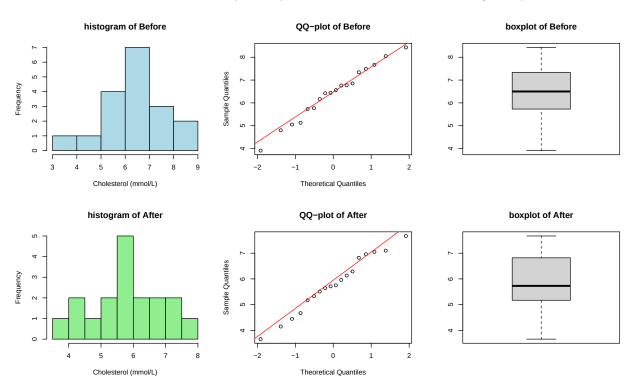
Group11

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Exercise 1

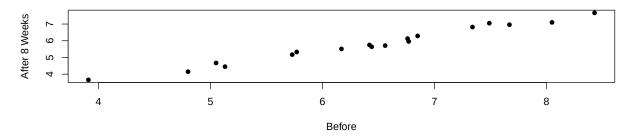
(a) We draw several figures to check the characteristics of the dataset. According to the Q-Q plots and from the shape of histogram, we can assume that the data Before and After 8 weeks both follow a normal distribution, since the points are approximately on a straight line, although the sample size is relatively small. Besides, we also check the scale, symmetry and outliers of the dataset through boxplot.



Then we also plot the scatter figures, we can see that there is approximately linear relation between these two values.

```
plot(data$Before, data$After8weeks,
    main = "Scatter Plot of Before vs. After 8 Weeks",
    xlab = "Before", ylab = "After 8 Weeks",
    col = "black", pch = 16)
```

Scatter Plot of Before vs. After 8 Weeks



The correlation of the two columns can also be calculated by Pearson as normality is assumed. The result is that the value of correlation is 0.991, which can infer that there is significant correlation.

(b) First, the data are paired as it is an experiment with two numerical outcomes per experimental unit. Therefore, the permutation test is applicable since permutation test is suitable for paired samples. However, Mann-Whitney test is not applicable, because it is utilized to compare two independent samples.

We already know that the data stems from a approximately normal distribution, so we can conduct **paired t-test** as follows:

```
t.test(data$Before,data$After8weeks, paired=TRUE)
```

```
##
## Paired t-test
##
## data: data$Before and data$After8weeks
## t = 15, df = 17, p-value = 3e-11
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.540 0.718
## sample estimates:
## mean difference
## 0.629
```

The p-value is 3e-11, so we can know taht H_0 can be rejected. H_0 means no significant difference between Before and After8weeks. In this case, there is significant difference between Before and After8weeks.

Then, as the sample size is 18, relatively small, we prefer non-Parametric tests. we can use **permutation test** as follows: The test statistic we choose is mean difference.

```
mean_difference = function(x,y) {mean(x-y)}
original_diff <- mean(data$Before - data$After8weeks)
B=1000; tstar=numeric(B)
for (i in 1:B) {
   md_star=t(apply(cbind(data$Before,data$After8weeks),1,sample))
   tstar[i]=mean_difference(data$Before,data$After8weeks)
}
myt = mean_difference(data$Before,data$After8weeks);
pl=sum(tstar<myt)/B;pr=sum(tstar>myt)/B;p=2*min(pl,pr); p
```

[1] 0

In the permutation test, we can reject H_0 and conclude that the diet with low fat margarine indeed has a significant effect.

(c) First, we construct the a 97%-CI for μ based on normality. As $X_1, \ldots, X_{18} \sim N(\mu, \sigma^2)$, we can calculate the t-confidence interval of level $1 - \alpha$ for μ . We use mean instead of median as the estimator because the outlier do not exists from conclusion in (a).

```
x = data$After8weeks;n = length(x);x_mean = mean(x);x_sd = sd(x);
alpha = 0.03;t = qt(1-alpha/2, df = n-1)
CI_normal = c(
    x_mean - t * (x_sd / sqrt(n)),
    x_mean + t * (x_sd / sqrt(n)))
cat("97% CI based on normality: [", round(CI_normal[1], 3),
    ",", round(CI_normal[2], 3), "]\n")
```

```
## 97% CI based on normality: [ 5.16 , 6.39 ]
```

Bootstrap CI is an alternative to determine CI's for non-normal sample, we implement bootstrap CI as follows:

```
B = 1000;x_mean = mean(data$After8weeks);Tstar = numeric(B)
for(i in 1:B){
    Xstar=sample(data$After8weeks,replace=TRUE)
    Tstar[i]=mean(Xstar)
}
Tstar15 = quantile(Tstar, 0.015);Tstar985 = quantile(Tstar, 0.985)
CI_bootstrap = c(2*x_mean-Tstar985,2*x_mean-Tstar15)
cat("Bootstrap 97% CI: [", round(CI_bootstrap[1], 3),
    ",", round(CI_bootstrap[2], 3), "]\n")
```

```
## Bootstrap 97% CI: [ 5.27 , 6.34 ]
```

As bootstrap CI will always yield a different interval when repeating, we run the chunk several time. And the conclusion is that the difference between the two confidence intervals is slight. From our perspective, this is be because the column after8weeks data steams form a normal distribution.

(d) We use the maximum of the sample as the test statistic and set B as 1000 to run the bootstrap test. We repeated the test a few times to see if the result is stable.

```
n=length(data$After8weeks)
t=max(data$After8weeks)
B=1000;tstar=numeric(B)
theta_values = seq(3, 12, by = 0.1)
p_values = sapply(theta_values, function(theta) {
   tstar = numeric(B)
   for (i in 1:B){
        xstar = runif(n, min = 3, max = theta)
        tstar[i] = max(xstar)
   }
   pl = sum(tstar < t) / B;pr = sum(tstar > t) / B
   p = 2 * min(pl, pr)
   return(p)
})
theta_valid = theta_values[p_values > 0.05]
cat("For HO cannot be rejected: [", min(theta_valid), max(theta_valid), "]\n")
```

```
## For HO cannot be rejected: [ 7.7 8.6 ]
```

Kolmogorov-Smirnov test can not be applied as it aims at two independent samples, but our data is paired.

(e) Since we are verifying the median now, the Sign test can be applied as follows:

```
n = length(data$After8weeks);below_6 = sum(data$After8weeks < 6);
binom.test(below_6,n,p=0.5,alt="l")
##</pre>
```

```
##
## Exact binomial test
##
## data: below_6 and n
## number of successes = 11, number of trials = 18, p-value = 0.9
## alternative hypothesis: true probability of success is less than 0.5
## 95 percent confidence interval:
## 0.000 0.801
## sample estimates:
## probability of success
## 0.611
```

We can conclude that H_0 cannot be rejected, since the p-value is higher than 0.05. We can set the original hypothesis H_0 as the percentage of cholesterol levels below 4.5 is less than 25%, while H_1 means the percentage of cholesterol levels below 4.5 is more than or equal to 25%.

```
below_4.5 = sum(data$After8weeks <= 4.5);n = length(data$After8weeks)
binom.test(below_4.5,n,p=0.25,alt="g")</pre>
```

```
##
## Exact binomial test
##
## data: below_4.5 and n
## number of successes = 3, number of trials = 18, p-value = 0.9
## alternative hypothesis: true probability of success is greater than 0.25
## 95 percent confidence interval:
## 0.047 1.000
## sample estimates:
## probability of success
## 0.167
```

Therefore, we cannot reject the original hypothesis H_0 since the p-value is larger than 0.05. We can conclude that the percentage of cholesterol levels below 4.5 is at most 25%.

Exercise 2

(a) Before we perform relevant ANOVA models, we need to check whether the data meets the necessary assumptions. For example, we need to check normality for numeric columns.

```
df <- read.table("crops.txt", header = TRUE)
df$County <- as.factor(df$County);df$Related <- as.factor(df$Related)
shapiro.test(df$Crops);shapiro.test(df$Size)</pre>
```

```
##
## Shapiro-Wilk normality test
##
## data: df$Crops
## W = 1, p-value = 0.3
##
## Shapiro-Wilk normality test
##
## data: df$Size
## W = 0.9, p-value = 0.001
```

From the result, the normaliy is doubtful for Crops.

We also check variance equality using leveneTest before applying anova model. As we can see p>0.05, thus we can apply anova models.

```
library(car)
## Loading required package: carData
leveneTest(Crops ~ County, data = df);leveneTest(Crops ~ Related, data = df);
## Levene's Test for Homogeneity of Variance (center = median)
         Df F value Pr(>F)
## group 2
                0.4
                      0.67
         27
##
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
##
## group 1
               1.09
                       0.3
##
         28
leveneTest(Crops ~ County:Related, data = df)
## Levene's Test for Homogeneity of Variance (center = median)
         Df F value Pr(>F)
## group 5
                0.3
                      0.91
##
         24
```

ANOVA model Without taking Size into account, there are two main factors: Country and Related, we also include the interaction factor: County+Related.

Tests the independent effects

Interaction model

```
anova_model2 <- aov(Crops ~ County*Related, data = df) ;summary(anova_model2)</pre>
```

Since for both model, all p-values are large (>0.05), we do not reject the null hypotheses, meaning there is no strong evidence that County or Related significantly impact Crops. As there is no significance for the interaction model, we choose the additive model to estimate the data.

```
new_farm <- data.frame(
   County = factor(3, levels = c(1, 2, 3)),
   Related = factor("no", levels = c("no", "yes"))
)
predicted_crops <- predict(anova_model1, newdata = new_farm, interval = "confidence");predicted_crops
## fit lwr upr
## 1 7760 6017 9504</pre>
```

The predicted crop yield for a farm in County 3 with no related factor is 7,760. However, the result may not be trustful.

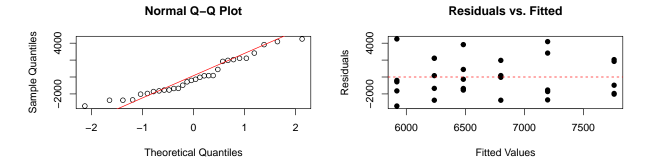
Here we do the post-hoc check using TukeyHSD.

TukeyHSD(anova_model2)

```
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
##
## Fit: aov(formula = Crops ~ County * Related, data = df)
##
## $County
##
       diff
              lwr upr p adj
## 2-1 -317 -3003 2369 0.953
## 3-1 960 -1726 3646 0.650
## 3-2 1277 -1409 3963 0.472
##
## $Related
##
          diff
                 lwr upr p adj
## yes-no -563 -2376 1249 0.527
##
## $'County:Related'
##
                diff
                       lwr upr p adj
## 2:no-1:no
                  93 -4610 4796 1.000
## 3:no-1:no
                 851 -3852 5554 0.993
              -362 -5065 4341 1.000
## 1:yes-1:no
## 2:yes-1:no -1090 -5792 3613 0.978
## 3:yes-1:no
                706 -3997 5409 0.997
## 3:no-2:no
                758 -3945 5461 0.996
## 1:yes-2:no -455 -5158 4248 1.000
```

The Tukey test results confirm that the differences in means are not statistically significant.

Examining residuals can help assess if the model assumptions hold. We can see that normality is doubtful for residuals. Also, we can see from the right picture that it shows non-random pattern, which suggests that the model is not well-fitted and may omit variables.



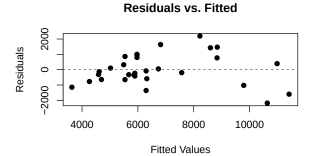
(b) ANOVA models only assumes that crops only depend on categorical factors, put size into perspective, we consider different ANCOVA models:

Tests the independent effects

```
ancova_main <- aov(Crops ~ County + Related + Size, data = df);summary(ancova_main)</pre>
##
               Df
                    Sum Sq Mean Sq F value
                                             Pr(>F)
                2 8.84e+06 4.42e+06
                                       3.71
## County
                                              0.039 *
## Related
                1 2.38e+06 2.38e+06
                                       2.00
                                              0.170
                1 1.10e+08 1.10e+08
                                      92.68 6.9e-10 ***
## Size
## Residuals
               25 2.98e+07 1.19e+06
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

We check the model assumptions. Here we can see from the residuals that they are approximately normally distributed.

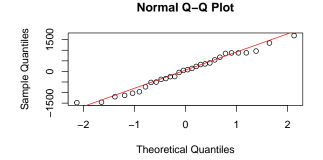
Normal Q-Q Plot Sample On Theoretical Quantiles

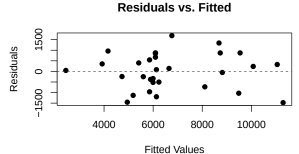


The interaction of Size X county

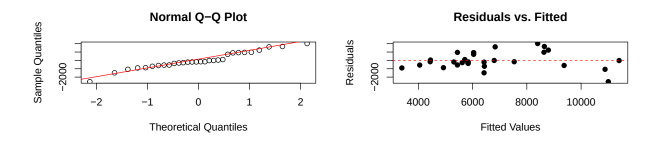
```
ancova_county_size <- aov(Crops ~ County * Size + Related, data = df);summary(ancova_county_size)</pre>
```

```
##
                    Sum Sq Mean Sq F value Pr(>F)
               Df
## County
                2 8.84e+06 4.42e+06
                                       5.01
                                            0.016 *
## Size
                1 1.11e+08 1.11e+08 126.47 8e-11 ***
## Related
                1 1.38e+06 1.38e+06
                                       1.57 0.223
## County:Size 2 9.53e+06 4.76e+06
                                       5.40 0.012 *
              23 2.03e+07 8.82e+05
## Residuals
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
par(mfrow = c(1,2))
residuals_county_size <- resid(ancova_county_size)</pre>
fitted_values_county_size <- fitted(ancova_county_size)</pre>
qqnorm(residuals_county_size);qqline(residuals_county_size, col="red")
plot(fitted values county size, residuals county size, main = "Residuals vs. Fitted",
     xlab = "Fitted Values", ylab = "Residuals", pch = 19)
abline(h = 0, col = "red", lty = 2)
```





```
ancova_related_size <- aov(Crops ~ County + Related * Size, data = df);summary(ancova_related_size)
##
                Df
                     Sum Sq Mean Sq F value Pr(>F)
## County
                 2 8.84e+06 4.42e+06
                                         3.73
                                                0.039 *
## Related
                 1 2.38e+06 2.38e+06
                                        2.01
                                                0.169
                 1 1.10e+08 1.10e+08
                                       93.21 9.7e-10 ***
## Related:Size 1 1.35e+06 1.35e+06
                                                0.296
                                         1.14
## Residuals
                24 2.85e+07 1.19e+06
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
par(mfrow = c(1,2))
residuals_related_size <- resid(ancova_related_size)</pre>
fitted_values_related_size <- fitted(ancova_related_size)</pre>
qqnorm(residuals_related_size);qqline(residuals_related_size, col="red")
plot(fitted_values_related_size,residuals_related_size , main = "Residuals vs. Fitted",
     xlab = "Fitted Values", ylab = "Residuals", pch = 19)
abline(h = 0, col = "red", lty = 2)
```



shapiro.test(residuals(ancova_main)); shapiro.test(residuals(ancova_related_size));

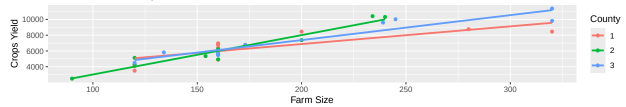
```
##
   Shapiro-Wilk normality test
##
##
## data: residuals(ancova_main)
## W = 1, p-value = 1
##
##
   Shapiro-Wilk normality test
##
## data: residuals(ancova_related_size)
## W = 1, p-value = 0.3
shapiro.test(residuals(ancova_county_size))
##
   Shapiro-Wilk normality test
##
##
## data: residuals(ancova county size)
## W = 1, p-value = 0.7
```

We can see from p-values that Size \times County interaction model has the lowest Residual Sum of Squares, from QQplot the normality is doubtful. But using Shapiro-Wilk normality test suggests that normality assumption holds. Thus we choose the Size \times County interaction model.

(c) We can review the ANCOVA Model Results.

Significant County (p < 0.05) \rightarrow County influences on Crops, meaning crop yields differ across counties. Significant Size (p < 0.05) \rightarrow Farm Size has a strong effect on Crops, which makes sense—larger farms generally produce more. Significant Related (p > 0.05) \rightarrow Whether the land lord and tenant are related does not significantly impact the crops. Significant County × Size (p < 0.05) \rightarrow This suggests that the effect of farm size on crop yield depends on the county. The figure blow confirms this conclusion, as the slopes are different.

Effect of Size on Crops Across Counties



(d)

```
new_farm <- data.frame(
   County = factor(2, levels = c(1, 2, 3)),
   Related = factor("yes", levels = c("no", "yes")),
   Size = 165
)
predicted_crops <- predict(ancova_county_size, newdata = new_farm, interval = "confidence")
predicted_crops</pre>
```

```
## fit lwr upr
## 1 6141 5428 6855
```

We estimate the error variance.

```
error_variance <- sum(residuals(ancova_county_size)^2) / df.residual(ancova_county_size) error_variance
```

[1] 881623

Exercise 3

(a)

```
library(MASS)
blocks = rep(1:6, each = 4)

random_N = unlist(tapply(rep(c(1, 1, 0, 0), 6), blocks, sample))
random_P = unlist(tapply(rep(c(1, 1, 0, 0), 6), blocks, sample))
random_K = unlist(tapply(rep(c(1, 1, 0, 0), 6), blocks, sample))
```

```
N_matrix <- matrix(random_N, nrow = 6, byrow = TRUE)
P_matrix <- matrix(random_P, nrow = 6, byrow = TRUE)
K_matrix <- matrix(random_K, nrow = 6, byrow = TRUE)
random_result = cbind(blocks, N_matrix, P_matrix, K_matrix)</pre>
```

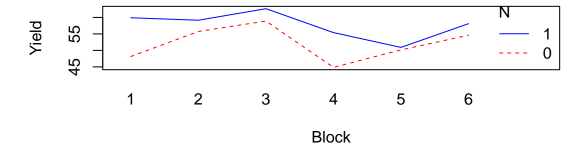
Warning in cbind(blocks, N_matrix, P_matrix, K_matrix): number of rows of ## result is not a multiple of vector length (arg 1)

```
##
         Block N1 N2 N3 N4 P1 P2 P3 P4 K1 K2 K3
## [1,]
                 1
                    0
                        1
                           0
                              0
                                  1
                                     1
                                         0
## [2,]
                 1
                        1
                           0
                                  0
                                     1
                                         0
## [3,]
                    1
                           0
                              1
                                  0
                                     1
## [4,]
                    0
                1
                        1
                           0
                                         0
## [5,]
             2
                1
                    0
                        0
                           1
                              0
## [6,]
```

(b) By generating the **interaction plot**, we can know that the mean yield of blocks using N is always higher than the blocks without using N. As interaction shows up as nonparallel curves, it looks like interaction seems to be not present.

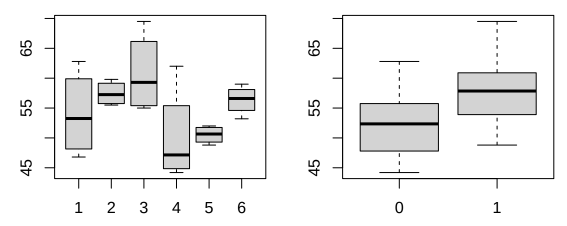
```
interaction.plot(npk$block, npk$N, npk$yield, xlab = 'Block', ylab = "Yield",
trace.label = "N", col = c("red", "blue"),
main = "The influence of using N or not on yield")
```

The influence of using N or not on yield



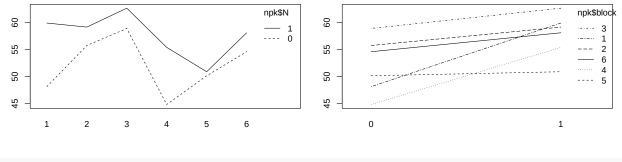
(c) First we look at the boxplot.

```
par(mfrow = c(1,2), mar = c(2, 3, 2, 1))
boxplot(npk$yield~npk$block);boxplot(npk$yield~npk$N);
```



From the interaction, we can observe that the lines seem parallel, so interaction might be no present.

```
par(mfrow = c(1,2), mar = c(3, 3, 2, 1))
interaction.plot(npk$block, npk$N, npk$yield);interaction.plot(npk$N, npk$block, npk$yield)
```



```
npk$block = as.factor(npk$block); npk$N = as.factor(npk$N);
npkaov = lm(yield~block*N, data=npk);anova(npkaov)
```

```
## Analysis of Variance Table
##
## Response: yield
##
             Df Sum Sq Mean Sq F value Pr(>F)
              5
                                   3.36
                                          0.04 *
## block
                   343
                           68.7
## N
              1
                   189
                          189.3
                                   9.26
                                          0.01 *
## block:N
              5
                    99
                           19.7
                                   0.96
                                          0.48
## Residuals 12
                   245
                           20.4
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
```

According to the result, there is no evidence for interaction between block and N, since the p-value of it is 0.48(>0.05). As the result of no interaction, we can remove interaction term from the model and fit the additive model.

```
npk$block = as.factor(npk$block); npk$N = as.factor(npk$N);
npkaov = lm(yield~block+N, data=npk);anova(npkaov)
```

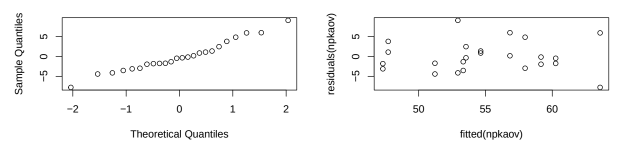
```
## Analysis of Variance Table
##
## Response: yield
##
             Df Sum Sq Mean Sq F value Pr(>F)
## block
              5
                   343
                           68.7
                                   3.40 0.0262 *
                   189
                          189.3
                                   9.36 0.0071 **
## N
              1
## Residuals 17
                           20.2
                   344
## ---
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

Then we can observe the p-value of each factor, it is known that both factors have a main effect in the additive model, since the p-values are smaller than 0.05.

We check the model assumptions. The Q-Q plot shows the normality.

```
par(mfrow = c(1,2))
qqnorm(residuals(npkaov)); plot(fitted(npkaov),residuals(npkaov))
```

Normal Q-Q Plot



The inclusion of Block factor is sensible for this model, because the block effect is significant in both models (p < 0.05), which means that different blocks have different average yields.

In this scenario, it's not suitable to use Friedman test, as from the dataset we can see that it is a replicated Complete Block Design(N>1). But actually we can do the aggregate operation to meet the requirement.

(d) From the results, we can know the pair values of each pair are all higher than 0.05, which means that there is no evidence for interaction in this pairs (N:block, P:block and K:block). Here, we can see N, K and block all present main effects, but we still cannot conclude this now.

```
model_N_block <- lm(yield ~ N * block + P + K, data = npk)
model_P_block <- lm(yield ~ P * block + N + K, data = npk)
model_K_block <- lm(yield ~ K * block + N + P, data = npk)
anova(model_N_block); anova(model_P_block); anova(model_K_block)</pre>
```

```
## Analysis of Variance Table
## Response: yield
             Df Sum Sq Mean Sq F value Pr(>F)
##
                           189.3
                                   13.36 0.0044 **
## N
               1
                    189
## block
               5
                    343
                            68.7
                                    4.85 0.0164 *
## P
               1
                      8
                             8.4
                                    0.59 0.4590
## K
               1
                     95
                            95.2
                                    6.72 0.0268 *
                                    1.39 0.3066
## N:block
               5
                     99
                            19.7
```

```
## Residuals 10
                           14.2
                    142
## ---
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' ' 1
  Analysis of Variance Table
##
##
## Response: yield
##
             Df Sum Sq Mean Sq F value Pr(>F)
## P
                      8
                            8.4
              1
                                   0.50 0.4966
## block
              5
                    343
                           68.7
                                   4.07 0.0282 *
                          189.3
## N
              1
                    189
                                  11.21 0.0074 **
## K
              1
                    95
                           95.2
                                   5.64 0.0389 *
              5
                    71
                           14.3
## P:block
                                   0.85 0.5473
## Residuals 10
                    169
                           16.9
## ---
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
## Analysis of Variance Table
##
## Response: yield
##
             Df Sum Sq Mean Sq F value Pr(>F)
                                   5.60 0.0395 *
## K
                    95
                           95.2
              1
## block
              5
                    343
                           68.7
                                   4.04 0.0288 *
## N
                    189
                          189.3
              1
                                  11.14 0.0075 **
## P
              1
                      8
                            8.4
                                   0.49 0.4980
              5
                    70
                           14.1
                                   0.83 0.5583
## K:block
## Residuals 10
                           17.0
                    170
## ---
                   0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' 1
## Signif. codes:
additive_model <- lm(npk$yield ~ npk$N + npk$block + npk$P + npk$K)
anova(additive_model)
## Analysis of Variance Table
##
## Response: npk$yield
##
             Df Sum Sq Mean Sq F value Pr(>F)
                    189
                          189.3
                                  11.82 0.0037 **
## npk$N
              1
              5
                    343
                           68.7
                                   4.29 0.0127 *
## npk$block
## npk$P
              1
                      8
                            8.4
                                   0.52 0.4800
## npk$K
              1
                    95
                           95.2
                                   5.95 0.0277 *
## Residuals 15
                           16.0
                    240
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

Therefore, we can conclude that P, K and block present significant effects in additive model, while P has no main effect in the additive model. Because in the first anova test, we cannot see interaction in each pair, so the additive model is better than the previous model with interaction terms.

- (e) According to the results in \mathbf{d} , we know N and K has the main effects in the additive model, because the p-value of N and K are both lower than 0.05. Hence, we can conclude that the combination of N and K can contribute to the yield most.
- (f) Our main question of interest is whether nitrogen N has an effect on yield. The mixed model processes the block variable as a random effect.

library(lme4) ## Loading required package: Matrix mixed_model <- lmer(yield ~ N + P + K + (1|block), REML=FALSE, data=npk) summary(mixed_model) ## Linear mixed model fit by maximum likelihood ['lmerMod'] ## Formula: yield ~ N + P + K + (1 | block) Data: npk ## ## ## AIC BIC logLik deviance df.resid ## 151.0 158.1 -69.5139.0 18 ## ## Scaled residuals: ## Min 1Q Median ЗQ ## -1.9706 -0.6831 0.0554 0.7124 1.4716 ## ## Random effects: ## Groups Name Variance Std.Dev. ## block (Intercept) 11.0 3.31 3.65 ## Residual 13.3 ## Number of obs: 24, groups: block, 6 ## Fixed effects: Estimate Std. Error t value 27.15 ## (Intercept) 54.65 2.01 ## N1 5.62 1.49 3.77 ## P1 -1.181.49 -0.79## K1 -3.98 1.49 -2.67## ## Correlation of Fixed Effects: (Intr) N1 P1 ## N1 -0.370 ## P1 -0.370 0.000 ## K1 -0.370 0.000 0.000

Then, we create a model without factor N and test the model inside the full one. Therefore, the Pr(>Chisq) is 0.0012, which means that the effect of N factor is significant. The results are the same as the fixed model.

```
mixed_without_N <- lmer(yield ~ P + K + (1|block), REML=FALSE,data=npk)
anova(mixed_model, mixed_without_N)</pre>
```

```
## Data: npk
## Models:
## mixed_without_N: yield ~ P + K + (1 | block)
## mixed_model: yield ~ N + P + K + (1 | block)
##
                  npar AIC BIC logLik deviance Chisq Df Pr(>Chisq)
## mixed_without_N
                     5 160 165 -74.7
                                            150
## mixed_model
                     6 151 158 -69.5
                                            139
                                               10.5 1
                                                            0.0012 **
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```