# hw\_06.Rmd

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#### Q1

Fit a linear model to the given data

```
# Given data
x = c(110.5, 105.4, 118.1, 104.5, 93.6, 84.1, 77.8, 75.6)
y = c(5.755, 5.939, 6.010, 6.545, 6.730, 6.750, 6.899, 7.862)
# Given equation corresponds to basic linear regression model.
# To fit the model
linear_model <- lm(y ~ x)</pre>
# Checking the summary of the model results
summary(linear_model)
##
## Call:
## lm(formula = y \sim x)
## Residuals:
                  1Q
                     Median
                                    3Q
## -0.34626 -0.27605 -0.09448 0.27023 0.53495
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 10.137455
                           0.842265 12.036
                                               2e-05 ***
               -0.037175
## x
                           0.008653 -4.296 0.00512 **
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.3624 on 6 degrees of freedom
## Multiple R-squared: 0.7547, Adjusted R-squared: 0.7138
## F-statistic: 18.46 on 1 and 6 DF, p-value: 0.005116
```

a) Least squares estimates of the slope.

Interpretation: For each additional unit increase in plant height, the estimated change in grain yield is approximately equal to  $(\hat{\beta}_1)$  units. The sign of  $(\hat{\beta}_1)$  indicates the direction of the relationship. If  $(\hat{\beta}_1)$  is positive, it suggests a positive correlation, meaning higher plant heights are associated with higher grain yields. If  $(\hat{\beta}_1)$  is negative, it suggests a negative correlation.

#### b) Perform F-test and then T-test.

## Coefficients:

## (Intercept) 10.137455

-0.037175

##

## x

## ---

```
# First let us do a F-test. F-test can be conducted by using the idea of ANOVA
anova(linear_model)
## Analysis of Variance Table
##
## Response: y
##
            Df Sum Sq Mean Sq F value
                                          Pr(>F)
## x
              1 2.42357 2.42357 18.455 0.005116 **
## Residuals 6 0.78794 0.13132
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
# Now we can check for T-test, Summary of the model-fitting will have the p-values which can be used to
summary(linear_model)
##
## Call:
## lm(formula = y \sim x)
##
## Residuals:
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -0.34626 -0.27605 -0.09448 0.27023 0.53495
##
```

2e-05 \*\*\*

Estimate Std. Error t value Pr(>|t|)

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

## Residual standard error: 0.3624 on 6 degrees of freedom
## Multiple R-squared: 0.7547, Adjusted R-squared: 0.7138
## F-statistic: 18.46 on 1 and 6 DF, p-value: 0.005116

0.842265 12.036

0.008653 -4.296 0.00512 \*\*

In both the cases, F-test and T-test, p value is less than 0.05 which provides evidence to reject the null hypothesis of  $H_0: \beta_1 = 0$ . This suggests that there is a significant linear relationship between the predictor variable (plant height) and the response variable (grain yield).

c) Construct a 95% CI by hand and compare to what R gives.

```
# ALpha value is given
alpha <- 0.05
n <- length(x)
# Extracting the values from the summary of model fitting
intercept_estimate <- coef(linear_model)[1]</pre>
SE_intercept <- summary(linear_model)$coefficients[1, "Std. Error"]
# Getting critical t value
t_critical <- qt(alpha/2, n-2)
# Calculting the intereval as upper and lower boundry
lower_bound <- intercept_estimate - t_critical * SE_intercept</pre>
upper_bound <- intercept_estimate + t_critical * SE_intercept</pre>
# Displaying the results
lower_bound # Calculated by hand
## (Intercept)
       12.1984
upper_bound # Calcualted by hand
## (Intercept)
      8.076507
confint(linear_model) # Calculated by R
##
                      2.5 %
                                 97.5 %
## (Intercept) 8.07650745 12.19840320
## x
               -0.05834895 -0.01600043
```

### d) Raw residuals.

```
# Raw residuals can be extracted from the model summary
residuals <- residuals(linear_model)

residuals

## 1 2 3 4 5 6 7

## -0.2746519 -0.2802428 0.2628757 0.2922999 0.0720958 -0.2610638 -0.3462643

## 8

## 0.5349514
```

#### e) Estimate of the error variance.

```
# The estimate of the error variance is obtained as the mean squared residual from the regression model
# We can obtain the same in R with following code
# Calculate the estimate of the error variance
error_variance <- sum(residuals^2) / (length(x) - 2)
# Display the result
error_variance
## [1] 0.1313228</pre>
```

#### f) Expected yield of the rice variety

```
# Given values
x_0 <- 100
alpha <- 0.05

# Calculate the expected yield
expected_yield <- coef(linear_model)[1] + coef(linear_model)[2] * x_0

# Calculate the standard error of the predicted values
SE_expected_yield <- sqrt(error_variance * (1/n + (x_0 - mean(x))^2 / sum((x - mean(x))^2)))

# Calculate the critical t-value
t_critical <- qt(alpha/2, length(x) - 2)

# Calculate the confidence interval
lower_bound <- expected_yield - t_critical * SE_expected_yield
upper_bound <- expected_yield + t_critical * SE_expected_yield

# Display the results
expected_yield</pre>
```

```
## (Intercept)
## 6.419986

lower_bound

## (Intercept)
## 6.743651

upper_bound

## (Intercept)
## 6.096321
```

## g) Prediction of the yield of new rice variety

```
# Calculate the standard error of the prediction
SE_prediction <- sqrt(error_variance * (1 + 1/n + (x_0 - mean(x))^2 / sum((x - mean(x))^2)))
# Calculate the prediction interval
lower_bound_prediction <- expected_yield - t_critical * SE_prediction
upper_bound_prediction <- expected_yield + t_critical * SE_prediction

# Display the results
lower_bound_prediction

## (Intercept)
## 7.363934

upper_bound_prediction

## (Intercept)
## 5.476038</pre>
```

Comparing the results from f, new variety of rice has a wider 95% prediction interval.

## h) Compute R2 and interpret the results

```
R_squared <- summary(linear_model) r.squared
R_squared
```

## [1] 0.7546518

Interpretation: A higher r square suggests that the linear regression model does a good job of explaining the variability in grain yield based on plant height.

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#### $\mathbf{Q2}$

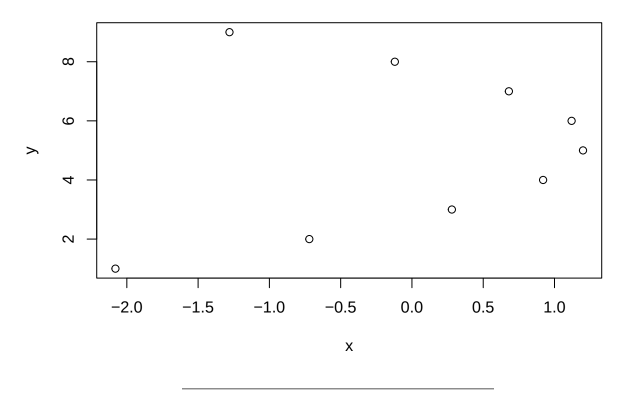
#### Answers

```
# Given artificial data
x \leftarrow c(1, 2, 3, 4, 5, 6, 7, 8, 9)
y \leftarrow c(-2.08, -0.72, 0.28, 0.92, 1.20, 1.12, 0.68, -0.12, -1.28)
# Fitting a linear model
linear_model <- lm(y ~ x)</pre>
summary(linear_model)
##
## Call:
## lm(formula = y \sim x)
## Residuals:
## Min 1Q Median 3Q
                                 Max
## -1.68 -0.42 0.48 1.02
                                1.20
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.5000
                         0.8674 -0.576
                                              0.582
                0.1000
                            0.1541 0.649
                                              0.537
## Residual standard error: 1.194 on 7 degrees of freedom
## Multiple R-squared: 0.05672, Adjusted R-squared: -0.07804
## F-statistic: 0.4209 on 1 and 7 DF, p-value: 0.5372
```

#### a) Plot y vs x

```
plot(y, x, main = "Scatterplot of y vs. x", xlab = "x", ylab = "y")
```

# Scatterplot of y vs. x

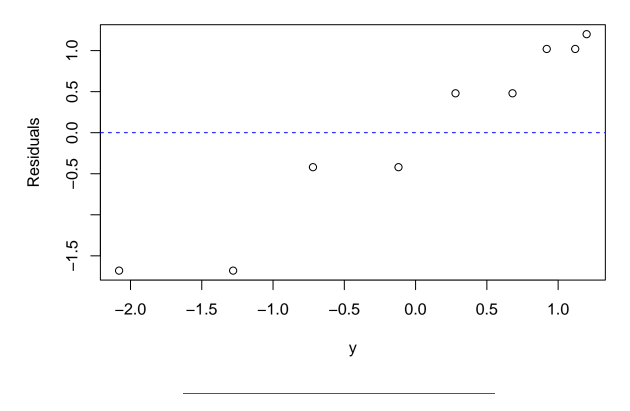


# b) Plot the raw residuals vs. y

```
raw_residuals <- residuals(linear_model)

plot(y, raw_residuals, main = "Residuals vs. y", xlab = "y", ylab = "Residuals")
abline(h = 0, col = "blue", lty = 2)</pre>
```

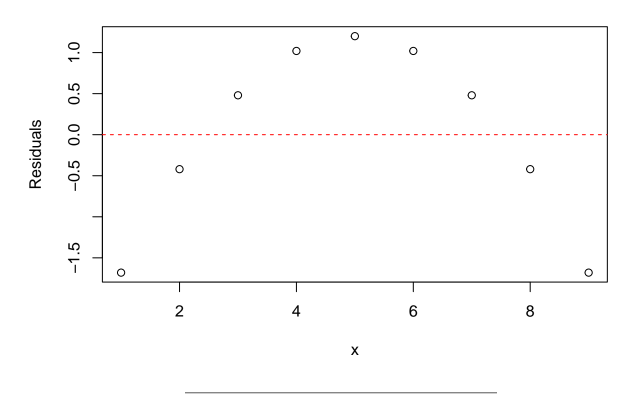
# Residuals vs. y



# c) Plot raw residuals vs $\mathbf{x}$

```
# Plot residuals against x
plot(x, raw_residuals, main = "Residuals vs. x", xlab = "x", ylab = "Residuals")
abline(h = 0, col = "red", lty = 2)
```

### Residuals vs. x

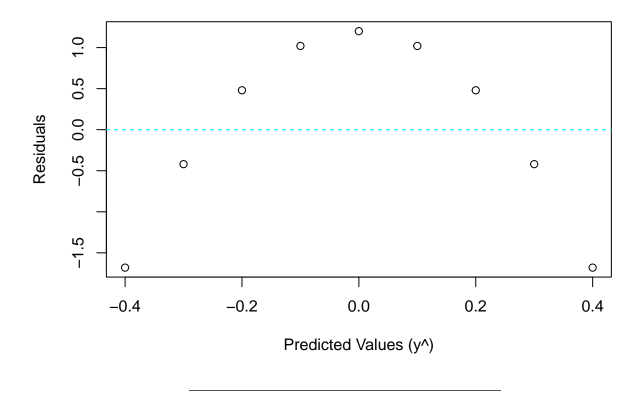


# d) plot raw residuals vs y^

```
predicted_values <- predict(linear_model)

plot(predicted_values, raw_residuals, main = "Residuals vs. Predicted Values", xlab = "Predicted Values abline(h = 0, col = "cyan", lty = 2)</pre>
```

#### **Residuals vs. Predicted Values**



### e) Which explains the better model fit?

While (b) and (c) provide valuable information, (d) Residuals vs. y^ gives a better indication of the lack of fit as it directly assesses the performance of the model in predicting the response variable y. If there is a pattern or trend in (d), it suggests that the linear model might not be appropriate for capturing the underlying relationship in the data.