

**STAT1006**

**Regression and Nonparametric Inference**

**Semester 2, 2024**

**Final Project Report**

**LINEAR REGRESSION ANALYSIS FOR SULFUR DIOXIDE CONTENT IN THE ATMOSPHERE**

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Bachelor of Science (Data Science)

**Declaration**

The work presented in this report is my own work and all references are duly acknowledged.

This work has not been submitted, in whole or in part, in respect of any academic award at Curtin University or elsewhere.



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# Introduction

This study employs simple machine learning techniques, i.e., linear regression analysis, to investigate a dataset containing various factors, with the aim of coming up with the best predictive model to predict the amount of SO₂ content in the air from these factors. SO₂ levels are used as an indicator of the amount of air pollution in many urban areas, and high amounts of SO₂ in the air generally equate to high levels of air pollution, which can be harmful to people’s health and the environment at large with possible respiratory problems and the formation of acid rain. This study would be helpful in understanding how different ecological and human factors affect the amount of air pollution and can help environmental scientists, among others, to plan ahead and put measures in place to control the amount of SO₂ emissions in urban areas.

The study aims to find out which of these factors contribute the most to the increase in SO₂ emissions, how well can we predict the amount of SO₂ content in the air with these explanatory factors, and which linear regression model would be best suited for this task. The report will cover the exploratory analysis of the dataset, simple linear regression with only one explanatory variable, and multiple linear regression with multiple explanatory variables and the SO₂ content as the target variable. Lastly, there will be a comparison of the predictive abilities of the different models to decide which one is the best predictive model.

The dataset is relatively small compared to the ideal size for regression and other machine learning applications, as it only contains forty one observations factored into seven variables – SO₂ , manu, temp, popul, wind, precip, predays, and Region:

1. **SO₂ :** mean amount of sulphur dioxide in the air (mg/cm3 ). This is the variable we are trying to predict, i.e., the response variable.
2. **Temp:** mean annual temperature (in Fahrenheit)
3. **Manu:** number of manufacturing factories with 20 or more workers
4. **Popul:** the population size in thousands as of 1970 census
5. **Wind:** mean annual wind speed (in miles per hour)
6. **Precip:** mean annual amount of precipitation (in inches)
7. **Predays:** the average number of days with precipitation annually
8. **Region:** the region in question (in USA)

# Exploratory Data Analysis (EDA)

To begin with, I checked the dataset for any issues that might need to be cleaned. All of the data entries were correct with no inconsistent entries or missing values in any of the columns, the variables had the right data types, and there were no duplicate records. I then conducted some exploratory data analysis to get a better understanding of the dataset and the variables. Firstly, I opened the dataset and a quick glimpse into the first six observations of the dataset also revealed the data types for each variable. All of the variables were numerical except Region, which was a categorical variable containing the region of each observation. I then checked the five-number summaries of the variables to gain some insight into the spread (quartiles), location (median), and range (minimum and maximum values) of each of the variables. Variable ‘manu’ had the largest range, and ‘popul’ had the largest mean value. I could also see how the variables would spread if they were plotted on a boxplot from the minimum value, the first quartile, the median (second quartile), third quartile, and the maximum value. Seeing as Region was the only categorical variable in the dataset, I wanted to see how the SO₂ levels vary by region, so I plotted the histograms for SO₂ by Region. The amount of SO₂ content in the air was highest in the East, followed by North, South, then the West, in descending order as shown in Fig. 1 below.

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**Fig. 1: Histogram of SO₂ content by Region**

To get a better look at how the values were spread and possible outliers for the SO₂ values in each region, I plotted boxplots of the two variables. I could see that the SO₂ values in the East were right-skewed with no outliers, left-skewed in the North with two possible outliers, slightly right-skewed in the South with three possible outliers, and left-skewed in the West with one possible outlier, as shown in Fig.2 below.

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**Fig. 2: Boxplot of SO₂ content by Region**

I then executed a function to check for the number of possible outlier observations in each variable in the original dataset and found that SO₂ had 3, temp had 1, manu had 4, popul had 3, wind had no outliers, precip had 2, and predays had 3. Since the dataset was very small, I was able to manually check each of the outliers in each variable and confirmed that they were genuine datapoints, they just happened to be larger compared to the other observations in their respective variables. I therefore opted not to drop them. However, later on in the analysis, I did try to perform some transformations to see if they could help alleviate the influence of these potential outliers on the final model.

Lastly, to check for the association and correlation between the variables, I plotted the correlation plot. This provided great insight into which variable had the strongest correlation with the target variable (SO₂ ) for the subsequent simple linear regression model, the level of multicollinearity in the dataset, i.e., which variables had very strong correlations between themselves, and the nature of the relationship between the different variables. I also got the Pearson’s correlation coefficient values for the correlation between each of the variables, and with the SO₂ target variable, which was crucial in deciding which variable would be best appropriate for modelling the simple linear regression model. The correlation plot is shown in Fig. 3 below.

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**Fig. 3: Correlation plot**

From the plot above, I could see that many of the variables did not fulfill the assumption of linearity with the response variable, SO₂. I explored different transformations to try and improve the linearity of the variable relationships, stabilize the variances, and possibly improve the normality of the data. I applied log, square root and cube root transformations to both the explanatory and response variables, for the simple linear model, and to all variables for the multiple linear models, but this only made the variables less linear and affected the models’ performances, so I opted to proceed with the dataset as it was, with the aim of using the cross validation to deal with this issue.

# Linear Regression modelling

With any machine learning model, such as the linear regression model in this report, it is advisable to employ model validation methods to ensure that the model generalizes well to new, unseen data, which helps avoid overfitting or underfitting. Model validation also helps assess the model’s predictive performance on the unseen datasets, ensuring that the relationships identified by the model on the training data are also identified in other datasets. There are a few methods of model validation, including splitting the original dataset into a training and testing split, cross validation (K-fold or Leave One Out method), bootstrapping, and examining the model performance metrics. These methods are generally applicable in most cases, but certain factors, such as the size of the dataset, the purpose of your analysis, and the computational resources available to you, can sometimes determine which methods you have to use. Since the dataset in this report is very small (only 41 observations), it would be better to use cross validation methods like k-fold instead of splitting the dataset, which would significantly reduce the data size for training the model.

Therefore, for my analysis in this report, I employed K-fold cross validation for both the simple and multiple linear regression models. This made sure that each data point, shuffled across different folds, is used for both training and validating the model, and also reduced the variance in performance estimation as it averages the performance across different folds and multiple repeats to give a more reliable performance measure. I defined a function, ‘cv\_error’, that would create 10 folds from all the observations in the dataset to make sure we had enough training data. This function would then split the data into training and testing sets for all the folds, fit the linear model on the training split, perform prediction on the test split, and calculate the mean squared error (MSE) for each fold. I also define the function ‘repeated cv\_error’ which would then repeat this process five times to get a range of possible MSE values, and then return the average of all MSE values. Getting the average of many repeats would make the results more reliable and provide a better assessment of each of the models’ predictive abilities.

## Simple Linear Regression

For the simple linear model, I was looking for coefficients of the model that would fit the formula **y^ (y hat) = a + bx**, where **b** is the slope of the least squares line, and **a** is the intercept, i.e., where the least squares line crosses the y-axis, or the value of y when x is 0.

Following the exploratory analysis, I identified that the variable ‘manu’ had the highest correlation with the target variable, ‘SO₂ ’, and would therefore be the best predictor for the simple linear model. I created the simple linear model with ‘SO₂ ’ as the y variable and ‘manu’ as the x variable and plotted the model with the line of best fit as shown in Fig. 4 below.

A graph with a line and dots

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**Fig. 4: Simple Linear model with least squares line**

**Results**

From the summary and visual plot in Fig. 4, I could see that the equation of the least squares line was equal to **y^(y hat) = 17.61 + 0.03x**. In other words:

**SO₂ = 17.61 + 0.03 manu**

The slope of the line, b, is 0.03. This means that each additional manufacturing enterprise with 20 or more workers is predicted to increase the SO₂ content in the air by 0.03 micrograms per cubic meter. The positive coefficient for ‘manu’ also showed that there is a positive association between ‘manu’ and SO₂ levels. The intercept, a, is 17.61. This means that when the number of manufacturing enterprises is zero, the SO₂ content in the air is 17.61 micrograms per cubic meter.

The R-squared (R^2) value for the model was 0.4157, which means that around 42% of the variability in the SO₂ levels was explained by the predictor variable , ‘manu’. The F-statistic, 27.75 with p-value 5.36×10−6 indicates the model’s overall significance and this value shows that the model as a whole was a better fit than just modelling with the intercept alone. The five-number summary of the residuals showed some variability in prediction errors, with the comparatively large maximum residual value, 67.18, indicating probable outlier effect or some potentially large deviations that could affect the performance of the model.

**Diagnostics checking**

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Fig. 5: Diagnostic plots for the Simple Linear Regression model

1. (TOP LEFT: Residuals vs Fitted): There seems to be a slightly upward trend, which might indicate that the variance of residuals is not constant, thus violating the homoscedasticity assumption.

2. (BOTTOM LEFT: Scale Location): There is a slightly upward trend, suggesting non-constant variance in the residual errors, thus violating the homoscedasticity assumption.

3. (TOP RIGHT: Normal Q-Q): Most the points fall approximately along this reference line, except a small minority of potential outliers on the top end, so we can assume normality.

4. (BOTTOM RIGHT: Residuals vs Leverage): The plot shows that there are a few points with relatively higher leverage, which could be potential influential outliers.

## Multiple Linear Regression

I then conducted multiple linear regression analyses with 1) all subsets, 2) forward stepwise, and 3) backward stepwise selection methods to compare how well each one would perform and to identify which one could best predict the SO₂ content in the air given multiple explanatory variables.

### All subsets

The goal for this model was to evaluate all possible combinations of explanatory variables to determine which combination had the best predictive performance. This method is typically more extensive as it exhausts all possible combinations to find the most optimal one for the prediction of the target variable. I then used the Mean Square Error (MSE) values calculated using the k-fold cross validation function to assess the predictive performance of each combination to determine how many predictor variables would have the best prediction of SO₂ content in the air. The MSE values for each combination of variables were as follows:

|  |  |
| --- | --- |
| **Number of predictors** | **MSE value** |
| 1 | 319.221 |
| 2 | 256.577 |
| 3 | 257.116 |
| 4 | 258.751 |
| 5 | 226.602 |
| 6 | 252.179 |

**Table 1: All Subsets MSE values**

As a general rule, the lower the MSE value, the better the model is at predicting the target variable. Therefore, from Table 1 above, the model with 5 predictors seemed to be the best model, as it has the lowest MSE value. The table also showed that the model with just a single predictor has the worst performance, which was in line with the MSE value for the simple linear regression model discussed later in the report.

### Backward stepwise

I then proceeded to create a multiple linear model which starts off fitted with all possible predictors and removes them one by one using the AIC (Akaike Information Criteria) to inform the algorithm when to stop the removal of predictors. The AIC helps ensure there is a balance between model complexity and model fit by penalizing models with more predictor variables, thus reducing possible overfitting and prioritizing models with lower AIC values. The model started with all 6 numerical predictor variables, and using the AIC, was able to reduce the number of predictor variables to 5.

**Results**

From the summary of the model, I could see that the formula, **y^ (y hat) = 100.15 – 1.12x1 + 0.06x2 – 0.04x3 – 3.08x4 + 0.42x5**, was equal to:

**SO₂ = 100.15 – 1.12 temp + 0.06 manu – 0.04 popul – 3.08 wind + 0.42 precip**

* The intercept is 100.15, which means when all predictor variables are equal to zero, the SO₂ content in the air is 100.15 micrograms per cubic meter.
* For each Fahrenheit increase in the average annual temperature, the SO₂ content in the air decreases by 1.12 micrograms per cubic meter.
* The SO₂ content in the air increases by 0.06 micrograms per cubic meter for each additional manufacturing enterprise employing 20 or more workers.
* For each additional thousand people increase in population size, the SO₂ content in the air is predicted to decrease by 0.039 micrograms per cubic meter.
* For each additional mph increase in average annual wind speed, the SO₂ content in the air decreases by 3.08 micrograms per cubic meter. However, the relationship between the two variables is marginally significant as shown by the p-value, suggesting a weak association between them.
* Lastly, the SO₂ content in the air is predicted to decrease by 0.419 micrograms per cubic meter for each additional inch in the average annual precipitation.

The R-squared (R^2) value showed that the backward stepwise model was able to explain approximately 66.85% of the variability in the SO₂ content in the air. The final model includes the predictors temp, manu, popul, wind, and precip, with ‘manu’ showing the strongest positive association with SO₂. The F-statistic, 14.12 with p-value 1.41×10−7 also showed that the model was statistically significant and indicates that the chosen predictor variables explain a significant proportion of the variability in SO₂ content in the air.

**Diagnostics checking**

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Fig. 6: Diagnostic plots for Backward stepwise Multiple Linear model

1. (TOP LEFT: Residuals vs Fitted): There seems to be a downward, then upward trend, which might indicate that the variance of residuals is not constant, thus violating the homoscedasticity assumption.

2. (BOTTOM LEFT: Scale Location): There seems to be a upward, then downward trend, suggesting non-constant variance in the residual errors, thus violating the homoscedasticity assumption.

3. (TOP RIGHT: Normal Q-Q): Most the points fall approximately along this reference line, except a small minority of potential outliers on the top end, so we can assume normality.

4. (BOTTOM RIGHT: Residuals vs Leverage): The plot shows that there are a few points with relatively higher leverage, which could be potential influential outliers.

### Forward stepwise

I then created a multiple linear model which starts off fitted with only the intercept, then incrementally adds predictors one by one using the AIC (Akaike Information Criteria) to inform the algorithm when to stop adding predictors to the model. The model started with only the intercept, and using the AIC, was able to add 3 predictor variables to arrive at the final model.

**Results**

From the summary of the model, I could see that the formula, **y^ (y hat) = 6.97 + 0.07x1 - 0.05x2 + 0.16x3**, was equal to:

**SO₂ = 6.97 + 0.07 manu - 0.05 popul + 0.16 predays**

* For each additional manufacturing enterprise employing 20 or more workers, the SO₂ content in the air increases by 0.07 micrograms per cubic meter.
* For each additional thousand people increase in population size, the SO₂ content in the air is predicted to decrease by 0.05 micrograms per cubic meter.
* Lastly, the SO₂ content in the air is predicted to increase by 0.16 micrograms per cubic meter for each additional day with precipitation annually.

The R-squared (R^2) value showed that the backward stepwise model was able to explain approximately 61.74% of the variability in the SO₂ content in the air. The final model includes the predictors manu, popul, and predays, with ‘manu’ showing the strongest positive association with SO₂. The F-statistic, 19.9 with p-value 7.54×10−8 also showed that the model was statistically significant and indicates that the chosen predictor variables explain a significant proportion of the variability in SO₂ content in the air.

## Predictive ability

In order to compare the predictive abilities of the models discussed above, I used the ‘cv\_error’ function I created earlier to create 10 folds from all the observations in the dataset, ensuring I had enough training data. This function would split the data into training and testing sets for all the folds, fit the linear model on the training split, perform prediction on the test split, and calculate the mean squared error (MSE) for each fold. For reliability of results, I then used the function ‘repeated cv\_error’, which repeated this process five times to get a range of possible MSE values, and then return the average of all MSE values. The average MSE values of the models were as follows:

|  |  |  |
| --- | --- | --- |
| **Model** | **MSE for single run** | **Average MSE for 5 runs** |
| Simple Linear | 348.513 | 330.879 |
| Backward stepwise | 242.549 | 239.482 |
| Forward stepwise | 241.113 | 247.772 |

**Table 2: Mean Square Error values for all models**

As mentioned earlier, the lower the MSE value, the better the model is at predicting the response variable given the list of predictor variables. Therefore, using the average MSE value for each model, I concluded that the multiple linear regression model with the backward stepwise method was the best predictor of SO₂ content in the air. The model was able to explain a significant amount of the variability in SO₂ values using 5 predictor variables – temp, manu, popul, wind, precip.

# Discussion

This study was able to provide insight into some of the key factors that affect the amount of SO₂ content in the air. The linear regression models used in the analyses were able to identify the main factors that influence the SO₂ levels, with the number of manufacturing enterprises with more than twenty workers emerging as a primary contributor, suggesting that the increase in industrial activity in a region coincides with increased levels of air pollution. Since I chose the backward stepwise model, this suggests that the number of manufacturing factories, the population size, and the number of days with precipitation annually have the most significant influence on the amount of SO₂ in the air, whether negative or positive.

There were, however, a few limitations of the study, including the small dataset, which limited the statistical power of the regression models and possibly affected how well the resulting model would generalize to new, unseen data. Additionally, the non-linearity of relationships between predictors and the target variable also affected the accuracy of the analysis. For future analyses, selecting a larger dataset would help to account for some of these issues, and if lack of normality is still an issue, consideration could be taken to use nonlinear models that are more robust to nonlinearity in data.

In conclusion, despite the limitations of the study, it still provides valuable insight into how different ecological and industrial factors affect the amount of SO₂ in the air and in turn affect air pollution, and it can be used to inform environmental scientists, governments, etc., on how to set in place regulations that would help curb the increase in air pollution. It also provides a guideline for which machine learning methods would be better suited for the purposes of predicting the amount of air pollution given a list of similar factors.

# Appendices (R Code)

knitr::opts\_chunk$set(echo = TRUE)  
# importing the necessary libraries   
library(dplyr)  
library(Hmisc)  
library(ggplot2)  
## opening the datasets  
pollution <- read.csv("Pollution.csv")  
View(pollution)  
## getting a glimpse of the dataset   
head(pollution)  
## five-number summary   
summary(pollution)  
## checking for missing values  
missing <- colSums(is.na(pollution))  
missing  
## histograms for SO2 levels by region  
ggplot(pollution, aes(x = factor(region), y = SO2)) +  
 geom\_col(fill = "darkgreen") +  
 labs(title = "SO₂ Levels by Region", x = "Region", y = "SO₂ Levels") +  
 theme\_minimal()

A graph of a number of levels

Description automatically generated

# boxplot to see the spread, median and potential outliers   
ggplot(pollution, aes(x = factor(region), y = SO2)) +  
 geom\_boxplot(fill = "darkgreen", color = "black") +  
 labs(title = "SO₂ Levels by Region", x = "Region", y = "SO₂ Levels") +  
 theme\_minimal()

A graph with green and black squares

Description automatically generated

## checking the correlation between numeric variables   
num\_pollution <- pollution[,c(-8)]  
pairs(num\_pollution) #shows the association between variables

A chart of a graph

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# corrplot   
library(PerformanceAnalytics) ## install this  
chart.Correlation(num\_pollution)

A graph of numbers and graphs

Description automatically generated with medium confidence

# Sample data: Assuming your data frame is called num\_pollution  
# Apply the function to each numeric column  
outlier\_counts <- sapply(num\_pollution, function(x) {  
 if (is.numeric(x)) {  
 Q1 <- quantile(x, 0.25, na.rm = TRUE)  
 Q3 <- quantile(x, 0.75, na.rm = TRUE)  
 IQR <- Q3 - Q1  
 lower\_bound <- Q1 - 1.5 \* IQR  
 upper\_bound <- Q3 + 1.5 \* IQR  
 sum(x < lower\_bound | x > upper\_bound) # Count outliers  
 } else {  
 NA # Skip non-numeric columns  
 }  
})  
  
# Remove any NA values from the result  
outlier\_counts <- outlier\_counts[!is.na(outlier\_counts)]  
  
# Display the count of outliers for each variable  
print(outlier\_counts)  
  
# Refined k-fold cross-validation function  
cv\_error <- function(formula, data, k = 10) {  
 #set.seed(123) #for reproducibility  
 n <- nrow(data)  
 folds <- sample(rep(1:k, length.out = n)) # Create folds  
 errors <- numeric(k) # To store MSE for each fold  
   
 for (i in 1:k) {  
 # Split data into training and testing sets based on fold  
 test\_index <- which(folds == i)  
 train\_data <- data[-test\_index, ]  
 test\_data <- data[test\_index, ]  
   
 # Fit the model on the training data  
 model <- lm(formula, data = train\_data)  
   
 # Predict on the test data  
 predictions <- predict(model, newdata = test\_data)  
   
 # Calculate mean squared error for this fold  
 errors[i] <- mean((test\_data$SO2 - predictions)^2)  
 }  
   
 # Return average cross-validated MSE  
 mean(errors)  
}  
  
# New function to perform repeated k-fold cross-validation and average the MSE  
repeated\_cv\_error <- function(formula, data, k = 10, repeats = 5) {  
 set.seed(123) # For reproducibility across multiple rounds  
 mse\_values <- numeric(repeats)  
   
 for (i in 1:repeats) {  
 mse\_values[i] <- cv\_error(formula, data, k)  
 }  
   
 mean(mse\_values) # Return the average MSE over all repeats  
}  
  
simple\_model <- lm(SO2~manu, data=num\_pollution)  
summary(simple\_model)  
# Run k-fold cross-validation on a simple linear model with SO2 ~ manu  
simple\_mse <- cv\_error(SO2 ~ manu, data = num\_pollution, k = 10)  
print(paste("Cross-validated MSE for Simple Linear Regression:", round(simple\_mse, 3)))  
  
# getting the average MSE for 10 folds and 5 repeats  
simple\_avgmse <- repeated\_cv\_error(SO2~manu, data = num\_pollution, k = 10, repeats = 5)  
print(paste("Average MSE for Simple Model:", round(simple\_avgmse, 3)))  
# loading the necessary pakckages for plotting SLR line of best fit  
library(ggplot2)  
library(ggpubr)  
  
# Plot the data using ggplot2 and include the regression line with the equation  
ggplot(num\_pollution, aes(x = manu, y = SO2)) +  
 geom\_point(color = rgb(0.5, 0, 1, alpha = 0.4), size = 3) + # Scatter plot  
 geom\_smooth(method = "lm", se = FALSE, color = "blue", linetype = "solid") + # Regression line  
 labs(x = "Manufacturers", y = "SO2 Levels") + # Axis labels  
 theme\_minimal() # for a cleaner look

## `geom\_smooth()` using formula = 'y ~ x'

A graph with dots and numbers

Description automatically generated

# diagnostics checking  
par(mfrow = c(2, 2)) # Set up for 4 plots  
plot(simple\_model)

A group of graphs showing different values

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# load the package  
library(leaps)  
  
# Fit all possible models  
all\_subsets <- regsubsets(SO2 ~ temp + manu + popul + wind + precip + predays, data = num\_pollution, nvmax = 6)  
#summary(all\_subsets)  
# loading required package  
if (!requireNamespace("leaps", quietly = TRUE)) install.packages("leaps")  
library(leaps)  
  
# Cross-validate each model in subset selection  
mse\_values <- sapply(1:6, function(i) {  
 # Build formula dynamically with the best predictors of the subset  
 predictors <- names(coef(all\_subsets, i))  
   
 # Exclude intercept from the predictors  
 if (length(predictors) > 1) {  
 formula <- as.formula(paste("SO2 ~", paste(predictors[-1], collapse = " + ")))  
 } else {  
 formula <- as.formula("SO2 ~ 1") # Only intercept model  
 }  
   
 # Perform cross-validation and get MSE  
 cv\_error(formula, data = num\_pollution, k = 10)  
})  
  
# Display cross-validated MSE for each model  
mse\_results <- data.frame(Num\_Predictors = 1:6, MSE = round(mse\_values, 3))  
print(mse\_results)  
  
# Fit the full model with all predictors  
full\_model <- lm(SO2 ~ temp + manu + popul + wind + precip + predays, data = num\_pollution)  
  
# Perform backward selection  
backward\_model <- step(full\_model, direction = "backward")  
  
# Cross-validate the backward selection model  
backward\_mse <- cv\_error(formula(backward\_model), data = num\_pollution, k = 10)  
print(paste("Cross-validated MSE for Backward Selection Model:", round(backward\_mse, 3)))  
  
# getting the average MSE for 10 folds and 5 repeats  
bwd\_avgmse <- repeated\_cv\_error(formula(backward\_model), data = num\_pollution, k = 10, repeats = 5)  
print(paste("Average MSE for Backward Model:", round(bwd\_avgmse, 3)))  
# bw model coefficients  
summary(backward\_model)  
# diagnostics checking  
par(mfrow = c(2, 2)) # Set up for 4 plots  
plot(backward\_model)

A group of graphs showing different values

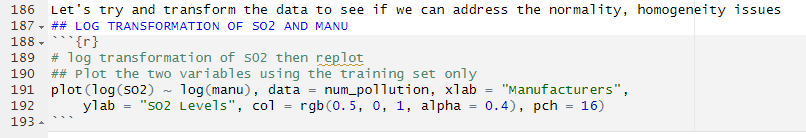
Description automatically generated

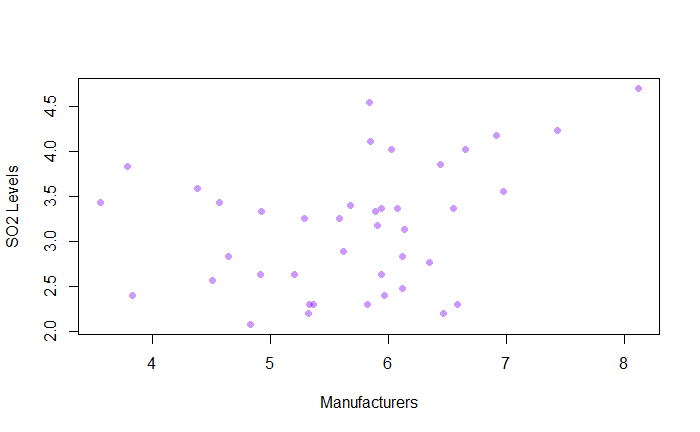
# Fit a minimal model (intercept only)  
minimal\_model <- lm(SO2 ~ 1, data = num\_pollution)  
  
# Perform forward selection  
forward\_model <- step(minimal\_model, direction = "forward", scope = formula(full\_model))  
  
# Cross-validate the forward selection model  
forward\_mse <- cv\_error(formula(forward\_model), data = num\_pollution, k = 10)  
print(paste("Cross-validated MSE for Forward Selection Model:", round(forward\_mse, 3)))  
  
# getting the average MSE for 10 folds and 5 repeats  
fwd\_avgmse <- repeated\_cv\_error(formula(forward\_model), data = num\_pollution, k = 10, repeats = 5)  
print(paste("Average MSE for Forward Model:", round(fwd\_avgmse, 3)))  
# fw model coefficients   
summary(forward\_model)  
# diagnostics checking  
par(mfrow = c(2, 2)) # Set up for 4 plots  
plot(forward\_model)

A group of graphs showing different values

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Failed attempt with transformation (sqrt and cube root transformations performed just as poorly).





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Description automatically generated

A screenshot of a computer

Description automatically generated