    189.3
                           189.28
                                     9.360 0.0071 **
                17
                    343.8
                            20.22
## Residuals
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

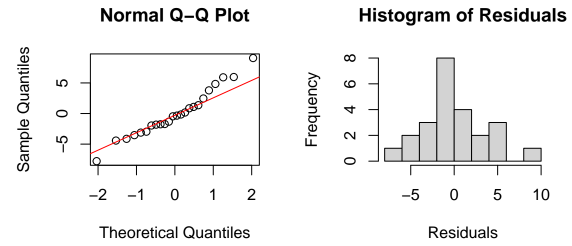
We then compute an interaction model.

```
##
               Df Sum Sq Mean Sq F value Pr(>F)
## block
                    343.3
                            68.66
                                     3.359 0.0397 *
## N
                 1
                    189.3
                           189.28
                                     9.261 0.0102 *
                            19.70
                                     0.964 0.4769
## block:N
                 5
                     98.5
## Residuals
                12
                    245.3
                            20.44
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

With a p-value of 0.0071 and 0.0262 respectively in the main effects model, the ANOVA signifies that both the N additive and block factor have statistically significant effects on the yield of peas with nitrogen having a larger effect compared to the block. However, we might say that the block factor merely introduces variability in the samples and do not represent fixed system effects. Including the block factor is, however, important to understand the amount of variability it may cause. In

the interaction model both factors again have a p-value below 0.05. Additionally, we notice that there is no statistically significant effect on the *yield* from an interaction between N and *yield*.

To solidify our findings we perform an analysis on the residuals of the main effects model with a normality test and a plot of the distribution with a histogram.



We note that there is a slight deviation at the tail of the normality test which we investigate this further with a Shapiro Wilk test.

```
##
## Shapiro-Wilk normality test
##
## data: residuals(two_way_anova_main)
## W = 0.96937, p-value = 0.6514
```

From the Shapiro-Wilk test we receive a p-value greater than 0.05, combined with the Q-Q and histogram it is likely normally distributed.

Tests indicate a normal distribution of the residuals and the data is continuous. We can therefore say that a non-parametric designed test such as a Friedman test is not applicable. Additionally, We have limited in-block variation due to a randomized block design with continuous data which is more applicable to the ANOVA model design.

Section d

To investigate the effects of other variables and any other possible interactions we explore more models. We first test for the presence of additional main effects from all additives and the block factor.

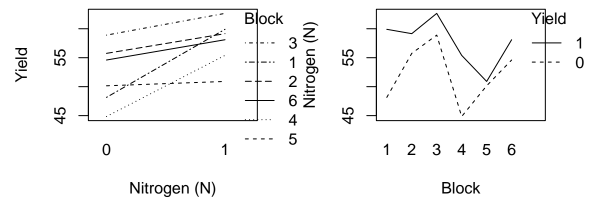
```
##
               Df Sum Sq Mean Sq F value Pr(>F)
## N
                    189.3
                           189.28
                                   11.821 0.00366 **
                 1
                      8.4
## P
                             8.40
                                     0.525 0.47999
## K
                 1
                     95.2
                            95.20
                                     5.946 0.02767 *
## block
                 5
                    343.3
                            68.66
                                     4.288 0.01272 *
## Residuals
                    240.2
                15
                            16.01
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

We now learn that potassium has a statistically significant effect on the yield with a p-value of 0.02767. From this main effects model we therefore learn that only P does not contribute significantly, we will therefore omit P in subsequent models.

Next, we explore the interaction between potassium and the other remaining main effect variables N and block.

```
##
               Df Sum Sq Mean Sq F value Pr(>F)
## N
                    189.3
                           189.28
                                   13.037 0.00476 **
## K
                     95.2
                            95.20
                                     6.557 0.02834 *
                 1
## block
                 5
                    343.3
                            68.66
                                     4.729 0.01777 *
                     33.1
                            33.14
## N:K
                 1
                                     2.282 0.16179
                 5
                     70.3
                                     0.968 0.48123
## K:block
                            14.05
## Residuals
                    145.2
                            14.52
                10
## ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

From the data we learn that there is no significant interaction between the different factors. We further explore this with an interaction plot.



From this plot we only observe a minor possible interaction between block and N. However, due to the overall linearity we suspect the interaction to be mostly negligible.

We will therefore approach a main effects model where we leave out phosphor to omit arbitrary data and prevent overfitting. The main effect model will include factors N, K and block

```
# NOTE: P could be omitted due to statistical insignificance
model_3 <- aov(yield ~ N + K + block, data = npk)
summary(model_3)
## Df Sum Sq Mean Sq F value Pr(>F)
```

```
Df Sum Sq Mean Sq F value Pr(>F)
## N
                   189.3
                           189.28
                                  12.183 0.00302 **
## K
                1
                     95.2
                            95.20
                                    6.128 0.02487 *
## block
                5
                   343.3
                            68.66
                                    4.419 0.01017 *
                   248.6
                            15.54
## Residuals
               16
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Section e

To investigate the influence of the factors on the *yield*, we will perform a Tukey's Honest Significant Difference (HSD) test.

```
# Apply TukeyHSD
tukey_results <- TukeyHSD(model_3)</pre>
tukey_results
     Tukey multiple comparisons of means
##
##
       95% family-wise confidence level
##
## Fit: aov(formula = yield ~ N + K + block, data = npk)
##
## $N
##
           diff
                      lwr
                               upr
                                       p adj
## 1-0 5.616667 2.205368 9.027965 0.0030242
##
## $K
##
            diff
                        lwr
                                   upr
                                           p adj
## 1-0 -3.983333 -7.394632 -0.5720346 0.0248744
##
## $block
##
          diff
                       lwr
                                 upr
                                         p adj
                -5.555691 12.405691 0.8168210
## 2-1
         3.425
## 3-1
         6.750
                -2.230691 15.730691 0.2062520
## 4-1
        -3.900 -12.880691
                            5.080691 0.7270343
## 5-1
        -3.500 -12.480691 5.480691 0.8035575
## 6-1
         2.325
                -6.655691 11.305691 0.9565038
## 3-2
         3.325
                -5.655691 12.305691 0.8338711
## 4-2
        -7.325 -16.305691
                            1.655691 0.1465878
## 5-2
        -6.925 -15.905691
                            2.055691 0.1863101
        -1.100 -10.080691
## 6-2
                            7.880691 0.9985112
## 4-3 -10.650 -19.630691 -1.669309 0.0156223
## 5-3 -10.250 -19.230691 -1.269309 0.0207261
        -4.425 -13.405691
## 6-3
                            4.555691 0.6173347
## 5-4
         0.400
                -8.580691
                            9.380691 0.9999896
## 6-4
         6.225
                -2.755691 15.205691 0.2761166
## 6-5
         5.825
                -3.155691 14.805691 0.3396777
```

From the TukeyHSD results we see that K has a significant negative impact on yield. Therefore we can say that the best combination for model 3 (where we do not consider P) is N=1 and K=0. Additionally, the block factor introduces variability with block 3 having the overall biggest positive impact and block 1 a relatively lower yield.

Section f

Finally, we create a mixed effects analysis for our chosen main effect model with factors N, K and block where we model block as a random effect.

```
## Linear mixed model fit by REML ['lmerMod']
## Formula: yield ~ N + K + (1 | block)
##
      Data: npk
##
## REML criterion at convergence: 131.4
##
## Scaled residuals:
##
        Min
                   10
                        Median
                                     3Q
                                              Max
## -1.67459 -0.51552 0.09225
                               0.51006
                                         1.52224
##
## Random effects:
    Groups
##
             Name
                          Variance Std.Dev.
    block
             (Intercept) 13.28
                                   3.644
##
                          15.54
##
    Residual
                                   3.942
##
  Number of obs: 24, groups: block, 6
##
## Fixed effects:
               Estimate Std. Error t value
##
                 54.058
                              2.039
                                     26.519
## (Intercept)
## N1
                  5.617
                              1.609
                                       3.490
## K1
                  -3.983
                              1.609
                                     -2.475
##
## Correlation of Fixed Effects:
##
      (Intr) N1
## N1 -0.395
## K1 -0.395
             0.000
```

Similarly to the additive model, we can conclude that N has a significant (5.617) positive effect on the yield with a p-value of 0.00302, whereas K has a statistically significant negative effect on the yield (-3.983) with a p-value of 0.02487. This analysis shows that blocks have significant variability in data, with a random effects variance of 13.16. This mixed effects model further solidifies the stance that N has a significant impact on yield. We further note that the mixed effects model may provide a better representation, since blocks introduce random variance and not fixed systemic effects such as N, P and K.