Monte Carlo simulation

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1 Introduction

Regression is a tool in statistics used to understand the relationship between one dependent variable and one or more independent variables. For regression to be both accurate and reliable, a set of assumptions needs to be met. These assumptions are independence of errors, linearity, homoscedasticity, normality of errors, multicollinearity, and correct model specifications. If these assumptions are not met, it can introduce bias and inaccuracies in the regression model.

1.1 Independence of Errors

This assumption states that the errors from the model are not correlated with each other but are independent. This means that one error cannot be used to predict the next one.

1.2 Linearity

This assumption states that the relationship between the independent variables and the parameters, also known as the coefficients, is linear. This does not mean that the regression model itself has to be linear. For example, in polynomial regression, the relationship between the independent variables and the coefficients is still linear, but the independent variables can be transformed using powers.

1.3 Homoscedasticity

This is the assumption of constant variance across errors for all levels of the independent variables.

1.4 Normality of Errors

This assumption states that the errors between the model and the observed values, also called residuals, are normally distributed. If this is not met, it may result in a biased model and a worse model fit.

1.5 Multicollinearity

This occurs when two or more independent variables in a regression model are highly correlated with each other. This means that changes in one independent variable are associated with changes in another, making it difficult to determine the individual effect of each independent variable on the dependent variable.

1.6 Correct Model Specifications

This assumes that the provided dependent variables for the models are the correct ones. If one is missing or the model is overfitting, this may result in incorrect coefficients and introduce errors into the model.

2 Statistical theory

2.1 Probability space

The **sample space**, *S*, is the set of all possible outcomes.

If a sample space contains a finite number of possibilities or an unending sequence with as many elements as there are whole numbers, it is called a **discrete** sample space.

Example: When rolling a standard six-sided die form the discrete sample space, the possible outcomes are S = 1, 2, 3, 4, 5, 6

If a sample space contains an infinite number of possibilities equal to the number of points on a line segment, it is called a **continuous sample space**. Example: Measuring the heights of people in a population. This is a continuous sample space, because height can take any real value within a given range. An **event** is a subset, $A \subseteq S$, of the sample space. The event is the amount that contains all possible events. An example of a discrete event could be rolling a die and getting an uneven number, this would be the event A = 1, 3, 5. For the continuous event, it could be that a person is between 160 cm and 170 cm tall.

The probability of an event A, P(A), is the sum of the weights of all sample points in A. The probability of the whole sample space is 1, P(S) = 1 The probability of any event being between 0 and 1,0 < P(A) < 1 The probability of the empty set being 0, $P(\emptyset) = 0$

Probability of mutually exclusive events If A and B are mutually exclusive, $A \cap B = \emptyset$, then $P(A \cup B) = P(A) + P(B)$

Where A and B never occur at the same time, so their union is equal to the two events added together.

Probability of union

$$P(A \cup B) = P(A) + P(B) - P(A \cap B)$$

Here, the union of the two events is A added to B, but minus their common event, since it otherwise would be added twice.

Two events A and B are independent, if

$$P(A|B) = P(A)$$

The equivalent definition to this is:

Two events A and B are independent if and only if

 $P(A \cap B) = P(A)P(B)$ This says that the probability of both event A and B happening, is equal to the product of the two events.

2.2 Random variables

test A random variable is defined as a function that associates a real number with each element in the sample space. We use capital letters to denote a random variable, for example X, and then the corresponding small letter, in this case x, for one of its values. As an example we roll a dice 3 times, which gives us a sample space of the different combinations. Each point in the sample space gets a numerical value assigned between 0 and 3. These values are random quantities and are assumed by the random variable X, which for example could assume the amount of 5's rolled. In that case X(5,1,2)=1 and X(3,6,1)=0.

A random variable X can be discrete, which means that its set of possible outcomes is countable. The dice example is a discrete random variable, because you can count how many times 5 is rolled. The outcomes of some statistical experiments may be neither finite nor countable. For example when something is measured such as temperature or speed where the set of possible values is an entire interval of numbers, it is not discrete. The random variable X then takes values on a continues scale, which therefore is called a continuous random variable.

2.2.1 Discrete random variable

A discrete random variable assumes each of its values with a certain probability. Frequently, it is convenient to represent all the probabilities of a random variable X by a formula. Let X be a discrete random variable which can take the values $x_1, x_2, ...$ Then the distribution of X is given by the probability function:

$$f(x_i) = P(X = x_i), \quad i = 1, 2, ...$$

For a discrete random variable this function is also called the probability mass function, where following holds for each possible outcome x:

- P(X = x) = f(x).
- f(x) > 0,
- $\sum_{x} f(x) = 1$.

In addition to the probability mass function f, the discrete random variable X also has a distribution function F(x) given by:

$$F(x) = P(X \le x) = \sum_{x_i \le x} f(x_i), \quad x \in \mathbf{R}.$$

This helps decide the probability that the random variable assumes a value equal to or smaller than x. Its sums up the probability mass functions's values.

The mean of a discrete variabel X, with a distribution function $f(x_i)$ is given by:

$$\mu = E(X) = \sum_{i} x_i P(X = x_i) = \sum_{i} x_i f(x_i).$$

The mean is typically the expected value. It is a weighed average of the possible values of X. The values are weighed by its probability in the sample space.

In addition to the mean, we should also mention the variance. The variance is the mean squared distance between the values of the variable and the mean value. It is given by:

$$\sigma^2 = E[(X - \mu)^2] = \sum_i (x_i - \mu)^2 f(x_i).$$

The variance indicates whether the values of X are far from the mean values or close. A high variance means that the values of X have a high probability of being far from the mean values and vice versa. Along with the variance, the standard deviation is also often used. It is given by the square root of the variance:

$$\sigma = +\sqrt{\sigma^2}$$
.

The advantage of the standard deviation over the variance is that it is measured in the same units as X.

2.2.2 Continuous random variable

Contrary to a discrete random variable, a continuous random variable can take values that are not countable. A continuous random variable can take infinetly many possible values within a certain range or interval. For a continuous random variable X the distribution is given by the probability density function f, which satisfies:

- f(x) is defined for all x in \mathbf{R} ,
- $f(x) \ge 0$ for all x in \mathbf{R} ,
- $\int_{-\infty}^{\infty} f(x) dx = 1$.

Condition 3. ensures that $P(-\infty < X < \infty) = 1$, which means that the probability of the random variable X being between $-\infty$ and ∞ is 100%. Furthermore the probability of X assuming a specific value a is zero, in other words: P(X=a)=0. That means that the values of the density function should not be interpreted as a probability of a given outcome. Instead the probability of X is found by integrating over the probability density function. So, the probability that a continuous random variable X lies between the values a and b is:

$$P(a < X < b) = \int_a^b f(x) dx.$$

A continuous random variable X also has a distribution function F(x), that also predicts whether X assumes a value equal to or smaller than x. For a continuous random variable it is again given by integrating over the probability density function in the interval from $-\infty$ to x:

$$F(x) = P(X \le x) = \int_{-\infty}^{x} f(y) \ dy$$

. That also means that P(a < X < b) can be calculated by F(b) - F(a).

For a continuous random variabel X the mean, variance and standard deviation the same interpretation applies. Just given by different formulars, which are:

$$\mu = E(X) = \int_{-\infty}^{\infty} x f(x) dx$$

and

$$\sigma^2 = E[(X - \mu)^2] = \int_{-\infty}^{\infty} (x - \mu)^2 f(x) dx,$$

(The standard deviation is still given by til square root of the variance).

If we are interested in certain parameters of a population distribution, we can look at a sample. From this, we can make a **point estimate**.

Examples of this are,

 \bar{x} is a point estimate of μ s is a point estimate of σ

This is often supplemented with a confidence interval

This is an interval around the point estimate, where we are confident that the population parameter is located.

For μ , we have different ways of estimating it. We can use the sample mean \bar{X} , or the average X_T of the sample upper and lower quartiles. But in this case, we have to look out for **bias**. If the distribution of a population is skewed, then X_T is biased. The result of this is, that in the long run, this estimator will systematically over or under estimate the value of μ . This is written as, $E(X_T) \neq \mu$.

It is generally preferred that the estimator is **unbiased**. In this case, \bar{X} is an unbiased estimate of the population mean μ .

The standard error of \bar{X} is $\frac{\sigma}{\sqrt{n}}$. Here, the standard error decreases, when the sample size increases. If an estimator has this property, it is called **consistent**. If we compare, the estimator X_T is also consistent, but has a greater variance than \bar{X} .

It is generally preferred that the estimator has the smallest possible variance, and in that case it is called efficient. So \bar{X} is an efficient estimator.

When estimating a parameter, the symbol $\hat{}$ is used above it. For μ , $\hat{\mu} = \bar{X}$.

We can calculate \bar{X} using the following formula,

$$\bar{X} = \frac{1}{n} \sum_{i=1}^{n} X_i$$

For the variance σ , we can estimate it by using the formula for S^2 ,

$$S^{2} = \frac{1}{n-1} \sum_{i=1}^{n} (X_{i} - \bar{X})$$

i

2.3 Probability distribution

Data can come in various distributions depending on different parameters such as degrees of freedom. The distribution is the shape of the data and it will have an effect on statistical models. Therefore it is important to have an understanding of distributions.

2.3.1 Normal distribution

In the world of statistics, the most common distribution is the normal distribution. It is constructed as a bell shape. The normal distribution is a continuous distribution, with this density function:

$$n(x; \mu, \sigma) = \frac{1}{\sqrt{2\pi\sigma}} e^{-\frac{(x-\mu)}{2\sigma^2}}$$

The distribution is dependent on the $\text{mean}(\mu)$ and the standard deviation (σ) , where changes to the mean will result in a change in the positioning of the normal distribution. Whereas a change in the standard deviation will change the spread of the curve. The normal distribution also always contains an area under the curve that is equal to one. This is to ensure that the normal distribution correctly models probability.

There is a special case of the normal distribution, called the standard normal distribution, where the mean is zero and the standard deviation is one. All variations of a normal distribution can be standardized by a transformation of the distribution, using the Z-score formula.

$$Z = \frac{X - \mu}{\sigma}$$

Z in the Z-score represents the amount of standard deviations a given X value, deviates from the mean.

2.3.2 The central limit theorem

A very effective theorem in statistics is the central limit theorem. This theorem states that if a random sample \overline{X} , with the size n, is taken from a population with a mean and a finite variance, then as n goes towards infinity, the distribution will resemble a normal distribution. If used with the Z-score formula, the distribution will resemble a standard normal distribution. The formula for the Z-score, when in conjunction with the central limit theorem, looks like this:

$$Z = \frac{\overline{X} - \mu}{\sigma / \sqrt{n}}$$

Where \overline{X} is a random sample of size n and μ is the mean of the true population. The standard error i represented by σ/\sqrt{n} , where σ is the standard deviation and n is the sample size. Usually the standard deviation is unknown, for these situations it's possible to use the estimator S^2 . This estimates the variance of the population from the variance of the sample, by this formula:

$$s^2 = \sum_{i=1}^n \frac{(x_i - \overline{x})}{n-1}$$

The square root of the variance is the standard deviation, therefore the square root of the estimator S^2 would be the estimated standard deviation. The problem with using the estimator S^2 , is that with small samples the variance is small and therefore it contains a lot of bias. In this situation the t-distribution would be used instead of the normal distribution, because the t-distribution takes the bias into account the bias of the standard deviation. It does this by having thicker tails, meaning that the probability of more extreme values are higher.

2.3.3 The t-distribution

The t-distribution is shaped as the standard normal distribution, in a bell shape and symmetrical around the mean of zero, the difference is that the t-distribution is more variable. This comes from the fact that the t-distribution is dependent on the degrees of freedom. When the degrees of freedom surpasses 30, the rule of thumb is that the distribution will resemble a normal distribution. So before 30 degrees of freedom, the distribution contains more variance. The t-distribution will come to resemble the standard normal distribution, when in surpasses 30 degrees of freedom, this makes sense, since the two distributions have the same formula:

$$T = \frac{\overline{X} - \mu}{S/\sqrt{n}}$$

The only difference is the estimated standard deviation S.

2.4 Statistical methods

2.4.1 Confidence intervals

The confidence interval is a good tool to use, when trying to estimate a parameter of a population. Its used to create an interval, where the parameter has a probability to lie inside of. This probability is called the confidence level and it's a chosen value, usually the chosen confidence level is either 95% or 99%. The confidence interval will become bigger with a larger confidence level. A good confidence interval is small with a large confidence level, this will usually occur when the sample size is large. The chosen confidence level relates to an α -value, where as an example the chosen confidence level is 95%, then the α -value would be 5% or normally written as 0.05. The α -value will sometimes be needed to find the critical value, that is used to calculate the margin of error, as an example it's used when trying to find the critical value of the confidence interval, when working with a t-distribution.

To set up a confidence interval, the margin of error needs to be computed and then that will be both added and subtracted from the point estimate. This will give the values of the outer bounds of the interval. The margin of error is calculated from this formula:

 $Margin_of_error = critical_value \pm standard_error$

The standard error will change depending on which parameter that the confidence interval is estimating, but the general formula for the standard error is:

$$\frac{\sigma}{\sqrt{n}}$$

An example of computing a confidence interval of the mean while working with a standard normal distribution, then the formula for the confidence interval would be this:

$$P(-z_{\alpha/2} < Z < z_{\alpha/2}) = 1 - \alpha$$

Where $1-\alpha$ is the confidence level. As it's the mean that is being estimated, then instead of Z-score, then μ must be isolated and that is done by multiplying $\frac{\sigma}{\sqrt{n}}$ and subtracting \bar{X} on all sides, then multiplying all side by -1 to remove the minus sign. So the formula for a confidence interval of the mean will look like this:

$$P(\bar{X} - z_{\alpha/2} \frac{\sigma}{\sqrt{n}} < \mu < \bar{X} + z_{\alpha/2} \frac{\sigma}{\sqrt{n}}) = 1 - \alpha$$

This formula will give the upper and lower bounds of the confidence interval.

The interpretation of a confidence interval

To interpret a confidence interval, it would be incorrect to interpret the confidence level of some value x, as the probability of the true parameter being inside of the interval. The reason behind this is that the computed interval is static, so either the value x is inside the interval or it's not. So the correct way of interpreting the confidence interval is by taking multiple samples and computing the confidence interval for all samples, then the value x would reside inside 95% of the confidence intervals. Kilde for fortolkningen af kofidense intervaller:

 $http://www.drhuang.com/science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_statistics_for_engineering_and_the_science/mathematics/book/probability_and_science/mathem$

2.4.2 Hypothesis testing

A hypothesis test is used to test an assumption about a population. This is done from a sample of the population, as the information about the population is usually hard to come by. A hypothesis test is set up, by having a null hypothesis and an alternate hypothesis.

$$H_0 = Null\ hypothesis$$

$$H_a = Alternate\ hypothesis$$

When working with hypothesis testing, the hypothesis H_0 is usually represented as the status quo, where as the hypothesis H_a is represented as the opposition. It is also important to note that there is only two outcomes of a hypothesis test, either H_0 is rejected in favor of H_a or H_0 is failed to be rejected. Therefore in no situation can H_0 be stated to be an absolute truth, as there might be other samples where H_0 will be rejected. Therefore in a hypothesis test H_0 needs to be the thing that can be rejected and if H_0 gets rejected, then H_a will become the new status quo until proven otherwise.

In a hypothesis test H_0 will be the assumption that a parameter for two populations is the same, where as H_a can be either one of three assumptions, depending on the intention of the hypothesis test.

$$H_0: \theta = \theta_0$$

1.
$$H_a: \theta \neq \theta_0$$

$$2. H_a: \theta < \theta_0$$

3.
$$H_a: \theta > \theta_0$$

When the direction of the rejection is not important and also is unknown, then (1) will be the case. This scenario sets up a two-tailed-test, where the hypothesis test is used to reject H_0 if H_a is either significantly larger or smaller than H_0 , this means that the critical area is on both sides of the difference of θ and θ_0 . Either (2) or (3) will set up a one-tailed-test, where depending on what is important, either the hypothesis test is used to determine if H_a is significantly bigger or smaller than H_0 . This means that the critical area only spans one side

of the difference between θ and θ_0 .

Error in hypothesis testing

When making a hypothesis test there is four different possible outcomes. The results are separated by correct decisions and errors. There exist two types of hypothesis errors, called type 1 error and type 2 error. The type 1 error occurs when H_0 is mistakenly rejected and H_0 is true. Type 2 error is the opposite, where H_a is rejected and H_a is true. The types of outcomes occurring from a hypothesis test can be seen in Table 1

	H_0 is true	H_0 is false
Does not reject H_0	Correct decision	Type 2 error
Reject H_0	Type 1 error	Correct decision

Table 1: Outcomes of a hypothesis test

It is possible to compute the type 1 error

3 Polynomial regression

Linear regression is a model that estimates the relationship between a dependent variable, y, and one or more independent variables, x.

A reasonable relationship between the two in simple regression is the linear relationship:

$$Y = \beta_0 + \beta_1 x$$

Where β_0 is the intercept, and β_1 is the slope.

In a lot of cases, there will be more independent variables, so the relationship for multiple regression will look like this:

$$Y = \beta_0 + \beta_1 x_1 + \dots + \beta_n x_n$$

Where n is the number of independent variables. Linear models use the method of least squares of the residuals to estimate parameters, in order to find the best fitting line for the data.

In simple linear regression, the random error ϵ is included:

$$Y = \beta_0 + \beta_1 x + \epsilon$$

It is assumed that ϵ is distributed with $\epsilon = 0$ and ϵ) = σ^2 , and it has consistent variance, which is usually called the *homogeneous variance assumption*. The random error ϵ adds randomness to account for the natural variability in real data, making the model more realistic.

Polynomial Regression

Polynomial regression is a form of linear regression, but the relationship between x and y is an nth-degree polynomial. It fits a nonlinear relationship between the value of x and the corresponding conditional mean of y, meaning the model predicts the expected value of y given x.

That is why it is used when the relationship between the independent variable and the dependent variable is better represented by a curve rather than a straight line, since it can show the nonlinear patterns in the data.

3.1 Assumptions

3.1.1 Homoscedasticity

One of the assumptions of a polynomial regression is that homoscedasticity is fulfilled. Homoscedasticity is the assumption of constant error variance, where observations in a dataset would exhibit errors that have roughly the same spread across all levels of the independent variable.

If this assumption is not upheld, then this will cause the standard error to be

biased and therefore not trustworthy. This problem causes further testing involving this standard error to become wrong, an example is the hypothesis test. The reason for the assumption needs to be upheld, comes from how the regression is created. The regression is created via the ordinary least square method, that requires the assumption of homoscedasticity to be upheld.

A way to display homoscedasticity is through the variance-covariance matrix. The matrix shows whether the data contains homoscedasticity or heteroscedasticity through the diagonal values. If the matrix contains all the same values through the diagonal, then the assumption of homoscedasticity is upheld, else the data contains heteroscedasticity. This is a showcase of the variance-covariance matrix with homoscedasticity:

Every position in the matrix is calculated, then the diagonal will tell if the data contains homoscedasticity or heteroscedasticity. The positions that are not on the diagonal should be zero else the data contains another problem, that is autocorrelation, meaning that the observations in the data set are correlated.

Source: https://openpublishing.library.umass.edu/pare/article/id/1590/

3.1.2 No multicollinearity

Perfect multicollinearity is a term used for describing a perfect linear relationship between two or more independent variables. This relationship occurs when an independent variable can be perfectly predicted from other independent variables. In mathematical terms, this could be written as a linear regression:

$$X_1 = c + \beta_1 \cdot X_2 + \dots + \beta_n \cdot X_n$$

Where $X_1...X_n$ is all the independent variables that have a perfect linear relationship. The coefficients are represented by $\beta_1...\beta_n$ and they are the amount that X_1 changes when their relative independent variable changes. Lastly c is the intercept and represents the value of X_1 , when all other independent variables are zero.

The regression model can feel the effects of multicollinearity even without there being perfect multicollinearity. A strong linear relationship is enough to have an effect on the model. The problem caused by multicollinearity, is that as it increases the variance of the value that the coefficients can receive also increases. Where as perfect multicollinearity will make the model unable to estimate a value of one coefficient, due to the perfect linearity between the independent variables.

The VIF (variance inflation factor) is used to measure how much the variance of the coefficients are inflated due to multicollinearity. The formula for the VIF, is as followed:

$$\frac{1}{1-R^2}$$

Where R^2 is the coefficient of dertermination. This coefficient of dermination is calculated through a linear regression that is set up with an independent variable as the dependent variable and all the other independent variables staying

as independent variables. This is done for all independent variables, so each independent variable has a VIF-value. A rule of thumb is that a VIF-value greater that ten is significant and therefore needs to be dealt with.

 $Source: \ https://ekja.org/upload/pdf/kja-19087.pdf$

3.2 Assumptions

4 Pseudo random number generator

5 Pseudo Random Number Generator

To generate the dataset, we use a Pseudo-Random Number Generator (PRNG). PRNGs are algorithms that produce sequences of numbers that appear random but are actually deterministically generated from an initial seed value. While these numbers are not truly random, they are sufficiently unpredictable for many practical applications. Random numbers are widely used in fields such as statistics, game theory, cryp- tography, and simulations. These applications require numbers that behave as if they were random, yet can be reproduced when needed. This is where PRNGs come in—they allow for repeatable randomness, making them ideal for controlled experiments, testing, and security. This chapter will explore the key concepts behind PRNGs. Before going into the mechanics of these generators, it is important to first understand what 'random' means and the characteristics that define truly random numbers.

5.1 Properties of PRNGs

The quality of a PRNG is determined by several key factors that influence its use for different applications. Some of the properties of a good PRNG is properties: Independency, a large period and reproducibility The numbers produced by the PRNG should be statistically independent, en- suring that each generated value exhibits no correlation with previous numbers or other sequences. This implies that knowledge of previously generated numbers or sequences provides no advantage in predicting the next output. A PRNG operates within a specific interval before its sequence begins to repeat. A high-quality PRNG has a long interval, delaying repetition and enhancing its unpredictability. Conversely, a PRNG with a shorter period becomes more pre-dictable and less suitable for practical use. A key feature of a PRNG is its ability to reproduce the same sequence of numbers when given a specific seed. This property is particularly useful in testing and simulation scenarios, where it is essential to generate identical sequences multiple times for consistency and reproducibility In addition, a PRNG must be fast and efficient to prevent it from introducing performance bottlenecks within an application. The speed of number generation directly impacts computational efficiency, especially in applications requiring a large volume of random numbers. An inefficient PRNG can significantly slow down processes, undermining the overall performance of the system. Therefore, balancing randomness and efficiency is essential for practical applications

5.2 Linear Congruential Generator

Linear Congruential Sequence (LCS) is a commonly used approach to generate pseudo-random numbers. LCS generates a sequence of numbers using a linear recurrence relation LCS is expressed as:

$$X_{n+1} = (aX_n + c) \bmod m.$$

where X_0 is the seed, a is the multiplier, c is the increment, and m is the modulus.

Example: Given a = 5, c = 1, m = 16, and $X_0 = 7$:

$$X_1 = (5 \cdot 7 + 1) \mod 16 = 4$$

$$X_2 = (5 \cdot 4 + 1) \mod 16 = 5$$

$$X_3 = (5 \cdot 5 + 1) \mod 16 = 10$$

$$X_4 = (5 \cdot 10 + 1) \mod 16 = 3$$

This sequence has a period of 16. In an LCG, the period can be as large as m, but choosing parameters carefully is crucial to achieving long periods. Therefore

- 6 Monte Carlo
- 6.1 Assumptions

7 Problem statement

8 intro til data

9 intro til data

To determine whether the data set meets the assumption of homoscedasticity, a scatter plot between the dependent and independent variables is created. It becomes apparent that the variables displacement, weight, and acceleration are not homoscedastic. Furthermore, MPG, displacement, weight, and acceleration are continuous numeric variables, while cylinders, model year, and origin are discrete numeric variables. Additionally, a heat map of the correlation between variables is created to determine if multicollinearity is present among the independent variables. Here, it is apparent that weight, displacement, and cylinders are highly correlated, and that all three are highly correlated with MPG. It is also clear from observing the density functions of the independent variables that none of them, except acceleration, are normally distributed.

```
1 library(ggplot2)
2 library(dplyr)
3 library(gridExtra)
4 library (reshape2)
5 library(lmtest)
6 library (gridExtra)
7 #read data
8 data <- read.csv("auto-mpg.csv", na.strings = ".")</pre>
10 #calculet the NA-%
na_percentage <- function(column) {</pre>
    sum(is.na(column)) / length(column) * 100
12
13 }
15 #print the NA-%
  print(sapply(data, na_percentage))
_{18} # Convert horsepower to numeric, coercing non-numeric values to NA
19 data$horsepower <- as.numeric(data$horsepower)</pre>
20
21
22
23 # Remove rows with any NA values
24 data <- data %>% na.omit()
25
  "numeric_data <- data %>% select_if(is.numeric)
27
28 # Beregn korrelationer og smelt til format for ggplot2
29 data_korrelationer <- melt(cor(numeric_data, use = "complete.obs"))</pre>
30
  "Opret heatmap baseret p korrelationer
31
ggplot(data_korrelationer, aes(x = Var1, y = Var2, fill = abs(value
      ))) +
    geom_tile() +
33
    geom_text(aes(label = round(value, 2)), color = "black", size =
34
    scale_fill_gradient(low = "white", high = "red", limit = c(0, 1),
       name="|Correlation|") +
```

```
theme_minimal() +
36
    theme(axis.text.x = element_text(angle = 45, hjust = 1)) +
37
    labs(title = "Correlation Matrix Heatmap", x = "", y = "")
38
40 " Lav scatter plots for hver kolonne mod 'mpg'
41 scatter_plots <- lapply(names(numeric_data), function(col) {</pre>
    ggplot(numeric_data, aes_string(x = col, y = "mpg")) +
      geom_point(color = "blue") +
43
      ggtitle(paste("Scatter Plot of", col, "vs mpg")) +
45
      theme_minimal()
46 })
47
48 "bp_results <- sapply(names(numeric_data), function(col) {</pre>
    model <- lm(mpg ~ numeric_data[[col]], data = numeric_data)</pre>
    bp_test <- bptest(model)</pre>
    bp_test$p.value
51
52 })
53
54 #make df with P_Value
55 bp_df <- data.frame(Variable = names(bp_results), P_Value = bp_</pre>
      results)
57 print(bp_df)
59
60 arrange scatter plots and Breusch prage plot
61 do.call(grid.arrange, c(scatter_plots, ncol = 3))
_{\rm 63} "Lav histogrammer og density plots med mean og SD
64 plots <- lapply(names(numeric_data), function(col) {</pre>
    n_bins <- ceiling(log2(length(numeric_data[[col]])) + 1)</pre>
    mean_value <- mean(numeric_data[[col]])</pre>
66
    sd_value <- sd(numeric_data[[col]])</pre>
67
68
69
    ggplot(numeric_data, aes_string(x = col)) +
70
      geom_histogram(aes(y = ..density..), bins = n_bins, fill = "
      blue", color = "black") +
71
      geom_density(color = "red", size = 1) +
      geom_vline(aes(xintercept = mean_value), color = "green",
72
      linetype = "dashed", size = 1) +
      ggtitle(paste("Density, Mean and SD of", col)) +
74
      theme_minimal() +
      annotate("text", x = Inf, y = Inf, label = paste("Mean:", round
      76
77 })
79 do.call(grid.arrange, c(plots, ncol = 3))
```

Listing 1: intro til data

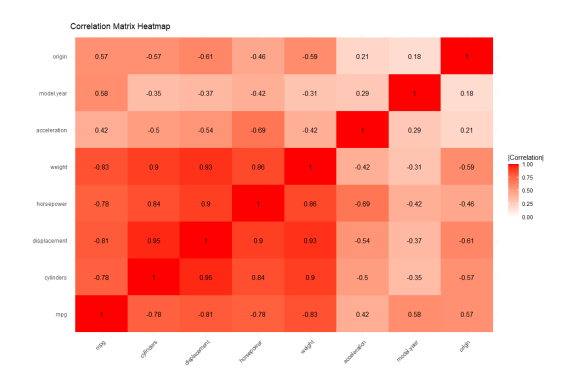


Figure 1: heatmap

10 Classical regression

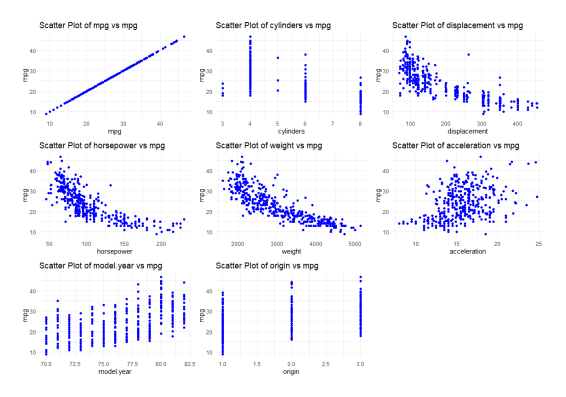


Figure 2: scatterplot

11 Monte Carlo regression

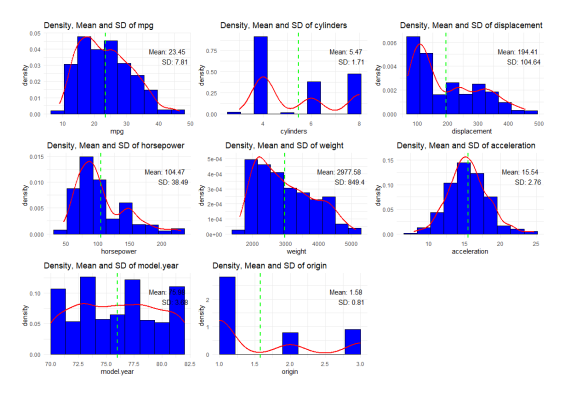


Figure 3: density, mean and sd

12 montecarlo bootstrapping

Monte Carlo bootstrapping is used to find the model for the mpg data. It starts by cleaning the data, then selecting the numeric columns, and defining the Monte Carlo bootstrapping function. After that, it defines the polynomial regression function using mpg as the dependent variable. Next, it defines the simulation function using clusters, thereby speeding up the computation time with the doParallel library. After defining the run simulation function, it runs the simulation with the set seed, simulation number, and sample size of the Monte Carlo bootstrapping method.

After that, it creates histograms of the simulation results. It calculates the mean result of the coefficients, and using these means, it calculates a new regression model and the R-squared of the new average model. Finally, it makes a scatter plot showing the final regression model results versus the results from the actual data set.

```
library(dplyr)
library(ggplot2)
library(foreach)
library(doParallel)
library(gridExtra)
```

```
6
7 # Set working directory and read data
8 setwd("C:/Users/Jonathan/Documents/GitHub/P2/R kode")
9 data <- read.csv("auto-mpg.csv", na.strings = ".")</pre>
11 # Convert horsepower to numeric, coercing non-numeric values to NA
data$horsepower <- as.numeric(data$horsepower)</pre>
# Remove rows with any NA values
data <- data %>% na.omit()
17 # Select numeric columns
numeric_data <- data[sapply(data, is.numeric)]</pre>
20 # Function to perform Monte Carlo bootstrapping
21 monte_carlo_bootstrap <- function(data, sample_size) {</pre>
   data %>%
      sample_n(sample_size, replace = TRUE)
23
24 }
25
26 # Fit polynomial regression function
27 fit_polynomial_regression <- function(data) {</pre>
    predictors <- setdiff(names(data), "mpg")</pre>
28
    formula <- reformulate(termlabels = paste0("poly(", predictors, "</pre>
       , 2)"), response = "mpg")
    lm(formula, data = data)
30
31 }
32
33 # Run simulations
34 run_simulations <- function(n_simulations, numeric_data, sample_
    num_cores <- detectCores() - 1</pre>
35
    cl <- makeCluster(num_cores)</pre>
36
    registerDoParallel(cl)
37
38
39
     clusterExport(cl, c("numeric_data", "fit_polynomial_regression",
      "monte_carlo_bootstrap", "sample_size"))
     clusterEvalQ(cl, library(dplyr))
40
41
    results <- foreach(i = 1:n_simulations, .combine = rbind, .
42
      packages = "dplyr") %dopar% {
      resampled_data <- monte_carlo_bootstrap(numeric_data, sample_</pre>
43
      size)
      model <- fit_polynomial_regression(resampled_data)</pre>
44
       c(coef(model), summary(model)$r.squared)
45
46
47
    stopCluster(cl)
48
49
    results_df <- as.data.frame(results)
50
    colnames(results_df) <- c(names(coef(fit_polynomial_regression(</pre>
51
      numeric_data))), "r_squared")
52
    return(results_df)
53
54 }
56 # Set seed and run simulations
```

```
57 set.seed(210)
58 n_simulations <- 10000</pre>
59 sample_size <- 300</pre>
60 results_df <- run_simulations(n_simulations, numeric_data, sample_
       size)
61
62 # Clean column names to remove special characters
63 clean_colnames <- function(df) {
     colnames(df) <- make.names(colnames(df), unique = TRUE)</pre>
65
    return(df)
66 }
67
68 results_df <- clean_colnames(results_df)
70 # Create histograms
71 create_histograms <- function(df) {</pre>
     plots <- lapply(names(df), function(col) {</pre>
       mean_value <- mean(df[[col]])</pre>
73
74
       sd_value <- sd(df[[col]])</pre>
75
       ggplot(df, aes_string(x = col)) +
76
         geom_histogram(aes(y = ..density..), bins = 30, fill = "blue"
77
        , color = "black", alpha = 0.7) +
78
         geom_density(color = "red", size = 1) +
         geom_vline(aes(xintercept = mean_value), color = "green",
79
       linetype = "dashed", size = 1) +
         ggtitle(paste("Density, Mean and SD of", col)) +
80
          theme_minimal() +
81
       annotate("text", x = Inf, y = Inf, label = paste("Mean:",
round(mean_value, 2), "\nSD:", round(sd_value, 2)),
82
                    hjust = 1.1, vjust = 2, color = "black", size = 4)
84
     })
85
     grid.arrange(grobs = plots, ncol = 2)
86
87 }
89 # Generate histograms for the simulation results
90 create_histograms(results_df)
91
_{92} # Calculate the mean of the coefficients from the Monte Carlo
       simulation
93 best_coefficients <- colMeans(results_df)</pre>
95 print(best_coefficients)
97 # Fit the final model using the original data
98 final_model <- fit_polynomial_regression(numeric_data)
_{\rm 100} # Create a function to apply averaged coefficients
apply_coefficients <- function(model, coefficients) {</pre>
    model$coefficients <- coefficients
102
103
     return(model)
104 }
105
# Apply the averaged coefficients to the final model
final_model <- apply_coefficients(final_model, best_coefficients)</pre>
```

```
# Predict using the final model
y_final_pred <- predict(final_model, newdata = numeric_data)</pre>
111
# Calculate R-squared
113 actual_values <- numeric_data$mpg</pre>
ss_total <- sum((actual_values - mean(actual_values))^2)
ss_residual <- sum((actual_values - y_final_pred)^2)
r_squared <- 1 - (ss_residual / ss_total)</pre>
118 # Print R-squared value
   cat("R-squared:", r_squared, "\n")
119
120
121 # Plot the final regression model
   ggplot(data.frame(Actual = actual_values, Predicted = y_final_pred)
       , aes(x = Actual, y = Predicted)) +
     geom_point(alpha = 0.7) +
123
     labs(title = paste("Final Regression Model (R-squared:", round(r_
       squared, 2), ")"),
         x = "Actual Values", y = "Predicted Values") +
125
     theme_minimal()
126
127
128 # Summarize models
129 klassisk_model <- fit_polynomial_regression(numeric_data)</pre>
130 summary(final_model)
131 summary(klassisk_model)
```

Listing 2: montecarlo bootstrapping

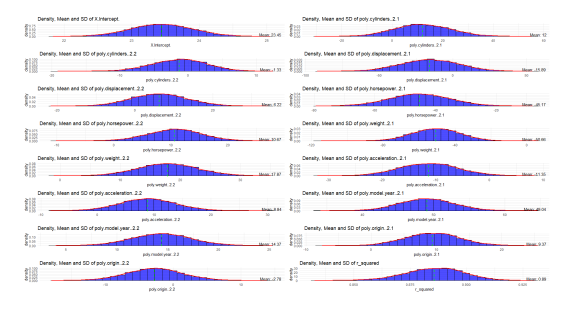


Figure 4: AQI gennem årene.

13 super syntetisk

14 super syntetisk

To show the consequences of violating the assumption of homoscedasticity, a dataset is generated with and without homoscedasticity. It starts with generating four normally distributed independent variables using a random number generator. Then, the dependent variable is generated as a function of the four independent variables. A regression model is fitted, and the model is tested. After this, homoscedasticity is violated by adding an error term that scales with the dependent variable. The model is tested again, and it becomes apparent what the consequences of violating the assumption of homoscedasticity are.

```
1 library(ggplot2)
2 library(gridExtra)
3 library(lmtest)
5 # Set seed for reproducibility
6 set.seed(223)
8 # Generate independent variables
9 n <- 250
x1 < rnorm(n, mean = 5, sd = 1)
x2 < - rnorm(n, mean = 10, sd = 4)
x3 \leftarrow rnorm(n, mean = 15, sd = 3)
13 \times 4 < - rnorm(n, mean = 20, sd = 5)
14
_{15} # Generate dependent variable with a polynomial relationship
_{16} y <- 3 + 2*x1 + 5*x1^2 + 1.5*x2 + 3*x3^2 + 2*x4 + 0.001*rnorm(n,
      mean = 0, sd = 1)
18 # Create a data frame
19 data <- data.frame(y, x1, x2, x3, x4)</pre>
21 # Fit a polynomial regression model
22 model <- lm(y \sim poly(x1, 2) + x2 + poly(x3, 2) + x4, data = data)
23
24 # Summary of the model
25 summary (model)
27 # Add an error term to x1 that scales with the corresponding y
x3_{new} < -x3 + rnorm(n, mean = 0, sd = 0.00254 * abs(y))
29
30 # Generate new dependent variable with the same polynomial
      relationship
y_{new} \leftarrow 3 + 2*x1 + 5*x1^2 + 1.5*x2 + 3*x3_{new}^2 + 2*x4 + 0.001*
      rnorm(n, mean = 0, sd = 1)
32
33 # Create a new data frame
34 data_new <- data.frame(y_new, x1 = x1, x2, x3_new, x4)</pre>
36 # Fit a new polynomial regression model
37 model_new <- lm(y_new \sim poly(x1, 2) + x2 + poly(x3, 2) + x4, data =
   data_new)
```

```
38
39 # Summary of the new model
40 summary (model_new)
^{42} # Diagnostic plots for the original model
_{43} par(mfrow = _{c}(2, 2))
44 plot(model)
45
46 # Diagnostic plots for the new model
_{47} par(mfrow = _{c}(2, 2))
48 plot(model_new)
49
_{\rm 50} # Function to create scatter plots for each numeric column against
      у,
51 scatter_plots <- function(data, color, title_prefix) {</pre>
    lapply(names(data)[-1], function(col) {
52
      ggplot(data, aes_string(x = "y", y = col)) +
53
        geom_point(color = color) +
54
55
         ggtitle(paste(title_prefix, col, "vs y")) +
         theme_minimal()
56
57
    })
58 }
59
60 # Scatter plots for normal data
61 scatter_plots_normal <- scatter_plots(data, "blue", "Normal Data:
      Scatter Plot of")
62 # Scatter plots for non-homoscedastic data
63 scatter_plots_non_homoscedastic <- scatter_plots(data_new, "red", "
      Non-Homoscedastic Data: Scatter Plot of")
64
# Arrange scatter plots in grids
66 grid.arrange(grobs = scatter_plots_normal, ncol = 2, top = "Scatter
       Plots for Normal Data")
67 grid.arrange(grobs = scatter_plots_non_homoscedastic, ncol = 2, top
       = "Scatter Plots for Non-Homoscedastic Data")
69 # Function to create histograms for each numeric column
70 create_histograms <- function(data, color, title_prefix) {</pre>
    lapply(names(data), function(col) {
71
72
      n_bins <- ceiling(log2(length(data[[col]])) + 1)</pre>
       ggplot(data, aes_string(x = col)) +
73
         geom_histogram(bins = n_bins, fill = color, color = "black")
74
         ggtitle(paste(title_prefix, col)) +
75
         theme_minimal()
76
    })
77
78 }
80 # Histograms for normal data
81 histograms_normal <- create_histograms(data, "blue", "Histogram of"
82 # Histograms for non-homoscedastic data
83 histograms_non_homoscedastic <- create_histograms(data_new, "red",</pre>
       "Histogram of")
85 # Arrange histograms in grids
```

Listing 3: super syntetisk data

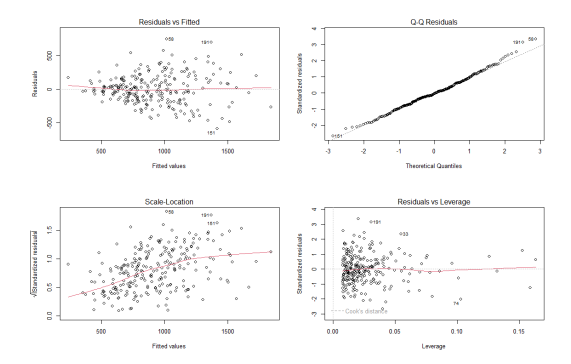


Figure 5: residualplot

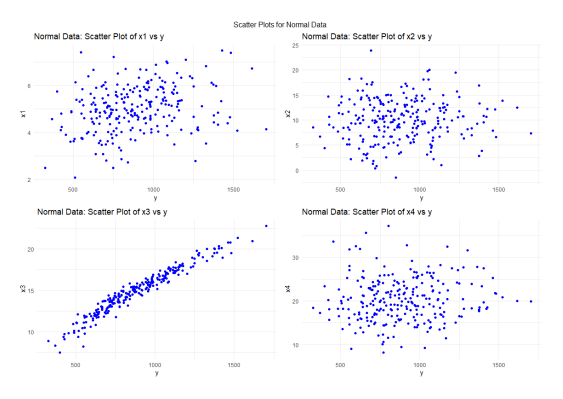


Figure 6: scatterplot.homo

15 Comparison between regressions

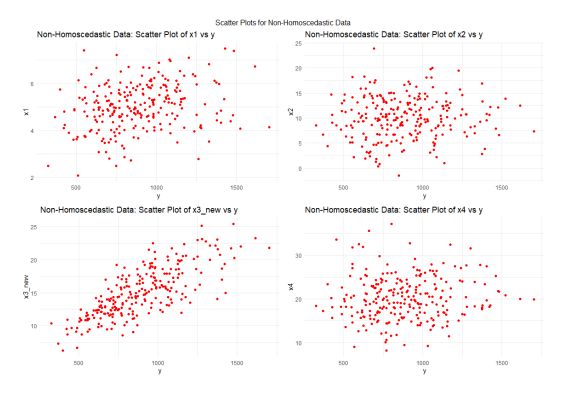


Figure 7: scatterplot.hetro

16 Discussion

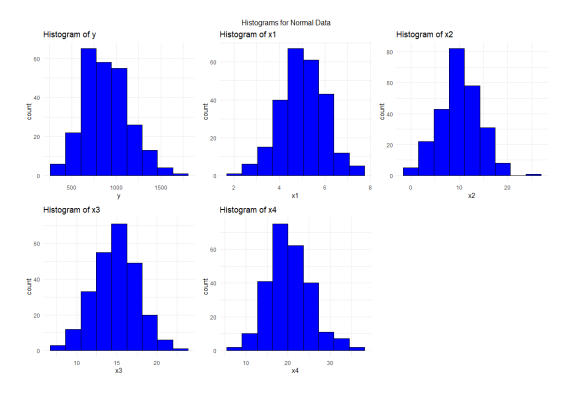


Figure 8: his.homo

17 Conclusion

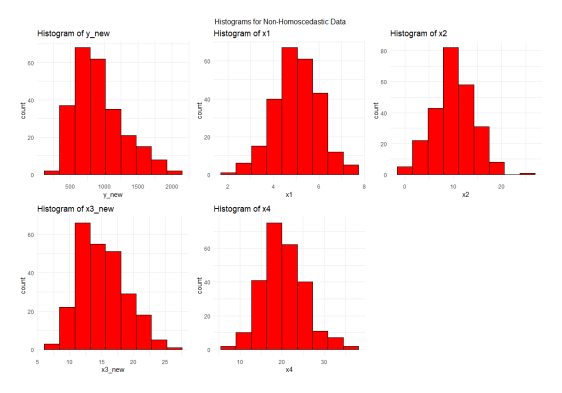


Figure 9: his.hetro.

18 Litteratur

19 latextables