MULTILINEAR REGRESSION RESEARCH ON **FACTORS** INFLUENCING LIFE EXPECTANCY ON DIFFERENT COUNTRIES



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ABSTRACT

Delving into the labyrinth of life expectancy disparities, we aim to unravel the intricate web of factors shaping longevity across diverse global landscapes. By conducting a comprehensive analysis across diverse countries, our research seeks to offer a nuanced perspective on global life expectancy dynamics, shedding light on the multifaceted determinants shaping both individual and population health outcomes.

To achieve this objective, we will undertake a rigorous examination of life expectancy data sourced from a multitude of authoritative outlets, including governmental records, international repositories, and scholarly literature. Employing robust statistical methodologies such as regression analysis and advanced data visualization techniques, we aim to pinpoint salient variables, unveil intricate relationships, and discern notable trends or anomalies.

Furthermore, our inquiry will extend beyond quantitative analysis to encompass a qualitative exploration of factors influencing life expectancy. This holistic approach will encompass an examination of healthcare infrastructure, socio-economic variables, educational attainment, and lifestyle choices. Through this comprehensive lens, our study aspires to furnish a profound understanding of the multifaceted determinants contributing to disparities in life expectancy across diverse global contexts.

INTRODUCTION

Life expectancy is a very important measure for assessing a country's overall. population health. Predicting the life expectancy helps the policy makers of a country. when making policies for a country hence life expectancy measure is essential. This measure then assists the policy makers of the country to decides the necessary social and health welfare changes in the country.

According to the United Nations, life expectancy at birth numbers corresponds to midyear estimations. They align with the relevant United Nations quinquennial population forecasts and fertility averages. Life tables are created using available mortality and data from civil registration. The World Health Organization supplies all the life tables, using data derived from the year 1,800 life tables based on the logit system.

A multitude of factors shape life expectancy, spanning various demographic, socioeconomic, and health-related aspects. These determinants exhibit notable diversity across nations owing to disparities in economic advancement, healthcare systems, social regulations, cultural practices, and epidemiological characteristics.

Hence, undertaking thorough analyses to pinpoint the primary influencers of life expectancy in diverse contexts is imperative for customizing interventions that efficiently cater to specific population health requirements.

The results of this study carry notable ramifications for public health policy and implementation.

Through understanding the factors influencing life expectancy, policymakers can craft tailored interventions to tackle primary causes of mortality and advocate for healthier behaviours, consequently, fostering enhancements in overall population health and welfare. Moreover, comprehending the determinants of life expectancy can facilitate comparisons between countries, pinpoint intervention areas, and aid in allocating resources effectively to optimize the outcomes of public health endeavours.

By doing this research, we hope to increase our knowledge of the factors that affect. life expectancy and make a valuable contribution to evidence-based strategies for improving population health and well-being in a range of sociodemographic settings.

We seek to promote more resilience against health difficulties in an increasingly interconnected society by facilitating informed decision-making and illuminating the intricate interactions between diseases and broader socioeconomic and environmental factors.

DATA CLEANING

Data Dictionary

Variable name	Variable description	Variable type
Country	Country, we observed	category
Year	Years we observed category	
Status	Developed or developing category	
Life expectancy	Life expectancy in age	numeric

Adult mortality	Adult mortality rate	numeric
Infant mortality	Infant mortality rate	numeric
Alcohol	Per capita alcohol consumption	numeric
Percentage expenditure	As percentage	numeric
Hepatitis B	Immunization coverage	categoric
Measles	Number of cases reported	numeric
BMI	Average body mass index	numeric
Under five deaths	Number of Under five deaths	numeric
Polio	Polio immunization coverage	categoric
Total expenditure	Government health expenditure	numeric
Diphtheria	Diphtheria coverage	categoric
HIV/AIDS	HIV/AIDS deaths	numeric
GDP	GDP/capita	numeric
Population	Population of the country	numeric

TABLE 1

Dataset

The Dataset that we used had 21 variables ,18 which are of numeric and 3 of which are a discrete data (category).

Variable selection criteria

The selection criteria for variables in the research encompass several key considerations to ensure a comprehensive understanding of the factors influencing life expectancy across different countries. Among the variables included are BMI (Body Mass Index), which reflects the population's overall health and nutrition status. Adult mortality rates provide insights into the prevalence of diseases and the effectiveness of healthcare systems in addressing them.

Similarly, infant death rates offer crucial information on maternal and child health, including access to prenatal and neonatal care.

Alcohol consumption rates serve as an indicator of lifestyle and behavioural factors that impact health outcomes. Percentage expenditure on healthcare highlights the level of investment in healthcare infrastructure and services, which directly affects access and quality of care. Disease specific factors such as hepatitis, measles, polio, and diphtheria vaccination rates shed light on the effectiveness of immunization programs and public health initiatives in preventing communicable diseases.

Moreover, the prevalence of HIV/AIDS serves as a critical indicator of the burden of infectious diseases and the effectiveness of prevention and treatment efforts. Economic indicators such as GDP (Gross Domestic Product) provide context on the overall wealth and development status of countries, which influences access to resources and healthcare services. Population size is included to account for demographic variations and scale effects in the analysis.

By incorporating these variables, researchers can explore the multifaceted determinants of life expectancy, ranging from individual health behaviours to broader socio-economic and healthcare system factors. This comprehensive approach enables the identification

of significant predictors and informs evidence-based public health policies aimed at improving population health outcomes worldwide. Then we proceeded to remove all variables that do not provide any additional information on predictors about life expectancy

The Country has a high cardinality problem, there are more countries repeated, every country has economic, social, and health-related data that has been included separately therefore, country variable does not provide any viable information that we can use to the analysis hence we removed it. The variable Year is a time series, and our analysis focuses on time independent economic, social and health related predictors of life expectancy, removing Country and Year helps on making dataset more streamlined for analysis. The Status variable is of no numeric data which would not affect our calculations as well as analysis in life expectancy. We then proceeded to remove Income composition of resource, Thinness of 5-9 years, Thinness 1-18 years, Total expenditure and under five deaths. These Factors were the least contributing factors towards our model.

The Polio, Diphtheria and Hepatitis B are to be regarded as discrete(categorically) since it's the coverage of immunization which is not continuous.

Exploratory Data Analysis

Model fitting.

```
> mydata=read.table("clipboard", header = TRUE)
> attach(mydata)
The following objects are masked from mydata (pos = 3):
    x1, x10, x11, x12, x2, x3, x4, x5, x6, x7, x8, x9, y
> f1 = lm(y ~ x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 + x10 + x11 + x12)
> f1

Call:
lm(formula = y ~ x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 + x10 + x11 + x12)

Coefficients:
(Intercept)    x1     x2     x3     x4     x5     x6     x7     x8     x9     x10     x11     x12
        6.379e+01    -2.484e-02    -6.620e-03     2.975e-01    -1.215e-04    -7.287e-03     1.714e-05     9.73
2e-02     2.293e-02     4.225e-02    -4.531e-01     1.463e-04     x12     5.963e-09
>
```

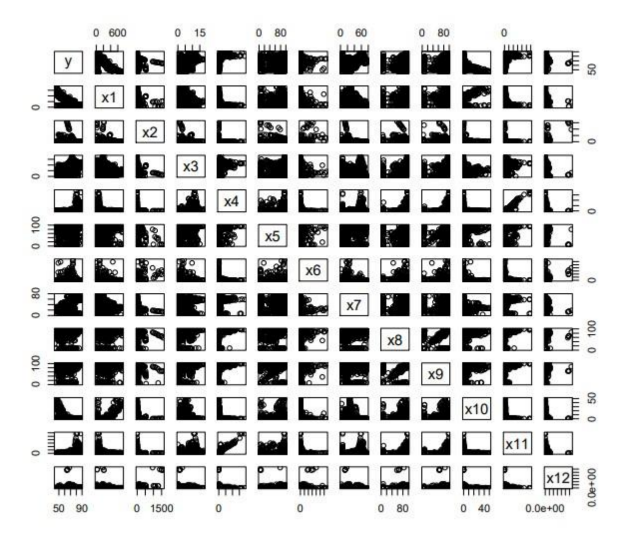
Summary of the initial model

```
> summary(f1)
Call:
lm(formula = y \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 +
    x10 + x11 + x12)
Residuals:
             1Q Median 3Q Max
.5064 0.2694 2.7378 17.6119
     Min
-21.2824 -2.5064
Coefficients:
               Estimate Std. Error t value Pr(>|t|)
2.975e-01 3.244e-02 9.171 < 2e-16 ***
x3
            -1.215e-04 2.230e-04 -0.545 0.585938
            -7.287e-03 5.515e-03 -1.321 0.186555
1.714e-05 1.297e-05 1.321 0.186553
9.732e-02 6.488e-03 14.999 < 2e-16 ***
x5
x6
x7
x8
             2.293e-02 6.369e-03 3.601 0.000326 ***
             4.225e-02 7.268e-03 5.813 7.37e-09 ***
x9
            -4.531e-01 2.203e-02 -20.565 < 2e-16 ***
1.463e-04 3.478e-05 4.206 2.74e-05 ***
5.963e-09 2.135e-09 2.792 0.005296 **
x10
x11
x12
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Residual standard error: 4.472 on 1636 degrees of freedom
Multiple R-squared: 0.7434,
                                  Adjusted R-squared: 0.7415
F-statistic: 395 on 12 and 1636 DF, p-value: < 2.2e-16
```

FIGURE 1

The Summary of the initial fitted model using 12 variables against Life expectancy $\dot{Y} = 6.379e + 01 - 2.484e - 02Adult Mortality - 6.620e - 01Infant Deaths + 2.975e - 01Alcohol - 1.215e 04Percentage Expenditure - 7.287e - 03Hepatitis B + 1.714e - 05Measles + 9.732e - 02BMI + 2.293e - 02Polio + 4.225e - 02Diphtheria - 4.531e - 01HIV/AIDS + 1.463e - 04GDP + 5.963e 09Population$

```
> mydata=read.table("clipboard", header = TRUE)
> attach (mydata)
The following objects are masked from mydata (pos = 3):
    x1, x10, x11, x12, x2, x3, x4, x5, x6, x7, x8, x9, y
The following objects are masked from mydata (pos = 6):
    x1, x10, x11, x12, x2, x3, x4, x5, x6, x7, x8, x9, y
> head (mydata)
     y x1 x2
              x3
                         x4 x5
                                x6
                                     x7 x8 x9 x10
                                                         x11
                                                                  x12
1 65.0 263 62 0.01 71.279624 65 1154 19.1 6 65 0.1 584.25921 33736494
2 59.9 271 64 0.01 73.523582 62 492 18.6 58 62 0.1 612.69651
3 59.9 268 66 0.01 73.219243 64 430 18.1 62 64 0.1 631.74498 31731688
4 59.5 272 69 0.01 78.184215 67 2787 17.6 67 67 0.1 669.95900 3696958
5 59.2 275 71 0.01 7.097109 68 3013 17.2 68 68 0.1 63.53723 2978599
6 58.8 279 74 0.01 79.679367 66 1989 16.7 66 66 0.1 553.32894 2883167
> plot (mydata)
>
```



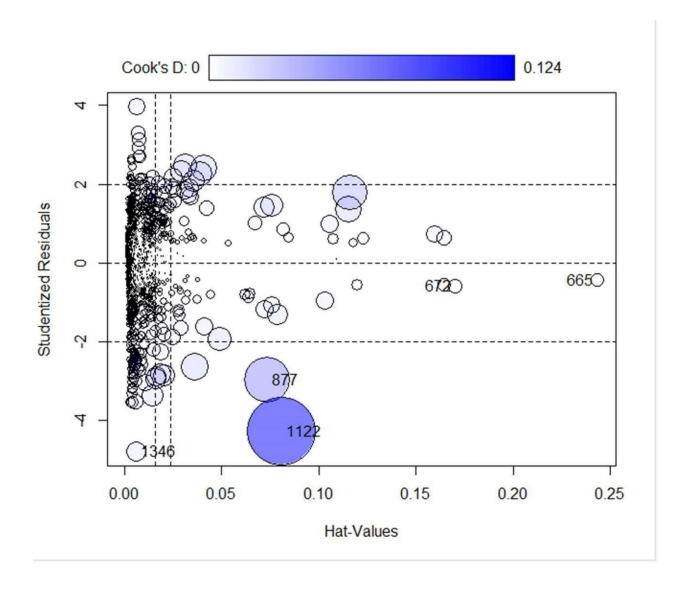
Life Expectancy is negatively related to the variables Adult Mortality, Alcohol, Infant deaths, Percentage Expenditure, Hepatitis B, and Measles, HIV/AIDS and Population and others are positively related to the life expectancy. Infant mortality is linked to life expectancy. The Newborn Deaths variable is expected to have a negative relationship with Life Expectancy since a higher number of baby deaths suggests that many infants die before reaching the age of one, reducing the country's life expectancy.

The Multiple R-squared is at 0.7434 close to 1 suggesting that the model is significant.

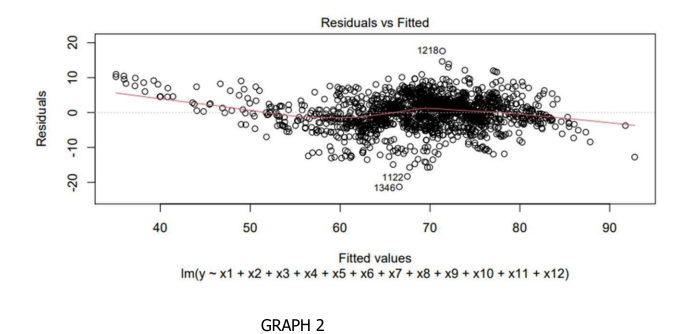
ASSUMPTION CHECKING

We started by checking for outliers using DFFITS and DFBETAS

Influence Plot

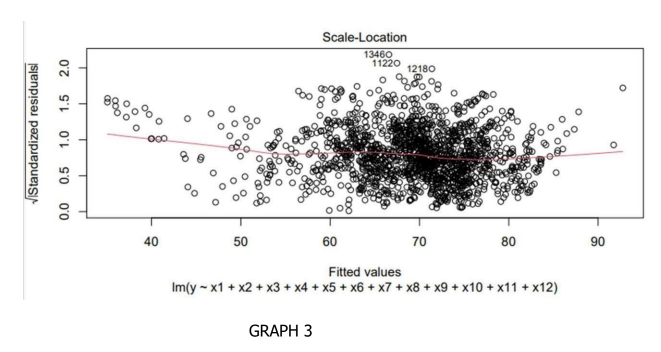


GRAPH 1



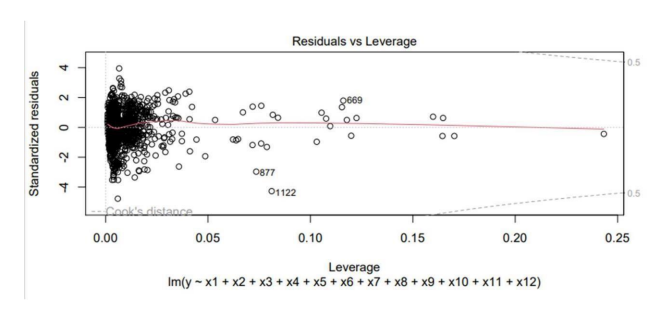
The scatterplot of the residuals indicates that most data points are closely clustered around the horizontal line at y = 0, suggesting a good fit between the model and the data without any systematic bias. However, a few outliers are observed, indicating instances where the model did not fit the data accurately. Despite these outliers, there's no discernible pattern to their distribution, making it challenging to pinpoint their cause. Overall, the model generally fits the data well.

Scale-Location



This plot shows a "fanning out" pattern of the residuals, which is typical of a model where the residual (error) variances are not all equal (heteroskedasticity).

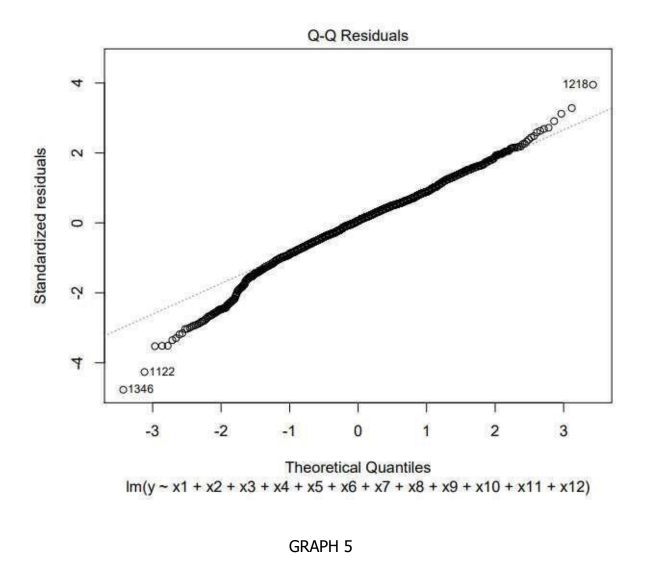
Residuals vs Leverage



GRAPH 4

The plot residual versus leverage indicates that most data points have little impact on the regression line. While there appears to be one point with potentially greater influence, without surpassing the Cook's distance threshold.

The Q-Q residual of the initial model



The scatterplot of the residuals shows a generally linear distribution with a few outliers, indicating a departure from perfect normality but not to an extreme extent. The observation that more data points lie above the line than below suggests a subtle positive skew in the residual distribution. Additionally, the Q-Q plot suggests that while the standardized residuals approximate a normal distribution, there are minor deviations from

perfect normality. Overall, while there are indications of non-normality in the residuals, they are not pronounced, implying that the model's assumptions are reasonably met.

```
> f1 = lm(y\sim x1+x2+x3+x4+x5+x6+x7+x8+x9+x10+x11+x12)
> p = 12
> n = length(y)
Call:
lm(formula = y \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 +
   x10 + x11 + x12)
Coefficients:
                    x1
x9
                                              x3
(Intercept)
                                                          x4 x5
                                                                                   x6
  6.379e+01 -2.484e-02 -6.620e-03 2.97
             x8
                                                   x11
                                       2.975e-01 -1.215e-04 -7.287e-03 1.714e-05
                                                                                          9.73
2e-02 2.293e-02 4.225e-02 -4.531e-01
                                             1.463e-04
       x12
  5.963e-09
> p
[1] 12
> n
[1] 1649
> leverage <- hatvalues(fl)
> threshold <- (2*p)/n
> threshold
[1] 0.01455428
> print (which (leverage > threshold))
12 49 74 80 81 82 83 84 86 121 133 141 145 1468 310 314 316 317 319 321 322 324 325 341 349 354 355
                                        86 121 133 141 145 148 228 229 230 231 232 2
12 49 74 80 81 82 83 84 86 121 133 141 145 14 68 310 314 316 317 319 321 322 324 325 341 349 354 355
                              83 84 86 121 133 141 145 148 228 229 230 231 232 2
 356 357 358 359 360 361 362 363 377 393 397 401 417 421 422 437 492 503 511 5
26 558 573 574 576 578 581 583 592 594 605 608 618 645
          358 359
                    360 361 362 363 377
                                            393 397
                                                      401
                                                                    422 437 492 503 511 5
26 558 573 574 576 578 581 583 592 594 605 608 618 645
 665 666 667 668 669 670 671 672 673 674 675 676
                                                                     706 712 713 719 803 8
29 837 838 844 845 846 847 848 866 871 872 874 877 879
     666 667 668 669 670 671 672 673 674 675 676
                                                           704
                                                                705 706 712 713 719 803 8
  837 838 844 845 846 847 848 866 871 872 874 877
                                                              879
 881 882 897 903 908 911 1022 1023 1054 1062 1063 1064 1088 1089 1090 1113 1114 1115 1116 11
17 1118 1119 1120 1121 1122 1128 1129 1132 1136 1188 1197 1231 1351
881 882 897 903 908 911 1022 1023 1054 1062 1063 1064 1088 1089 1090 1113 1114 1115 1116 11 17 1118 1119 1120 1121 1122 1128 1129 1132 1136 1188 1197 1231 1351
1354 1356 1357 1371 1372 1373 1374 1419 1420 1421 1422 1423 1424 1425 1426 1427 1428 1431 1432 14
33 1434 1435 1482 1483 1492 1493 1494 1501 1502 1504 1534 1535 1558
1354 1356 1357 1371 1372 1373 1374 1419 1420 1421 1422 1423 1424 1425 1426 1427 1428 1431 1432 14
33 1434 1435 1482 1483 1492 1493 1494 1501 1502 1504 1534 1535 1558
1561 1566 1573 1575 1580 1581 1596 1629 1631 1632 1639 1642 1643 1644 1645 1646 1647 1648 1649
1561 1566 1573 1575 1580 1581 1596 1629 1631 1632 1639 1642 1643 1644 1645 1646 1647 1648 1649
```

FIGURE 2

The observation number 12, 49, 74, \dots , 1649 were all identified as outliers, because their hat values are greater than 2p/n.

Non-constant variance

1. Linear relationship between the dependent and independent variables

Ho: Variance is constant

H1: Variance is not constant.

Breusch-Pagan test

FIGURE 3

We conducted a test using a Breusch-Pagan test, the Null hypothesis is rejected, the pvalue is less than 0.05 suggesting that the Variance is not constant, and we went on to do another test, White test.

White test

FIGURE 4

The Null hypothesis for the White test is rejected since the p-value is less than the 0.05 suggesting that the variance is not constant.

Normality

Normally distributed error component.

Ho: Residuals are Normally distributed

H1: Residuals are not Normally distributed.

Anderson-Darling test

```
> library(nortest)
> resids = f1$residuals
> ad.test(resids)

Anderson-Darling normality test

data: resids
A = 5.8909, p-value = 1.698e-14
```

FIGURE 5

Shapiro-Francia test

FIGURE 6

Both tests reject the null hypothesis of normally distributed residuals. Since in both test the p-value is less than 0.05, it is then suggested that the residuals are not normally distributed.

Correlation

```
> observed_values <- model1$fitted.values
> correlation <- cor(observed_values, model1$residuals)
> correlation
[1] 0.00000000000000001644619
> |
```

FIGURE 7

The correlation of the initial model is close to zero suggesting that the model predictions are not systematically biased in one direction or the other.

Correlated error (auto correction)

Ho: Residuals are uncorrelated

H1: Residuals are not uncorrelated.

Durbin-Watson test

```
> library(car)
> durbinWatsonTest(f1)
lag Autocorrelation D-W Statistic p-value
    1    0.5612034    0.8735383    0
Alternative hypothesis: rho != 0
>
```

FIGURE 8

The p-value is extremely low (p < 0.05), indicating strong evidence against the null hypothesis. Therefore, we reject the null hypothesis in favour of the alternative, suggesting that there is autocorrelation present in the residuals of your regression model for life expectancy.

The Normal Q-Q plot

Our assumptions about the model suggested that there is, non-linear relationship, heteroscedasticity, the residuals are not normally distributed which led us to go on and try to Transform the variables as well as the model. We started by using the Boxcox transformation and found out that there are little to no changes on our model.

The following are the results after we performed the BOXCOX.

```
> lambda <- bc$x[which.max(bc$y)]
> lambda
[1] 1.59596
> y2=(y^lambda-1)/lambda
> model2=lm (y2~x1+x2+x3+x4+x5+x6+x7+x8+x9+x10+x11+x12)
```

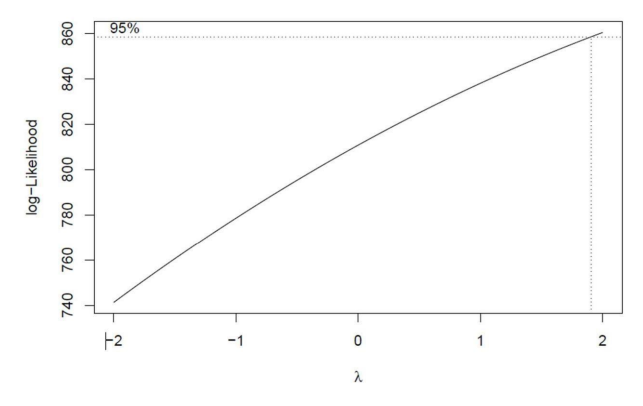
FIGURE 9

This is the summary after we performed the Box-cox transformation to our model, and we now have a new model.

```
> summary(model2)
Call:
lm(formula = y2 \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 +
   x10 + x11 + x12)
Residuals:
   Min
          1Q Median
                        3Q
                               Max
-237.981 -32.542
              2.841 33.625 237.181
Coefficients:
                        Std. Error t value
              Estimate
                                                 Pr(>|t|)
X1
X2
          X3
         -0.00097931225
X4
         X5
                                                 0.160778
                                                 0.180289
X6
          X7
          0.27746305512 0.07884942338 3.519
0.51156068639 0.08998803359 5.685
x8
                                                 0.000445
x9
                                             0.0000000155
X10
         -5.01447648302 0.27278199588 -18.383 < 0.0000000000000002
          0.00186891452 0.00043062017 4.340 0.0000151169
0.00000006950 0.00000002644 2.629 0.008648
X11
X12
(Intercept) ***
X1
         222
X2
         ***
X3
X4
X5
X6
         222
X7
x8
         222
x9
X10
         ***
         222
X11
X12
         22
Signif. codes: 0 'ana' 0.001 'an' 0.01 'a' 0.05 '.' 0.1 ' ' 1
Residual standard error: 55.37 on 1636 degrees of freedom
                        Adjusted R-squared: 0.7336
Multiple R-squared: 0.7355,
F-statistic: 379.2 on 12 and 1636 DF, p-value: < 0.0000000000000022
```

FIGURE 10

The Log-likelihood of the 2^{nd} model after we did Box-cox transformation.



GRAPH 6

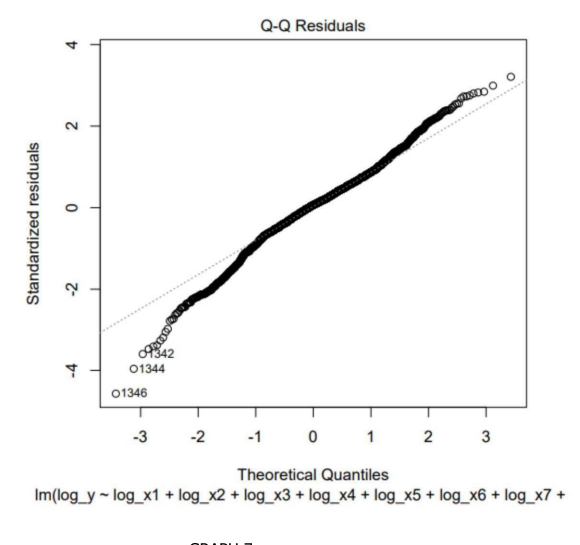
LOG TRANSFORMATION

```
> log_y <- log(mydata$y)
> log_x1 <- log(mydata$x1)
> log_x2 <- log(mydata$x2)</pre>
> log x3 <- log(mydata$x3)
> log_x4 <- log(mydata$x4)
> log x5 <- log(mydata$x5)
> log_x6 <- log(mydata$x6)
> log x7 <- log(mydata$x7)
> log x8 <- log(mydata$x8)
> log_x9 <- log(mydata$x9)
> log x10 <- log(mydata$x10)
> log_x11 <- log(mydata$x11)
> log x12 <- log(mydata$x12)
 > f2 = lm(log_y \sim log_x1 + log_x2 + log_x3 + log_x4 + log_x5 + log_x6 + log_x7 + log_x8 + log_x9 + log_x10 + log_x11 + log_x12) 
Call:
 lm(formula = log_y \sim log_x1 + log_x2 + log_x3 + log_x4 + log_x5 + log_x6 + log_x7 + log_x8 + log_x9 + log_x10 + log_x11 + log_x12) 
Coefficients:
                log_xl
                           log x2
                                       log_x3
                                                    log_x4
(Intercept)
                                                               log_x5
                                                                           log_x6
                                   log_x10
7.477e-03
             log_x8
                         log_x9
                                                log_x11
 log_x7
  4.049e+00 -1.195e-02 -1.143e-03
                                                 1.309e-02
                                                           -4.034e-03 -1.346e-03
.115e-02 3.332e-03 1.451e-02 -5.407e-02 -9.912e-05
   log x12
  6.659e-04
> summary(f2)
Call:
lm(formula = log y \sim log x1 + log x2 + log x3 + log x4 + log x5 +
     log x6 + log x7 + log x8 + log x9 + log x10 + log x11 + log x12)
Residuals:
                                          30
       Min
                    10
                          Median
-0.287705 -0.033737 0.004123 0.037379 0.202008
Coefficients:
                Estimate Std. Error t value Pr(>|t|)
(Intercept) 4.049e+00 2.109e-02 191.970 < 2e-16 ***
             -1.195e-02 1.574e-03 -7.592 5.25e-14 ***
log xl
log x2
              -1.143e-03 5.281e-04 -2.164 0.0306 *
log x3
              7.477e-03 7.821e-04
                                       9.560 < 2e-16 ***
log x4
              1.309e-02 1.763e-03 7.423 1.83e-13 ***
log x5
             -4.034e-03 2.632e-03 -1.532
                                                0.1256
log x6
             -1.346e-03 3.003e-04 -4.482 7.93e-06 ***
log x7
              1.115e-02 2.133e-03 5.229 1.93e-07 ***
log x8
              3.332e-03 2.981e-03 1.118 0.2638
              1.451e-02 3.345e-03 4.337 1.53e-05 ***
log x9
             -5.407e-02 1.149e-03 -47.042 < 2e-16 ***
log x10
              -9.912e-05 2.107e-03 -0.047
                                                 0.9625
log x11
              6.659e-04 6.544e-04 1.018 0.3090
log_x12
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Residual standard error: 0.06334 on 1636 degrees of freedom
Multiple R-squared: 0.7813,
                                    Adjusted R-squared: 0.7797
F-statistic: 487 on 12 and 1636 DF, p-value: < 2.2e-16
```

FIGURE 11

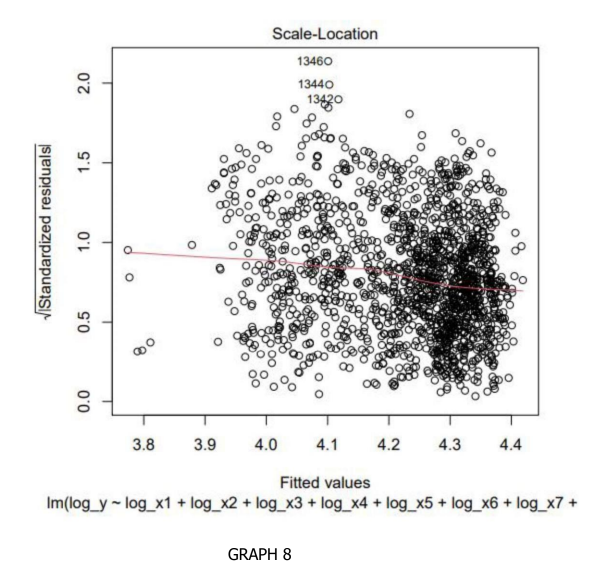
The residuals exhibit a scatter around zero, indicating a generally good fit between the model and the data. Although there are a few outliers suggesting instances where the model's fit is less accurate, their limited number suggests they have a minor impact on overall results. Notably, most residuals are positive, indicating a tendency for the model to underestimate actual values. This slight positive skew in the residuals suggests a consistent pattern of underestimation by the model. Despite these observations, most data points conform well to the model, implying its effectiveness in capturing the underlying relationships within the data.

Q-Q Residual of Log-Transformation



GRAPH 7

Scale-Location of Log-Transformation



After we fitted the 3rd model using log Transformation, we then proceeded to test our 3rd model to see if is adequate to be used or not.

We started by using Breusch-Pagan test and White test to test for heteroscedasticity in our fitted model.

The Breusch-Pagan test and White test are both used to detect heteroscedasticity in regression models.

- Null Hypothesis (Ho): There is homoscedasticity in the life expectancy regression model.
- Alternative Hypothesis (H1): There is heteroscedasticity in the life expectancy regression model.

FIGURE 12

The Breusch-Pagan test for 3rd model indicates significant evidence against the null hypothesis of homoscedasticity, suggesting that the residuals in 3rd model exhibit heteroscedasticity. This is supported by a low p-value of 2.091e-11. Similarly, the White test for 3rd model also provides significant evidence against the null hypothesis of homoscedasticity, with an even lower p-value of < 2.2e-16. In summary, both tests confirm that the 3rd model suffers from heteroscedasticity, indicating that the variability of life expectancy is not constant across different values of the independent variables in the model.

The following graphs were plotted after we did a transformation in our variables against fitted.

The Anderson-Darling and Shapiro-Francia tests were conducted on the residuals of the life expectancy regression model to assess their normality.

• Null Hypothesis (Ho): The residuals follow a normal distribution.

 Alternative Hypothesis (H1): The residuals do not follow a normal distribution. Anderson-Darling and Shapiro-Francia test

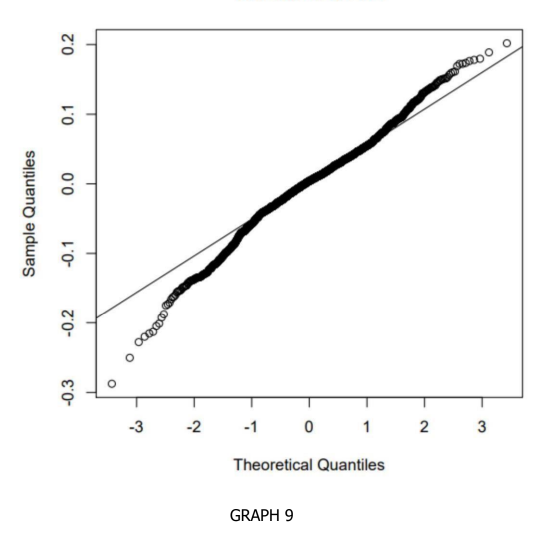
FIGURE 13

The normality of the residuals for 3rd model was assessed using the Anderson-Darling and Shapiro-Francia tests. Both tests indicated significant evidence against the null hypothesis of normality. The Anderson-Darling test yielded a p-value that is less than 2.2e-16, while the Shapiro Francia test resulted in a p-value of 2.107e-10. These results suggest that the residuals of the 3rd model deviate significantly from a normal distribution.

The following is the graph after we did log Transformation and obtained 3rd model.

Normal Q-Q Plot of Log-Transformation





Since the plotted points are not in a straight line, the residuals appear to be not normally distributed.

We then again did an Autocorrelation test.

The Durbin-Watson test assesses the presence of autocorrelation in the residuals of a regression model.

• Null Hypothesis (Ho): There is no autocorrelation present in the residuals (rho = 0).

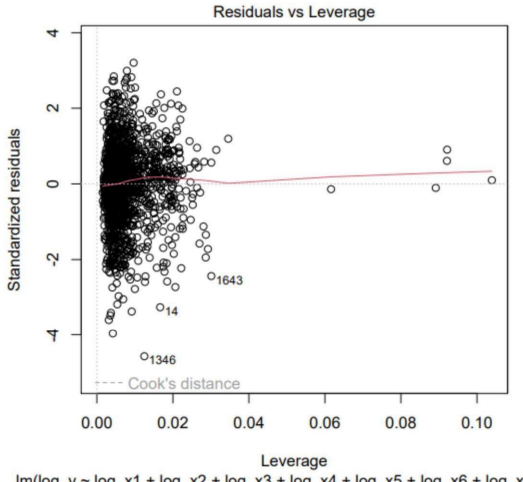
Alternative Hypothesis (H1): There is autocorrelation present in the residuals (rho ≠ 0).
 Durbin-Watson test

```
> library(car)
> durbinWatsonTest(f2)
lag Autocorrelation D-W Statistic p-value
    1    0.6915865    0.6162016    0
Alternative hypothesis: rho != 0
>
```

FIGURE 14

The Durbin-Watson test was conducted on the 3rd model to assess autocorrelation in the residuals. The test yielded a Durbin-Watson statistic of 0.6162016 with a corresponding p-value of 0, indicating significant evidence against the null hypothesis of no autocorrelation. This result suggests that there is autocorrelation present in the residuals of 3rd model.

Residuals vs Leverage of Log-Transformation



 $Im(log_y \sim log_x1 + log_x2 + log_x3 + log_x4 + log_x5 + log_x6 + log_x7 + log_x6 + log_x7 + log_x8 +$

GRAPH 10

Weighted least squares (WLS)

```
> fl
Call:
lm(formula = y \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 +
    x10 + x11 + x12)
Coefficients:
                                                                                      x5
                                                                                                     x6
(Intercept)
                         x1
                                         x2
                                                        x3
                                                                       \times 4
                  x8
                                                x10
   x7
                                 x9
                                                              x11
  6.379e+01
                -2.484e-02
                               -6.620e-03
                                                2.975e-01
                                                             -1.215e-04
                                                                            -7.287e-03
                                                                                             1.714e-05
                                                                                                            9.73
2e-02
         2.293e-02
                        4.225e-02
                                       -4.53le-01
                                                        1.463e-04
         x12
  5.963e-09
> x=abs(rnorm(100,0,3))
> y=3-2*x+rnorm(100,0,sapply(x,function(x){1+0.5*x^2}))
> plot(x,y,pch=16,cex=0.5)
> abline(a=3,b=-2,col="black")
> fit.ols=lm(y~x)
> fit.wls=lm(y~x, weights=1/(1+0.5*x^2))
> abline(fit.wls$coefficients,lty=3)
> legend("topright",c("ols","wls"),lty=c(2,3))
                                                                                       ols
                                                                                       wls
       20
       10
```

6

The coefficients of the two lines are shown below.

2

X

0

The comparison of coefficients between the OLS and WLS models reveals notable differences, particularly with larger intercept and slope values in the WLS model. This suggests a distinct approach in fitting the data, potentially addressing heteroscedasticity or unequal variances. While OLS assumes equal variance across all observations, WLS adjusts by assigning weights based on variance, giving more weight to observations with lower variance. Therefore, the WLS model places greater emphasis on certain data points, aiming to provide a more accurate fit by mitigating the effects of heteroscedasticity.

Variable selection

Backward Selection

```
> f1 = lm(y \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 + x10 + x11 + x12)
> summary(f1)
lm(formula = y \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 +
   x10 + x11 + x12)
Residuals:
            1Q Median
    Min
                              30
                                        Max
-21.2824 -2.5064 0.2694 2.7378 17.6119
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) 6.379e+01 6.194e-01 102.982 < 2e-16 ***
           -2.484e-02 1.128e-03 -22.012 < 2e-16 ***
x1
          -6.620e-03 1.434e-03 -4.617 4.19e-06 ***
x2
           2.975e-01 3.244e-02 9.171 < 2e-16
-1.215e-04 2.230e-04 -0.545 0.585938
x3
                                  9.171 < 2e-16 ***
x4
           -7.287e-03 5.515e-03 -1.321 0.186555
x5
           1.714e-05 1.297e-05 1.321 0.186553
            9.732e-02 6.488e-03 14.999 < 2e-16 ***
x7
                                 3.601 0.000326 ***
            2.293e-02
x8
                       6.369e-03
                                   5.813 7.37e-09 ***
x9
            4.225e-02 7.268e-03
          -4.531e-01 2.203e-02 -20.565 < 2e-16 ***
x10
            1.463e-04 3.478e-05 4.206 2.74e-05 ***
x11
x12
           5.963e-09 2.135e-09 2.792 0.005296 **
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' 1
Residual standard error: 4.472 on 1636 degrees of freedom
Multiple R-squared: 0.7434, Adjusted R-squared: 0.7415
F-statistic:
             395 on 12 and 1636 DF, p-value: < 2.2e-16
```

We fit model with all the parameters, we show estimation details then we remove x4, x5 and x6 because the p-values of these predictor is greater than 0.10.

```
 > f2 = 1m(y \sim x1 + x2 + x3 + x7 + x8 + x9 + x10 + x11 + x12) 
> summary(f2)
lm(formula = y \sim x1 + x2 + x3 + x7 + x8 + x9 + x10 + x11 + x12)
Residuals:
             10 Median
    Min
                               30
                                       Max
-21.3473 -2.5123 0.3134 2.7690 17.4916
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) 6.368e+01 6.133e-01 103.829 < 2e-16 ***
           -2.495e-02 1.127e-03 -22.139 < 2e-16 ***
x1
           -5.616e-03 1.288e-03 -4.362 1.37e-05 ***
x2
            3.003e-01 3.241e-02 9.267 < 2e-16 ***
x3
x7
            9.672e-02 6.477e-03 14.933 < 2e-16 ***
           2.185e-02 6.279e-03 3.479 0.000516 ***
x8
            3.829e-02 6.539e-03 5.855 5.76e-09 ***
x9
x10
          -4.518e-01 2.201e-02 -20.525 < 2e-16 ***
x11
           1.289e-04 1.098e-05 11.741 < 2e-16 ***
x12
           5.859e-09 2.131e-09 2.749 0.006046 **
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Residual standard error: 4.473 on 1639 degrees of freedom
Multiple R-squared: 0.7428,
                             Adjusted R-squared: 0.7414
F-statistic: 526 on 9 and 1639 DF, p-value: < 2.2e-16
>
```

After we remove x4, x5 and x6, we fit the model again. We stop the elimination because the pvalues of a remaining predictor is less than 0.1.

Mallows C_p statistic

```
> library(leaps)
> x = model.matrix(f1)[,-1]
> y = y
> f1 = leaps(x,y)
Swhich
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```
Slabel
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 [1] "(Intercept)" "1"
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                                                                                            "5"
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                                         "9"
       "7" "8"
                                                            "A"
                       "C"
Ssize
  [55]
[109] 12 12 13
$Cp
   [1] 1584.34944 2494.85208 2857.88508 3489.42023 3661.37328 3697.17853 3988.41626 4048.26029 447
6.41296 4549.02730 952.88533 1072.75952 1117.65474 1188.27618
  [15] 1200.38790 1303.15449 1335.68879 1354.36584 1462.48346 1483.96643 590.33501 606.39542 66
3.48121 698.15137 705.58730 714.84278 760.52806 768.13910 [29] 807.37484 831.84733 269.99442 288.87887 312.11035 404.45966 426.13327 442.25730 45 7.61055 476.69149 502.03337 503.48730 126.86590 141.26357
 [43] 157.72588 158.31521 164.43708 179.48448 188.77387 206.16465 228.68015 246.68753 3
8.02057 56.68904 63.19154 81.39104 108.61458 110.00394
[57] 116.68241 119.02095 128.30722 128.54673 25.94430 26.95482 39.39877 39.64424 3 9.89629 39.98191 43.23057 45.36696 50.35396 58.04173 [71] 16.39888 20.95122 27.23117 27.39132 27.43192 27.75488 27.87420 28.29474 2 8.56067 33.53107 10.83889 16.42411 16.94840 18.18874 [85] 20.95027 22.38773 22.51654 28.02287 28.57663 28.77987 10.97629 11.05817 1 2.59678 17.06276 18.14439 18.74879 22.44006 22.53339 [99] 23.89373 28.15810 11.29685 12.74608 12.74610 18.79646 23.96764 28.69228 3 2.31816 44.78934 95.09930 235.96706 13.00000
 > f1 = lm(y \sim x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9 + x10 + x11 + x12)
 > library(olsrr)
 > # Backward variable selection using AIC
 > ols step backward aic(fl)
```

Stepwise Summary

Step	Variable	AIC	SBC	SBIC	R2	Adj. R2
0	Full Model	9634.613	9710.324	4955.160	0.74342	0.74154
1	×4	9632.912	9703.215	4953.429	0.74338	0.74165
2	x5	9632.604	9697.499	4953.092	0.74311	0.74155
3	x6	9632.478	9691.965	4952.941	0.74282	0.74141

x4, x5, x6 eliminated.

- > # Forward variable selection using AIC
- > ols_step_forward_aic(f1)

Stepwise Summary

Step	Variable	AIC	SBC	SBIC	R2	Adj. R2
0	Base Model	11853.803	11864.619	7171.562	0.00000	0.00000
1	x1	10733.977	10750.201	6051.881	0.49354	0.49323
2	x7	10375.913	10397.544	5693.799	0.59289	0.59239
3	x10	10128.373	10155.413	5446.490	0.65006	0.64942
4	x11	9873.091	9905.538	5192.032	0.70061	0.69988
5	x9	9744.706	9782.562	5064.257	0.72337	0.72253
6	x3	9659.512	9702.775	4979.651	0.73762	0.73666
7	x8	9647.579	9696.250	4967.824	0.73983	0.73872
8	x2	9638.063	9692.142	4958.421	0.74164	0.74038
9	x12	9632.478	9691.965	4952.930	0.74282	0.74141

x1, x7, x10, x11, x9, x3, x8, x2, x12 selected.

- > # Forward variable selection using AIC
 > ols_step_forward_aic(f1)

Stepwise Summary

Step	Variable	AIC	SBC	SBIC	R2	Adj. R2
0	Base Model	11853.803	11864.619	7171.562	0.00000	0.00000
1	x1	10733.977	10750.201	6051.881	0.49354	0.49323
2	x7	10375.913	10397.544	5693.799	0.59289	0.59239
3	x10	10128.373	10155.413	5446.490	0.65006	0.64942
4	x11	9873.091	9905.538	5192.032	0.70061	0.69988
5	x9	9744.706	9782.562	5064.257	0.72337	0.72253
6	x3	9659.512	9702.775	4979.651	0.73762	0.73666
7	x8	9647.579	9696.250	4967.824	0.73983	0.73872
8	x2	9638.063	9692.142	4958.421	0.74164	0.74038
9	x12	9632.478	9691.965	4952.930	0.74282	0.74141

DISCUSSIONS

Adult mortality has an inverse relationship with life expectancy, indicating that higher adult mortality rates are associated with shorter average lifespans. Additionally, when infant mortality rates are high, life expectancy is reduced, reflecting a negative relationship.

There is a positive relationship between polio immunization coverage and life expectancy, as people who are immunized tend to live longer compared to those who contract polio without immunization.

People with HIV/AIDS have shorter lifespans, as indicated by our data, which focuses on the prevalence of the virus rather than immunization coverage. This results in a negative relationship with life expectancy, as those infected by the virus tend to die at a younger age. For instance, in South Africa from 2003 to 2008, the high mortality rates among both children and adults due to HIV/AIDS significantly reduced life expectancy.

A country's higher Gross Domestic Product (GDP) indicates that its citizens are likely to enjoy longer life spans, as increased economic production enables the country to invest in healthier initiatives and choices.

Multilinear model assumptions Findings

Our model does not meet the following.

Despite employing Box-cox and Log transformations, the residuals still do not follow a normal distribution. Furthermore, they are not evenly distributed around the line of best fit, as depicted in the figure. This deviation from normality can lead to underestimated standard errors for regression coefficients, inaccurate t-statistics, p-values, and ultimately, incorrect projected values. Given these violations of multi-linear regression assumptions, it is highly advised that we interpret the conclusions of our study with caution. The Heteroscedasticity remains present in our data.

CONCLUSIONS

The multilinear regression model was used to predict the Life expectancy of the country using social, environmental, and health-related factors influencing it. The factors/predictors that were used are as follows, adult mortality, Infant deaths, Alcohol, percentage expenditure, Polio, Hepatitis B, BMI, Measles, Population, Diphtheria, HIV/AIDS, and GDP to predict how they influence the life expectancy. Our predictors are under the significant level of 0.05.

The final model is.

Life expectancy = 6.368e+01 - 2.495e-02x15.616e-03x2 + 3.003e-01x3 + 9.672e-02x7 + 2.185e-02x8 + 3.829e-02x9 - 4.518e-01x10 + 1.289e-04x11 + 5.859e-09x12

Multiple R-squared is 74.3%, Adjusted R-squared is 74.2% after we refined the model,

This means that our model is significant since our Multiple R-square is 74.3%, we cannot really trust the model to do accurate predictions, for instance,

let's predict the life expectancy in Afghanistan in 2005, the country is developing.

Adult mortality = 280

Infant deaths = 80

Alcohol = 4.7

Percentage expenditure = 79.67936736

Hepatitis = 66

Measles = 1990

BMI = 55.8

Polio = 0

Diphtheria = 0

HIV/AIDS = 0.1

GDP = 189.681557

Population = 2.966463e6

Life expectancy =6.368e+01 -2.495e-02(280)-5.616e-03(80) +3.003e-01(4.7) +9.672e-02(55.8)+2.185e-02(0)+3.829e-02(0)-4.518e-01(0.1)+1.289e-04(189.681557)+5.859e-09(2.966463e6)

Life expectancy = 63 years

HIV/AIDS: HIV/AIDS is a life-threatening chronic illness. Countries with higher HIV/AIDS prevalence have shorter life expectancy. A clear result is the inverse link between adult mortality and life expectancy. A greater adult mortality rate corresponds to a shorter life expectancy. This is because a greater adult mortality rate suggests that the country has more health issues.

The more a country have immunization coverage against polio and diphtheria, the longer people live.

Despite refinement and transformation, the final model does not meet our assumptions. Hence, the residuals are not normally distributed, and the model exhibits heteroscedasticity, and there is no multicollinearity between independent variables. To ensure the accuracy of the p-value and t-statistic, the residuals in the final model are to follow a normal distribution around the fitted line. However, in our analysis, the residuals not only deviate from the fitted line but also show significant dispersion, indicating a lack of normality. This discrepancy may result in incorrect conclusions and needs careful consideration. These are only a few of the significant social, economic, and health-related factors that may be utilized to create a multiple linear regression model to forecast a country's life expectancy. The precise predictors that are most essential may differ based on the nation and the available data. This can result in underestimated standard errors for regression coefficients, incorrect t-statistics and p-values, and, ultimately, incorrect projected values.

GLOSSARY

Life expectancy: refers to the number of years a person is expected to live based on statistical average.

Adult Mortality: refers to the death rate among individuals within a specific age range typically considered to be adults, often defined as individuals aged 15 to 64 or 15 to 69 years old, depending on the context

Infant Deaths: Infant mortality refers to the death of a babies that occurs between the time it is born and 1 year of age.

Quinquennial: refers to a five-year period or event.

Percentage expenditure: refers to the portion or share of total spending allocated to a particular category, activity, or item, expressed as a percentage of the overall budget or expenditure.

Hepatitis B: this is a viral infection that primarily affects the liver. It is caused by the hepatitis B virus (HBV), which is transmitted through contact with the blood or other body fluids of an infected person

Measles: Measles is a very contagious disease that causes fever, a red rash, cough and watery eyes and mostly common in children.

BMI: BMI is used to estimate your body fat and establish whether you are underweight, at your healthy weight, overweight, or have obesity.

Polio immunization coverage: refers to the proportion of individuals within a population who have received the polio vaccine, typically measured as a percentage

Diphtheria immunization coverage: refers to the percentage of individuals within a population who have received vaccination against diphtheria

HIV/AIDS: this is a viral infection that weakens the immune system, making individuals more susceptible to various infections and diseases.

Variable changed name description.

Variable name	Renamed variable name
Life expectancy	У
Adult mortality	X1
Infant death	X2
Alcohol	X3
Percentage expenditure	X4
Hepatitis B	X5
Measles	X6
BMI	X7
Polio	X8
Diphtheria	Х9
HIV/AIDS	X10
GDP	X11
Population	X12

REFERENCES

Life expectancy dataset retrieved from the following GitHub website.

https://github.com/selva86/datasets/blob/master/Life_Expectancy_Data.csv

programming language for analysis

R and RStudio

Research was done with the help of the following websites and books.

https://en.wikipedia.org/wiki/Cook%27s_distance_https://onlinestatbook.com/2/transformations/box-

cox.html https://www.statology.org/how-to-calculate-mallows-cp-in-r/

https://stattrek.com/regression/influential-points#google_vignette

https://book.stat420.org/transformations.html#response-transformations Books

STA37W1 Linear Model notes

Applied Linear Model statistical analysis 5th edition by Richad A Johnson, Dean W. Wichern