MA50259: Statistical Design of Investigations

Coursework 2 (2024)

09618

Disclaimer:

AI software (RStudio built-in Co-Pilot and ChatGPT) are used in this coursework for code, RMarkdown formatting, explanation suggestions, grammar check, and debugging purposes.

Part 1: Barley Experiment

The data in table below show the yields (measured in bushels per acre) of five varieties of barley in an experiment carried out in a rural area in the US.

Place	Year	Excel	Compana	Drummond	Conlon	Kindred
1	1971	71.0	95.4	109.7	119.5	88.3
1	1972	90.7	102.3	79.4	86.2	80.2
2	1971	132.6	121.0	140.7	171.5	135.7
2	1972	99.4	105.5	120.2	157.7	102.1
3	1971	87.3	73.3	79.4	128.6	85.3
3	1972	102.5	111.4	100.5	131.9	122.6
4	1971	129.8	111.3	121.5	148.8	114.8
4	1972	96.9	60.3	83.2	118.5	72.4
5	1971	92.3	81.4	72.1	84.8	99.1
5	1972	65.2	48.9	91.5	72.8	77.4
6	1971	84.3	72.1	76.9	109.8	90.0
6	1972	63.3	68.4	62.7	97.8	92.2

(a) What would be the purpose of running these experiments in different locations and years? Also, comment on the design that would be appropriate for analysing the data above.

Answer:

 ${\it\# Reference: Lawson, J., 2014. *Design and Analysis of Experiments with R*. Florida: CRC Press.}$

So that variability caused by geographical and meteorological conditions can be removed from the error sum of squares.

Doing so meant that the conclusions can be generalised over the range of geographical and meteorological conditions that are being studied.

Some varieties might perform exceptionally well in one location but poorly in another, or some might have good years and bad years depending on external conditions.

An appropriate design for analysing the data would be a Randomised Complete Block Design (RCBD).

(b) Organise the data in an appropriate dataframe in R with barley yields as the response variable, places as the blocks and the five varieties of barley as the treatment effects. Use an appropriate model which includes block effects and treatment effects to perform the ANOVA and determine if there is a significant difference across barley varieties. Clearly state the assumptions you may need in the model and write the null and the alternative hypotheses considered in the ANOVA.

Answer:

```
# Create the dataframe
# Reference: https://www.math.mcgill.ca/dstephens/OldCourses/204-2008/Handouts/Math204-07-RBDANOVA.pdf
barley_data <- data.frame(</pre>
  Place = factor(rep(1:6, each = 2)), # Ensure 'Place' is treated as a factor
  Year = factor(rep(c(1971, 1972), times = 6)),
  Excel = c(71.0, 90.7, 132.6, 99.4, 87.3, 102.5, 129.8, 96.9, 92.3, 65.2, 84.3, 63.3)
  Compana = c(95.4, 102.3, 121.0, 105.5, 73.3, 111.4, 111.3, 60.3, 81.4, 48.9, 72.1, 68.4)
  Drummond = c(109.7, 79.4, 140.7, 120.2, 79.4, 100.5, 121.5, 83.2, 72.1, 91.5, 76.9, 62.7),
  Conlon = c(119.5, 86.2, 171.5, 157.7, 128.6, 131.9, 148.8, 118.5, 84.8, 72.8, 109.8, 97.8)
  Kindred = c(88.3, 80.2, 135.7, 102.1, 85.3, 122.6, 114.8, 72.4, 99.1, 77.4, 90.0, 92.2)
)
# Reshape data from wide to long format
barley_long <- barley_data %>%
  pivot longer(cols = Excel:Kindred, names to = "Variety", values to = "Yield")
# Fit the ANOVA model with block effects (place) and treatment effects
anova_model <- aov(Yield ~ Place + Variety, data = barley_long)</pre>
summary(anova_model)
               Df Sum Sq Mean Sq F value
##
                                           Pr(>F)
## Place
                   16921
                            3384
                                 11.348 2.41e-07 ***
                    7031
## Variety
                4
                            1758
                                   5.894 0.000576 ***
                  14912
                             298
## Residuals
               50
```

0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1 ## Signif. codes:

The model can be expressed as:

$$y_{ij} = \mu + b_i + \tau_j + \epsilon_{ij}$$

where:

- y_{ij} is the yield of the jth variety in the ith block,
- μ is the overall mean yield,
- b_i is the effect of the *i*th block (place),
- τ_i is the effect of the jth variety,

• ϵ_{ij} is the random error term.

The ANOVA model relies on several key assumptions:

- Independence: The observations within each group should be independent of each other.
- Normality: The response variable should be normally distributed for each group.
- Homogeneity of variances: The variances among different groups should be equal.

The hypotheses for the ANOVA are formulated as follows:

- H_0 (Null Hypothesis): $H_0: \tau_1 = \tau_2 = \tau_3 = \tau_4 = \tau_5 = 0$ (There are no differences between barley varieties in terms of yield)
- H_a (Alternative Hypothesis): H_a : At least one $\tau_j \neq 0$ (At least one barley variety has a different yield compared to the others)

The ANOVA results uses the F-statistic to test the null hypothesis.

The F-value obtained has its associated p-value, which will help us determine the significance

p-value is 0.000576, which is less than 0.05. Therefore, at a significance level of 0.05, we reject the null hypothesis. This suggests that there are statistically significant differences in the yields across the barley varieties.

(c) Omit the block effects in part (b) and use an appropriate model to perform the ANOVA and determine if there is a significant difference across barley varieties. Clearly state the assumptions you may need in the model and write the null and the alternative hypotheses considered in the ANOVA.

Answer:

```
# Recreate the data frame
data <- data.frame(</pre>
  # Place = rep(1:6, each = 2),
  # Year = rep(c(1971, 1972), times = 6),
  Place = factor(rep(1:6, each = 2)),
  Year = factor(rep(c(1971, 1972), times = 6)),
  Excel = c(71.0, 90.7, 132.6, 99.4, 87.3, 102.5, 129.8, 96.9, 92.3, 65.2, 84.3, 63.3)
  Compana = c(95.4, 102.3, 121.0, 105.5, 73.3, 111.4, 111.3, 60.3, 81.4, 48.9, 72.1, 68.4)
  Drummond = c(109.7, 79.4, 140.7, 120.2, 79.4, 100.5, 121.5, 83.2, 72.1, 91.5, 76.9, 62.7),
  Conlon = c(119.5, 86.2, 171.5, 157.7, 128.6, 131.9, 148.8, 118.5, 84.8, 72.8, 109.8, 97.8)
  Kindred = c(88.3, 80.2, 135.7, 102.1, 85.3, 122.6, 114.8, 72.4, 99.1, 77.4, 90.0, 92.2)
# Convert data to long format
data_long <- data %>%
  pivot_longer(cols = Excel:Kindred, names_to = "Variety", values_to = "Yield")
# Fit the ANOVA model considering only the Variety effect
model_no_blocks <- aov(Yield ~ Variety, data = data_long)</pre>
```

```
# Perform ANOVA
anova_results_no_blocks <- summary(model_no_blocks)
anova_results_no_blocks</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## Variety    4    7031   1757.7   3.037   0.0247 *
## Residuals    55   31833   578.8
## ---
## Signif. codes:    0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The model is now:

$$y_{ij} = \mu + \tau_j + \epsilon_{ij}$$

where:

- y_{ij} is the yield of the jth variety in the ith block,
- μ is the overall mean yield,
- τ_j is the effect of the jth variety,
- ϵ_{ij} is the random error term.

Assumptions of the ANOVA Model:

- Independence: The observations within each group should be independent of each other.
- Normality: The response variable should be normally distributed for each group.
- Homogeneity of variances: The variances among different groups should be equal.

The ANOVA results uses the F-statistic to test the null hypothesis.

The F-value obtained has its associated p-value, which will help us determine the significance

The hypotheses for the ANOVA are formulated as follows:

- H_0 (Null Hypothesis): $H_0: \tau_1 = \tau_2 = \tau_3 = \tau_4 = \tau_5 = 0$ (There are no differences between barley varieties in terms of yield)
- H_a (Alternative Hypothesis): H_a : At least one $\tau_j \neq 0$ (At least one barley variety has a different yield compared to the others)

Here are the results from the ANOVA performed without considering the block effects (Place):

p-value is 0.0247, which is less than 0.05. Therefore, at a significance level of 0.05, we reject the null hypothesis. This suggests that there are statistically significant differences in the yields across the barley varieties.

⁽d) Which of the two designs represented by the two models in part (b) and part (c) would be more efficient. Justify your answer.

Answer:

Reducing variance in an experimental design will increase the precision of the estimates of the effects being studied. The inclusion of block effects, as in part (b), aims to control for variability attributable to differences in environmental conditions, soil types, microclimates, etc., at different locations. By accounting for these block effects, we can observe:

From the ANOVA results:

- With Block Effects (Part B): Both the variety and the place had significant effects on the yields, with the model showing that variability due to location is significant.
- Without Block Effects (Part C): Only the variety effect was tested, and it showed significance but with a larger residual variance, because some of the variations due to the place effect have been unaccounted for.

Models that account for blocking factors are generally more robust across different environments because they adjust for potential factors that affects our experimental setup. This can make findings more generalisable across similar conditions beyond the scope of the experiment.

Part 2A: Chronic Respiratory Disease Study

An increase in deaths due to chronic respiratory disease (CRD) was observed in certain parts of the UK after the millennium. A case-control study was carried out to investigate the possible association between prescription of one particular medication, namely X12 and CRD deaths.

Both cases and controls were chosen among persons who were admitted to hospital for CRD. The cases comprised 257 persons who died of CRD; the controls were 570 persons who did not die of CRD. The data are presented in the following table.

X12 prescribed	Cases	Controls
Yes	130	200
No	127	370
Total	257	570

(a) Obtain the odds ratio between X12 prescription and CRD deaths and the corresponding 95% confidence interval from the above table.

Answer:

```
# Reference: https://www.youtube.com/watch?v=jRQ2nP7lAoU
# Reference: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8198228/
# Create a 2x2 table for the data
X12_table <- matrix(c(130, 127, 200, 370), nrow = 2, byrow = TRUE)</pre>
```

```
colnames(X12_table) <- c("Cases", "Controls")</pre>
rownames(X12_table) <- c("Yes", "No")</pre>
# Calculate the odds ratio
odds_ratio <- X12_table[1, 1] * X12_table[2, 2] / (X12_table[1, 2] * X12_table[2, 1])
# Round the odds ratio to two decimal places
odds ratio <- round(odds ratio, 2)
# Calculate the standard error of the log odds ratio
log_odds_ratio <- log(odds_ratio)</pre>
se_log_odds_ratio <- sqrt(1 / X12_table[1, 1] + 1 / X12_table[1, 2] +</pre>
                              1 / X12_table[2, 1] + 1 / X12_table[2, 2])
# Calculate the 95% confidence interval for the odds ratio
ci_lower <- exp(log_odds_ratio - 1.96 * se_log_odds_ratio)</pre>
ci_upper <- exp(log_odds_ratio + 1.96 * se_log_odds_ratio)</pre>
# Round the confidence interval to two decimal places
ci_lower <- format(round(ci_lower, 2), nsmall = 2)</pre>
ci_upper <- format(round(ci_upper, 2), nsmall = 2)</pre>
# Print the results
cat("The odds ratio (OR) between X12 prescription and CRD deaths is",
    odds ratio, ".\n")
```

The odds ratio (OR) between X12 prescription and CRD deaths is 1.89 .

```
cat("The 95% confidence interval for the odds ratio is (",
    ci_lower, ", ", ci_upper, ").\n")
```

 $\mbox{\tt \#\#}$ The 95% confidence interval for the odds ratio is (1.40 , $\ 2.55$).

The odds ratio (OR) is calculated as follows:

$$OR = \frac{\text{odds of exposure among cases}}{\text{odds of exposure among controls}}$$

Where:

- Odds of exposure among cases = $\frac{\text{Number of exposed cases}}{\text{Number of unexposed cases}} = \frac{130}{127}$
- Odds of exposure among controls = $\frac{\text{Number of exposed controls}}{\text{Number of unexposed controls}} = \frac{200}{370}$

We can calculate the OR using these ratios. Additionally, we will calculate the 95% confidence interval (CI) for the odds ratio using the formula:

$$\log(OR) \pm 1.96 \times SE(\log(OR))$$

where the standard error (SE) of the log odds ratio is:

$$SE(\log(OR)) = \sqrt{\frac{1}{a} + \frac{1}{b} + \frac{1}{c} + \frac{1}{d}}$$

with a, b, c, and d being the cell frequencies in the 2x2 table:

$$a = 130$$
 $b = 200$
 $c = 127$ $d = 370$

After computing the OR and its 95% CI, we get:

The odds ratio (OR) is 1.89. This indicates that the ratio of those prescribed with X12 to those not prescribed with X12 was 1.89 times higher in the group of that died (cases) compared to those who did not die (controls).

The null hypothesis:

$$H_0 : OR = 1$$

The alternative hypothesis:

$$H_a: OR \neq 1$$

The corresponding 95% confidence interval for the odds ratio ranges from approximately 1.40 to 2.55. This means we are 95% confident that the true odds ratio lies within this interval, suggesting a statistically significant association between X12 prescription and CRD deaths as the interval does not include 1. Thus, at a significance level of 0.05, we reject the null hypothesis.

(b) Test the null hypothesis of no association between X12 prescribed and CRD deaths. Interpret the results in the context of the study.

Answer

```
# Reference: https://vula.uct.ac.za/access/content/group/9c29ba04-b1ee-49b9-8c85-9a468b556ce2/DOH/Modul
# Define the 2x2 contingency table
contingency_table <- matrix(c(130, 127, 200, 370), nrow = 2, byrow = TRUE)

# Perform the chi-square test for independence
chi2_test <- chisq.test(contingency_table, correct = FALSE)

# Extract the test statistic and p-value
test_statistic <- chi2_test$statistic
p_value <- chi2_test$p.value

# Round the test statistics and p-value
test_statistic <- round(test_statistic, 2)
p_value <- format(p_value, scientific = TRUE, digits = 2)

# Print the results
cat("The chi-square test statistic is approximately", test_statistic, ".\n")</pre>
```

The chi-square test statistic is approximately 17.74 .

cat("The p-value is about", p_value, ".\n")

The p-value is about 2.5e-05 .

We can use the chi-square test for independence in a 2x2 contingency table.

In table format:

	Cases	Controls	Total
X12 prescribed	130	200	330
No	127	370	497
Total	257	570	827

Null Hypothesis (H_0) : There is no association between X12 prescription and CRD deaths.

Alternative Hypothesis (H_a) : There is an association between X12 prescription and CRD deaths.

The chi-square test works by:

$$E_{ij} = \frac{(\text{row total}_i \times \text{column total}_j)}{\text{total observations}}$$

where E_{ij} is the expected frequency for cell (i, j), and the row and column totals are the sums of the observed frequencies in the respective rows and columns.

The chi-square test statistic is calculated as:

$$\chi^2 = \sum \frac{(O_{ij} - E_{ij})^2}{E_{ij}}$$

where O_{ij} is the observed frequency for cell (i,j) and E_{ij} is the expected frequency for the same cell.

Alternatively, we can use the formula to calculate the chi-square (χ^2) statistic:

$$\chi^2 = \frac{(ad - bc)^2 \times (a + b + c + d)}{(a + b)(c + d)(a + c)(b + d)}$$

Where:

- a = 130 (Exposed cases)
- b = 200 (Exposed controls)
- c = 127 (Unexposed cases)
- d = 370 (Unexposed controls)

Next.

The degrees of freedom for a 2x2 contingency table are 1.

This is calculated as:

degrees of freedom = (number of rows -1) × (number of columns -1)

The p-value is then determined based on the chi-square statistic and the degrees of freedom using a chi-square distribution table (built-in to R chi-square test function).

Let's perform the chi-square test to determine if there is a significant association between X12 prescription and CRD deaths.

The chi-square test statistics (Without Yates' correction) is 17.74, and the p-value is about 2.5e-05.

This p-value is very small (less than 0.05), indicating that the test result is statistically significant. Therefore, at a significance level of 0.05, we reject the null hypothesis of no association between X12 prescription and CRD deaths. This suggests that there is a statistically significant association between being prescribed X12 and the likelihood of dying from chronic respiratory disease.

This finding could imply that X12 is either a risk factor for CRD mortality or is prescribed more frequently in more severe cases, or it could be associated with other confounding factors. Further investigation would be needed to clarify the nature of this association.

(c) There was a concern that cases and control may have differed according to the underlying severity of their CRD. Indeed, disease severity may be associated with a lifestyle habit such as smoking, and hence is a potential confounder. Accordingly, the data were stratified by variables associated with severity. One such indicator of severity is smoking habit in the previous year. The data, stratified by this variable, is shown in the following table

Non-smokers			Smokers		
X12 prescribed	Cases	Controls	X12 prescribed	Cases	Controls
Yes	74	170	Yes	56	30
No	100	267	No	27	103
Total	174	437	Total	83	133

Estimate the odds ratio and calculate the corresponding 95% confidence interval for each stratum.

Answer:

To estimate the odds ratio and calculate the corresponding 95% confidence interval for each stratum (non-smokers and smokers), we will use the counts provided in the table for both strata. The approach involves calculating the odds ratio within each stratum and then computing the confidence intervals similarly to the previous part.

For Non-Smokers:

a=74(Exposed cases)
b=170 (Exposed controls)
c=100 (Unexposed cases)
d=267 (Unexposed controls)

For Smokers:

a=56 (Exposed cases)
b=30 (Exposed controls)
c=27 (Unexposed cases)
d=103 (Unexposed controls)

We'll calculate the odds ratio for each stratum using the formula:

$$OR = \frac{\mathrm{ad}}{\mathrm{bc}}$$

And then calculate the 95% confidence interval using:

$$\log(OR) \pm 1.96 \times SE(\log(OR))$$

Where the standard error is:

$$SE(\log(OR)) = \sqrt{\frac{1}{a} + \frac{1}{b} + \frac{1}{c} + \frac{1}{d}}$$

The null hypothesis:

$$H_0 : OR = 1$$

The alternative hypothesis:

$$H_a: OR \neq 1$$

Let's perform these calculations for both strata.

```
# Create 2x2 contingency tables for non-smokers and smokers
non_smokers_table <- matrix(c(74, 100, 170, 267), nrow = 2, byrow = \frac{TRUE}{TRUE})
smokers_table \leftarrow matrix(c(56, 27, 30, 103), nrow = 2, byrow = TRUE)
# Calculate the odds ratio and 95% confidence interval for non-smokers
odds_ratio_non_smokers <- non_smokers_table[1, 1]* non_smokers_table[2, 2] /
  (non_smokers_table[1, 2] * non_smokers_table[2, 1])
log_odds_ratio_non_smokers <- log(odds_ratio_non_smokers)</pre>
se_log_odds_ratio_non_smokers <- sqrt(1 / non_smokers_table[1, 1] +</pre>
                                          1 / non_smokers_table[1, 2] +
                                          1 / non smokers table[2, 1] +
                                         1 / non_smokers_table[2, 2])
ci_lower_non_smokers <- exp(log_odds_ratio_non_smokers - 1.96 * se_log_odds_ratio_non_smokers)
ci_upper_non_smokers <- exp(log_odds_ratio_non_smokers + 1.96 * se_log_odds_ratio_non_smokers)
# Calculate the odds ratio and 95% confidence interval for smokers
odds_ratio_smokers <- smokers_table[1, 1] * smokers_table[2, 2] /
  (smokers_table[1, 2] * smokers_table[2, 1])
log_odds_ratio_smokers <- log(odds_ratio_smokers)</pre>
se_log_odds_ratio_smokers <- sqrt(1 / smokers_table[1, 1] +</pre>
                                     1 / smokers table[1, 2] +
                                     1 / smokers_table[2, 1] +
                                     1 / smokers_table[2, 2])
ci_lower_smokers <- exp(log_odds_ratio_smokers - 1.96 * se_log_odds_ratio_smokers)
ci_upper_smokers <- exp(log_odds_ratio_smokers + 1.96 * se_log_odds_ratio_smokers)</pre>
```

```
# Round the results
odds_ratio_non_smokers <- round(odds_ratio_non_smokers, 2)</pre>
ci_lower_non_smokers <- round(ci_lower_non_smokers, 2)</pre>
ci_upper_non_smokers <- round(ci_upper_non_smokers, 2)</pre>
odds ratio smokers <- round(odds ratio smokers, 2)
ci_lower_smokers <- round(ci_lower_smokers, 2)</pre>
ci_upper_smokers <- round(ci_upper_smokers, 2)</pre>
cat ("Non-Smokers Odds Ratio:", odds_ratio_non_smokers)
## Non-Smokers Odds Ratio: 1.16
cat ("\n")
# cat ("Non-Smokers CI:", ci_lower_non_smokers, ci_upper_non_smokers)
cat ("Non-Smokers CI:", "(", ci_lower_non_smokers, ci_upper_non_smokers,")")
## Non-Smokers CI: ( 0.81 1.66 )
cat ("\n")
cat ("Smokers Odds Ratio:", odds_ratio_smokers)
## Smokers Odds Ratio: 7.12
cat ("\n")
cat ("Smokers CI:", "(",ci_lower_smokers, ci_upper_smokers,")")
## Smokers CI: ( 3.86 13.15 )
```

Interpretation:

For non-smokers:

- The OR is 1.16.
- This indicates that the ratio of those prescribed with X12 to those not prescribed with X12 was about 1.16 times higher when we compare non-smokers group who died of CRD (cases) to non-smokers group who did not die of CRD (controls). Since confidence interval includes 1, the association is not statistically significant for non-smokers. At a significance level of 0.05, we fail to reject the null hypothesis.

For smokers:

- The OR is 7.12.
- This indicates that the ratio of those prescribed with X12 to those not prescribed with X12 was about 7.12 times higher when we compare among smokers who died of CRD (cases) compared to smokers who did not die of CRD (controls). The confidence interval does not include 1, suggesting a statistically significant association for smokers. At a significance level of 0.05, we reject the null hypothesis.

These results imply that the effect of X12 prescription on CRD mortality may be much more pronounced in smokers than in non-smokers.

(d) Calculate the overall regression summary odds ratio (together with its corresponding 95% confidence interval) when adjusting for smoking. Interpret the results in the context of the study.

Answer:

To calculate the overall adjusted odds ratio that accounts for smoking as a potential confounding factor, we can use the Mantel-Haenszel method. This method provides a weighted average of the odds ratios from each stratum (here, smokers and non-smokers), adjusting for the confounding effects of smoking.

The formula for the Mantel-Haenszel summary odds ratio is:

The Mantel-Haenszel odds ratio (MH OR) is given by:

MH OR =
$$\frac{\sum (a_i \cdot d_i/n_i)}{\sum (b_i \cdot c_i/n_i)}$$

Where n_i is the total number of individuals in the *i*-th stratum. We will also calculate the corresponding 95% confidence interval for the Mantel-Haenszel summary odds ratio. The standard error of the log(MH OR) is calculated as follows:

95% CI for a odds ratio (OR) ranges from OR/EF to OR*EF, where EF is the exposure factor;

Where $EF = \exp(1.96 \times SE)$, where exp is exponential and SE is the standard error of the odds ratio.

When odds ration is the Mantel-Haenszel summary odds ratio, the confidence interval is calculated as follows:

$$SE = \sqrt{\frac{V}{Q \times R}}$$

Where:

```
• Q = \sum (a_i \cdot b_i/n_i)

• R = \sum (c_i \cdot d_i/n_i)

• V = \sum (a_i + b_i) \cdot (c_i + d_i) \cdot (a_i + c_i) \cdot (b_i + d_i)/(n_i^2 \cdot (n_i - 1)).
```

Let's compute the Mantel-Haenszel summary odds ratio and its confidence interval.

```
# Find MH OR
a_ns <- 74  # Exposed cases
b_ns <- 170  # Exposed controls
c_ns <- 100  # Unexposed cases
d_ns <- 267  # Unexposed controls
n_ns <- a_ns + b_ns + c_ns + d_ns  # Total non-smokers

# Smokers Data
a_s <- 56  # Exposed cases
b_s <- 30  # Exposed controls
c_s <- 27  # Unexposed cases
d_s <- 103  # Unexposed controls</pre>
```

```
n_s <- a_s + b_s + c_s + d_s # Total smokers
# Calculate Mantel-Haenszel Odds Ratio
mh_numerator <- (a_ns * d_ns / n_ns) + (a_s * d_s / n_s)</pre>
mh_denominator \leftarrow (b_ns * c_ns / n_ns) + (b_s * c_s / n_s)
MH_OR <- mh_numerator / mh_denominator</pre>
cat("The Mantel-Haenszel summary odds ratio (MH OR) is approximately", round(MH_OR, 2), ".\n")
## The Mantel-Haenszel summary odds ratio (MH OR) is approximately 1.87.
# Reworked approach
# Data for non-smokers and smokers
# Non-Smokers Data
# Reference: https://www.ncss.com/wp-content/themes/ncss/pdf/Procedures/NCSS/Mantel-Haenszel Test.pdf
# Reference: https://www.youtube.com/watch?v=9bMxD3MGBik&list=PLLTSM0eKjC2cG57KjzfV0Cymduzv_mEvy&index=
# Calculate Q, R, V for Mantel-Haenszel SE calculation
Q \leftarrow (a_ns * b_ns / n_ns) + (a_s * b_s / n_s)
R \leftarrow (c_ns * d_ns / n_ns) + (c_s * d_s / n_s)
V \leftarrow ((a_ns + b_ns) * (c_ns + d_ns) * (a_ns + c_ns) * (b_ns + d_ns) / (n_ns^2 * (n_ns - 1))) +
    ((a_s + b_s) * (c_s + d_s) * (a_s + c_s) * (b_s + d_s) / (n_s^2 * (n_s - 1)))
# Calculate the more precise SE
SE \leftarrow sqrt(V / (Q * R))
# Exposure Factor and Confidence Interval using more precise SE
EF \leftarrow exp(1.96 * SE)
ci lower mh precise <- MH OR / EF
ci_upper_mh_precise <- MH_OR * EF</pre>
# Round the results
MH_OR <- round(MH_OR, 2)</pre>
ci_lower_mh_precise <- round(ci_lower_mh_precise, 2)</pre>
ci_upper_mh_precise <- round(ci_upper_mh_precise, 2)</pre>
cat("The Mantel-Haenszel summary odds ratio (MH OR) is approximately", MH_OR, ".\n")
## The Mantel-Haenszel summary odds ratio (MH OR) is approximately 1.87 .
cat("The 95% confidence interval for the Mantel-Haenszel summary odds ratio is (", ci_lower_mh_precise,
## The 95% confidence interval for the Mantel-Haenszel summary odds ratio is (1.36, 2.57).
Null Hypothesis:
                                       H_0: MH OR = 1
```

Alternative Hypothesis:

 $H_a: MH OR \neq 1$

The Mantel-Haenszel summary odds ratio (MH OR), when adjusting for smoking, is approximately 1.87. The corresponding 95% confidence interval ranges from 1.36 to 2.57.

The MH OR of approximately 1.87 suggests a stronger association between being prescribed medication X12 and the risk of dying from chronic respiratory disease (CRD), taking into account the smoking status as a stratifying factor.

The confidence interval (1.36 to 2.57) does not include 1, which further supports the statistical significance of this finding. At a significance level of 0.05, we reject the null hypothesis.

This analysis highlights that X12 prescription may increase the risk of CRD mortality irrespective of smoking status, though the effect is more pronounced in smokers as seen in the stratified analysis.

(e) Compare the odds ratios you obtained in parts (a) and (d). How did confounding by smoking affect the apparent direction of association between X12 and CRD deaths?

Answer:

In part (a), where we didn't account for smoking, the odds ratio (OR) calculated for the association between X12 prescription and CRD deaths was approximately 1.89.

In part (d), after adjusting for smoking using the Mantel-Haenszel method, the summary odds ratio was approximately 1.87. This odds ratio, though very similar in magnitude to the unadjusted odds ratio, now explicitly accounts for smoking as a confounding factor.

This analysis highlights that X12 prescription may increase the risk of CRD mortality irrespective of smoking status, though the effect is more pronounced in smokers as seen in the stratified analysis.

The magnitude of the association between X12 prescription and CRD deaths remained relatively unchanged (from 1.89 to 1.87) after adjusting for smoking. This suggests that the overall effect of X12 on the risk of CRD deaths is not heavily confounded by smoking, as the adjustment did not markedly alter the odds ratio.

Adjusting for smoking suggests that the relationship between presence of X12 prescription and increased mortality from CRD is robust and not merely a product of confounding due to differences in smoking habits among the cases and controls.

(f) Which other variables that you can think of could have been used as confounders to have a better picture of the association between X12 and CRD deaths? You should justify your answer.

Answer:

Socio-economic Status (SES): Socio-economic factors including income, education, and occupational exposures can influence health behaviours, lifestyle factors, access to healthcare, adherence to treatments, and overall health outcomes. Lower SES is often associated with poorer health outcomes and might influence the type of treatment received.

Age: Age is a fundamental determinant of health status and is strongly associated with both the likelihood of being prescribed certain medications and the risk of mortality from chronic diseases, including CRD. Older individuals may have both higher medication use and higher mortality rates.

Environmental Exposures: Exposure to pollutants or allergens, particularly in industrial or high-traffic areas, can exacerbate CRD and may influence both the severity of the disease and the treatment options prescribed.

Part 2B: Low Birth Weight Study

Consider a cohort study on birth weight (BW) of singleton babies and lifestyle habits (smoking, drinking, exercise etc.) of their mothers. Style 1 relates to unhealthy habits such as smoking/drinking and little exercise, and Style 2 relates to healthy habits including no smoking, no drinking and sufficient exercise. The number of babies born with low birth weight (LBW) within a 1-year period to Style 1 and Style 2 are 34 and 10 respectively. The total number of mothers who predominantly follow Style 1 and Style 2 are 335 and 320 respectively.

(a) Calculate the risk difference and the relative risk of LBW in the general population for the two lifestyle habits.

Answer:

First we organise the data into a table with header on Lifestyle, LBW, NBW, and Total.

Lifestyle	LBW	NBW	Total
Style 1	34	301	335
Style 2	10	310	320
Total	44	611	655

To analyse the data from the birth weight study, we need to calculate two key epidemiological measures: the Risk Difference (RD) and the Relative Risk (RR). Here's how we can calculate each:

Risk Difference (RD): This measures the absolute difference in risk (or probability) of an outcome between two groups. It is given by:

$$RD = P1 - P2$$

where P1 is the proportion of low birth weight babies among mothers with Style 1 and P2 is the proportion among mothers with Style 2.

Relative Risk (RR): This measures the ratio of the probability of an event occurring in the exposed group to the probability of the event occurring in the control group. It is calculated as:

$$RR = \frac{P1}{P2}$$

where P1 and P2 are as defined above.

Let's first calculate the proportion of low birth weight (LBW) babies for each lifestyle:

- Style 1: $P1 = \frac{\text{Number of LBW babies in Style 1}}{\text{Total number of mothers in Style 1}}$
- Style 2: $P2 = \frac{\text{Number of LBW babies in Style 2}}{\text{Total number of mothers in Style 2}}$

We will use these proportions to calculate the RD and RR. Given Data:

```
Total mothers in Style 1: 335
LBW babies in Style 2: 10
Total mothers in Style 2: 320
Now, let's compute the proportions, RD, and RR.
\# \ Reference: \ https://handbook-5-1.cochrane.org/chapter\_9/9\_2\_2\_4\_measure\_of\_absolute\_effect\_the\_risk\_diagrams and the substitution of the su
# Define the data
LBW_Style1 <- 34
Total_Style1 <- 335
LBW_Style2 <- 10
Total_Style2 <- 320
# Calculate the proportions of LBW babies for each lifestyle
P1 <- LBW_Style1 / Total_Style1
P2 <- LBW_Style2 / Total_Style2
# Calculate the Risk Difference (RD)
RD <- P1 - P2
# Calculate the Relative Risk (RR)
RR <- P1 / P2
P1 <- round(P1, 4)
P2 <- round(P2, 4)
RD <- round(RD, 4)
RR <- round(RR, 2)
cat("Proportion of LBW babies in Style 1:", P1)
## Proportion of LBW babies in Style 1: 0.1015
cat("\n")
cat("Proportion of LBW babies in Style 2:", P2)
## Proportion of LBW babies in Style 2: 0.0312
cat("\n")
cat("Risk Difference (RD):", RD)
## Risk Difference (RD): 0.0702
cat("\n")
cat("Relative Risk (RR):", RR)
```

LBW babies in Style 1: 34

Relative Risk (RR): 3.25

These results indicate that the risk of having a baby with low birth weight is 7.02% (RD = 0.0702) higher for mothers with unhealthy lifestyle habits (Style 1) compared to those with healthy lifestyle habits (Style 2). Furthermore, babies born to mothers with Style 1 have a relative risk 3.25 times greater of being of low birth weight compared to those from mothers with Style 2.

(b) Find a 95% confidence interval for the risk difference and the relative risk of LBW in the general population for the two lifestyle habits. Interpret the results in the context of the study.

Answer:

Reference:

To determine the confidence intervals (CI) for both the Risk Difference (RD) and the Relative Risk (RR), we can use standard methods for proportions:

Confidence Interval for Risk Difference (RD): The standard error (SE) for the RD between two independent proportions can be calculated by:

$$SE_{(RD)} = \sqrt{\frac{P_1(1-P_1)}{n_1} + \frac{P_2(1-P_2)}{n_2}}$$

The 95% confidence interval for RD is then:

$$RD \pm 1.96 \times SE_{(RD)}$$

Confidence Interval for Relative Risk (RR): We first compute the natural log of the RR (ln(RR))) and then find the standard error:

$$SE_{ln(RR)} = \sqrt{\frac{1 - P_1}{n_1 \times P_1} + \frac{1 - P_2}{n_2 \times P_2}}$$

The 95% CI for ln(RR) is:

$$ln(RR) \pm 1.96 \times SE_{(ln(RR))}$$

We then exponentiate the limits of this interval to get the CI for RR.

by the formula:

$$CI = e^{\ln(RR) \pm 1.96 \times SE_{\ln(RR)}}$$

Let's calculate these confidence intervals using the given data:

```
# References:
# https://www.youtube.com/watch?v=jRQ2nP7lAoU
# https://www.youtube.com/watch?v=BmXJ_jJIGEO
# https://sphweb.bumc.bu.edu/otlt/mph-modules/ep/ep713_randomerror/EP713_RandomError5.html
# Calculate the standard error for Risk Difference (RD)
SE_RD <- sqrt(P1 * (1 - P1) / Total_Style1 + P2 * (1 - P2) / Total_Style2)</pre>
```

```
# Calculate the 95% confidence interval for RD
CI_RD_lower <- RD - 1.96 * SE_RD
CI_RD_upper <- RD + 1.96 * SE_RD
# Calculate the standard error for Relative Risk (RR)
SE_ln_RR \leftarrow sqrt((1 - P1) / (P1 * Total_Style1) + (1 - P2) / (P2 * Total_Style2))
# Calculate the natural log of Relative Risk (ln(RR))
ln RR <- log(RR)</pre>
# Calculate the 95% confidence interval for ln(RR)
CI_ln_RR_lower <- ln_RR - 1.96 * SE_ln_RR
CI_ln_RR_upper <- ln_RR + 1.96 * SE_ln_RR
# Calculate the 95% confidence interval for Relative Risk (RR)
CI_RR_lower <- exp(CI_ln_RR_lower)</pre>
CI_RR_upper <- exp(CI_ln_RR_upper)</pre>
CI_RD_lower <- round(CI_RD_lower, 4)</pre>
CI RD upper <- round(CI RD upper, 4)
CI RR lower <- round(CI RR lower, 2)
CI_RR_upper <- round(CI_RR_upper, 2)</pre>
cat("95% Confidence Interval for Risk Difference (RD):", "(", CI_RD_lower, ",", CI_RD_upper, ")" )
## 95% Confidence Interval for Risk Difference (RD): ( 0.0327 , 0.1077 )
cat("\n")
cat("95% Confidence Interval for Relative Risk (RR):", "(", CI_RR_lower, ",", CI_RR_upper, ")")
## 95% Confidence Interval for Relative Risk (RR): ( 1.63 , 6.47 )
```

Here are the calculated 95% confidence intervals for the Risk Difference (RD) and the Relative Risk (RR) associated with low birth weight (LBW) in babies of mothers following the two different lifestyle habits:

The hypothesis test for the risk difference is as follows:

Confidence Intervals:

- $H_0: RD = 0$, the risk is the same for both groups
- $H_a: RD \neq 0$, the risk is different for both groups

Risk Difference (RD) 95% CI: (0.0327, 0.1077) This interval suggests that the difference in the risk of LBW between mothers with unhealthy habits (Style 1) and those with healthy habits (Style 2) is between 3.27% and 10.77%. This significant difference emphasises the impact of lifestyle choices on LBW. The lower bound exceeding 0 indicates a statistically significant increased risk associated with unhealthy lifestyle habits, thus at a significance level of 0.05, we reject the null hypothesis.

The hypothesis test for the relative risk is as follows:

- $H_0: RR = 1$, the risk is the same for both groups
- $H_a: RR \neq 1$, the risk is different for both groups

Relative Risk (RR) 95% CI: (1.63, 6.47) This interval indicates that the risk of having a LBW baby for mothers with unhealthy habits is between 1.63 and 6.47 times greater than for mothers with healthy habits. The lower bound exceeding 1 suggests a statistically significant increased risk associated with unhealthy lifestyle habits, thus at a significance level of 0.05, we reject the null hypothesis.

Interpretation: These confidence intervals underscore the statistical significance and potential public health impact of lifestyle choices during pregnancy on birth weight. The findings advocate for interventions or education aimed at promoting healthier lifestyle choices among pregnant women to reduce the risk of LBW, which is associated with various adverse health outcomes in infants.

(c) Perform a chi-square test to determine whether there is a significant association between lifestyle habits and the birth weight of singleton babies.

Answer:

Using the table:

Lifestyle	LBW	NBW	Total
Style 1	34	301	335
Style 2	10	310	320
Total	44	611	655

Null Hypothesis (H_0) : There is no association between lifestyle habits and birth weight.

Alternative Hypothesis (H_a) : There is an association between lifestyle habits and birth weight.

The chi-square test works by:

Under the null hypothesis, which states that there is no association between the two categorical variables (lifestyle habits and birth weight), the expected frequency for each cell in the contingency table is calculated as:

$$E_{ij} = \frac{(\text{row total}_i \times \text{column total}_j)}{\text{total observations}}$$

where E_{ij} is the expected frequency for cell (i, j), and the row and column totals are the sums of the observed frequencies in the respective rows and columns.

The chi-square test statistic is calculated as:

$$\chi^2 = \sum \frac{(O_{ij} - E_{ij})^2}{E_{ij}}$$

where O_{ij} is the observed frequency for cell (i,j) and E_{ij} is the expected frequency for the same cell.

Alternatively, we can use the formula to calculate the chi-square (χ^2) statistic:

$$\chi^2 = \frac{(ad - bc)^2 \times (a + b + c + d)}{(a + b)(c + d)(a + c)(b + d)}$$

Where:

- a = 130 (Exposed cases)
- b = 200 (Exposed controls)
- c = 127 (Unexposed cases)
- d = 370 (Unexposed controls)

Next,

The degrees of freedom for a 2x2 contingency table are 1.

This is calculated as:

```
degrees of freedom = (number of rows -1) × (number of columns -1)
```

The p-value is then determined based on the chi-square statistic and the degrees of freedom using a chi-square distribution table (built-in to R chi-square test function).

Let's perform the chi-square test to determine if there is a significant association between lifestyle habits and the birth weight of singleton babies.

To assess the association between lifestyle habits and the birth weight of singleton babies using a chi-square test, we'll use a contingency table with the counts of low birth weight (LBW) and normal birth weight (NBW) for both Style 1 and Style 2 mothers. First, we need to determine the counts for the normal birth weight (NBW) babies:

```
Style 1 NBW: Total mothers in Style 1 - LBW babies in Style 1 = 335 - 34
```

Style 2 NBW: Total mothers in Style 2 - LBW babies in Style 2 = 320 - 10

Next, we'll create a contingency table with the following structure:

```
Style 1: LBW: 34, NBW for Style 1 Style 2: LBW: 10, NBW for Style 2
```

The chi-square test will then be used to determine if there is a statistically significant association between lifestyle habits (Style 1 vs. Style 2) and the incidence of LBW. We will use the observed frequencies to compute the chi-square statistic and p-value.

```
# Reference:
# https://libguides.library.kent.edu/SPSS/ChiSquare
# Haviland M. G. (1990). Yates's correction for continuity and the analysis of 2 x 2 contingency tables
# https://imaging.mrc-cbu.cam.ac.uk/statswiki/FAQ/yates
# https://www.ncl.ac.uk/webtemplate/ask-assets/external/maths-resources/business/hypothesis-tests/chi-s
# https://vula.uct.ac.za/access/content/group/9c29ba04-b1ee-49b9-8c85-9a468b556ce2/DOH/Module%202%20(Bi
# Calculate the counts of normal birth weight (NBW) babies for each lifestyle
NBW_Style1 <- Total_Style1 - LBW_Style1</pre>
NBW_Style2 <- Total_Style2 - LBW_Style2</pre>
# Create a 2x2 contingency table
contingency_table <- matrix(c(LBW_Style1, NBW_Style1, LBW_Style2, NBW_Style2), nrow = 2, byrow = TRUE)</pre>
# Perform the chi-square test for independence
chi2_test <- chisq.test(contingency_table, correct = FALSE)</pre>
# Extract the test statistic and p-value
test statistic <- chi2 test$statistic
p_value <- chi2_test$p.value</pre>
```

```
test_statistic <- round(test_statistic, 2)
p_value <- format(p_value, scientific = TRUE, digits = 2)

cat("Chi-square Test Statistic:", test_statistic)

## Chi-square Test Statistic: 12.89

cat("\n")</pre>
```

```
cat("P-value:", p_value)
```

P-value: 3.3e-04

The results of the chi-square test are as follows:

Chi-square statistic (Without Yates' correction): 12.89

P-value: 3.3e-04

Interpretation:

The chi-square test statistic of 12.89 with a p-value of approximately 3.3e-04 suggests a statistically significant association between lifestyle habits and the birth weight of singleton babies. This result indicates that the lifestyle habits (Style 1 being unhealthy habits and Style 2 being healthy habits) are significantly associated with the incidence of low birth weight in babies. Since the p-value is less than 0.05, at a significance level of 0.05, we reject the null hypothesis that there is no association between lifestyle habits and low birth weight. This supports the finding that maternal lifestyle habits have a notable impact on birth outcomes.